Package 'immunarch'

October 15, 2025

```
Type Package
Title Multi-Modal Immune Repertoire Analytics for Immunotherapy and
     Vaccine Design in R
Version 0.10.3
Contact support@immunomind.com
Description A comprehensive analytics framework for building reproducible pipelines on T-
     cell and B-cell immune receptor repertoire data.
     Delivers multi-modal immune profiling (bulk, single-cell, CITE-
     seq/AbSeq, spatial, immunogenicity data),
     feature engineering (ML-ready feature tables and matrices), and biomarker discovery workflows
     (cohort comparisons, longitudinal tracking, repertoire similarity, enrichment).
     Provides a user-friendly interface to widely used AIRR methods —
     clonality/diversity, V(D)J usage, similarity, annotation, tracking, and many more.
     Think Scanpy or Seurat, but for AIRR data, a.k.a. Adaptive Immune Receptor Repertoire, VDJ-
     seq, RepSeq, or
     VDJ sequencing data. A successor to our previously published ``tcR" R package (Nazarov 2015).
License Apache License (>= 2.0)
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```

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Description

Get a column's name using the input alias

Usage

```
.quant_column_choice(x)
```

Arguments

x Character vector of length 1.

Value

A string with the column name.

Developer Examples

immunarch:::.quant_column_choice("count") immunarch:::.quant_column_choice("freq")

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aa_properties

Tables with amino acid properties

Description

Tables with amino acid properties

aa_table

Amino acid / codon table

Description

Amino acid / codon table

Usage

AA_TABLE

Format

An object of class table of length 65.

add_class

Add a new class attribute

Description

Add a new class attribute

Usage

```
add_class(.obj, .class)
```

Arguments

.obj

R object.

.class

String with the desired class name.

Value

Input object with additional class .class.

Developer Examples

```
tmp <- "abc" class(tmp) tmp <- immunarch:::add_class(tmp, "new_class") class(tmp)</pre>
```

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airr_clonality

Clonality - receptor overabundance statistics for immune repertoires

Description

[Experimental]

A family of functions to quantify **receptor overabundance** per repertoire. Helps in deciphering the structure and partition the repertoire.

Available functions:

Supported methods are the following.

airr_clonality_line - build ranked abundance lines: for each repertoire, take the top limit receptors by count and attach repertoire metadata. Useful for per-repertoire rank-abundance plots.

airr_clonality_rank - aggregate clonal space by **rank bins**. Receptors are ordered by proportion within each repertoire; each receptor is assigned to the smallest threshold in bins that contains its rank.

airr_clonality_prop - aggregate clonal space by **proportion bins**. Each receptor is assigned to a named bin according to its proportion (e.g., Hyperexpanded >= 1e-2, Large >= 1e-3, ...). Thresholds are matched in descending order; unmatched receptors fall into "Ultra-rare".

Usage

```
airr_clonality_line(
  idata,
  limit = 1e+05,
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_clonality_rank(
  idata,
 bins = c(10, 30, 100, 300, 1000, 10000, 1e+05),
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_clonality_prop(
  idata,
 bins = c(Hyperexpanded = 0.01, Large = 0.001, Medium = 1e-04, Small = 1e-05, Rare =
 autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
```

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Arguments

idata An ImmunData object.

limit Positive integer >= 10: maximum number of top receptors to keep **per reper**

toire (default 100000).

autojoin Logical. If TRUE, join repertoire metadata by the schema repertoire id. Change

the default behaviour by calling options(immunarch.autojoin = FALSE).

format String. One of "long" ("long" tibble with imd_repertoire_id, facet columns,

and value; useful for visualizations) or "wide" (wide/unmelted table of features, with each row corresponding to a specific repertoire / pair of repertoires;

useful for Machine Learning).

bins A **named** numeric vector of thresholds (e.g., c(Hyperexpanded = 1e-2, Large

= 1e-3, ...)). Names become bin labels and must be non-empty. Internally

sorted in descending order.

Value

airr_clonality_line:

A tibble with columns:

- repertoire_id repertoire identifier
- index rank within repertoire (1 = most abundant)
- count receptor count used for ranking
- plus any repertoire metadata columns carried from idata\$repertoires

airr_clonality_rank:

A tibble with

- repertoire_id
- clonal_rank_bin the rank threshold (e.g., 10, 100, ...)
- occupied_prop sum of proportion within the bin
- plus repertoire metadata columns from idata\$repertoires

airr_clonality_prop:

A tibble with

- repertoire_id
- clonal_prop_bin factor-like label from names(bins) or "Ultra-rare"
- occupied_prop sum of proportion within the bin
- plus repertoire metadata columns from idata\$repertoires

See Also

• Per-repertoire summaries: annotate clonality

• Data container: immundata::ImmunData

Examples

```
# Limit the number of threads used by the underlying DB for this session.
# Change this only if you know what you're doing (e.g., multi-user machines, shared CI/servers).
db_exec("SET threads TO 1")
# Load data
## Not run:
immdata <- get_test_idata() |> agg_repertoires("Therapy")
## End(Not run)
# airr_clonality_line
## Not run:
top_line <- airr_clonality_line(immdata, limit = 1000)</pre>
## End(Not run)
# airr_clonality_rank
## Not run:
rank_stat <- airr_clonality_rank(immdata, bins = c(10, 100))</pre>
## End(Not run)
# airr_clonality_prop
## Not run:
prop_stat <- airr_clonality_prop(immdata)</pre>
## End(Not run)
```

airr_diversity

Diversity - estimating the heterogeneity of immune repertoires

Description

[Experimental]

A family of functions to quantify **receptor diversity** per repertoire. A characteristic of a whole repertoire.

Available functions:

Supported methods are the following.

airr_diversity_dxx - **coverage diversity**: minimal number of top receptors needed to reach perc% of clonal space (by proportion). Great for spotting dominance/overexpansion and for quick, interpretable dashboards (e.g., D50 = receptors to cover half of the repertoire).

airr_diversity_chao1 - Chao1 estimator is a nonparameteric asymptotic estimator of species richness (number of species in a population). One of the most used methods for estimating immune repertoire diversity.

airr_diversity_shannon - Shannon entropy (base 2) per repertoire computed from proportion. Ideal when you want a single evenness-aware diversity score; pair with Pielou/Hill for samples with very different richness.

airr_diversity_pielou - Pielou's evenness H / log2(S) with richness S. Best when you need a **size-normalized** evenness score that's comparable across repertoires with different receptor counts.

airr_diversity_index - convenience alias for Hill number with q = 1 (exp(Shannon) using natural log). A solid **default single metric** that's relatively robust to rare-count noise and easy to compare across samples.

airr_diversity_hill - Hill numbers ("true diversity") for orders q \eqn{\in}{in} {0, 1, 2, ...}: q=0 richness, q=1 exp(Shannon), q>1 emphasizes abundant receptors. Perfect when you want a **diversity profile** that tunes sensitivity to rare vs. abundant clonotypes.

Usage

```
airr_diversity_dxx(
  idata,
  perc = 50,
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_diversity_chao1(
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_diversity_shannon(
  idata,
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_diversity_pielou(
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_diversity_index(
  idata,
```

```
autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_diversity_hill(
  idata,
 q = 0:5,
 autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
```

Arguments

format

idata An ImmunData object.

perc A number or numeric vector in (0, 100] (default 50), e.g. 50 for D50, 20 for

autojoin Logical. If TRUE, join repertoire metadata by the schema repertoire id. Change

the default behaviour by calling options(immunarch.autojoin = FALSE).

String. One of "long" ("long" tibble with imd_repertoire_id, facet columns,

and value; useful for visualizations) or "wide" (wide/unmelted table of features, with each row corresponding to a specific repertoire / pair of repertoires;

useful for Machine Learning).

A scalar or vector of non-negative orders. Defaults to 0:5. q

Value

airr_diversity_dxx:

A tibble with:

- imd_repertoire_id
- perc
- dxx minimal count of top receptors to reach perc%
- plus repertoire metadata from idata\$repertoires

airr_diversity_chao1:

A tibble with:

- imd_repertoire_id
- Estimator number of species
- SD standard deviation for the estimator value
- Conf. 95. lo CI 0.025
- Conf.95.hi CI 0.975
- plus repertoire metadata from idata\$repertoires

airr_diversity_shannon:

A tibble with:

• imd_repertoire_id

• shannon - entropy in bits

airr_diversity_pielou:

A tibble with:

- imd_repertoire_id
- shannon
- n_receptors
- pielou evenness in [0, 1] (NA if S <= 1)

airr_diversity_index:

A tibble with:

- imd_repertoire_id
- q = 1
- hill_number
- plus repertoire metadata from idata\$repertoires

airr_diversity_hill:

A tibble with:

- imd_repertoire_id
- q Hill order
- hill_number true diversity of order q
- plus repertoire metadata from idata\$repertoires

See Also

immundata::ImmunData

```
# Limit the number of threads used by the underlying DB for this session.
# Change this only if you know what you're doing (e.g., multi-user machines, shared CI/servers).
db_exec("SET threads TO 1")
# Load data
## Not run:
immdata <- get_test_idata() |> agg_repertoires("Therapy")

## End(Not run)

#
# airr_diversity_dxx
#
## Not run:
d50 <- airr_diversity_dxx(immdata, perc = 50)
d_multi <- airr_diversity_dxx(immdata, perc = c(20, 50, 80))

## End(Not run)

## End(Not run)</pre>
```

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```
# airr_diversity_chao1
## Not run:
chao <- airr_diversity_chao1(immdata)</pre>
## End(Not run)
# airr_diversity_shannon
## Not run:
sh <- airr_diversity_shannon(immdata)</pre>
## End(Not run)
# airr_diversity_pielou
## Not run:
pj <- airr_diversity_pielou(immdata)</pre>
## End(Not run)
# airr_diversity_index
## Not run:
idx <- airr_diversity_index(immdata)</pre>
## End(Not run)
# airr_diversity_hill
## Not run:
hill \leftarrow airr_diversity_hill(immdata, q = c(0, 1, 2))
## End(Not run)
```

airr_public

Public indices - pairwise repertoire overlap

Description

[Experimental]

A family of functions to quantify **public or shared receptors** between repertoire.

Available functions:

Supported methods are the following.

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airr_public_intersection - number of **shared receptors** between each pair of repertoires (intersection size). Handy for quick overlap heatmaps, QC of replicate similarity, or spotting donor-shared "public" clonotypes.

airr_public_jaccard - **Jaccard similarity** of receptor sets between repertoires $(A \cap B \mid A \cup B)$. Best when comparing cohorts with different sizes to get a scale-invariant overlap score.

Usage

```
airr_public_intersection(
   idata,
   autojoin = getOption("immundata.autojoin", TRUE),
   format = c("long", "wide")
)
airr_public_jaccard(
   idata,
   autojoin = getOption("immundata.autojoin", TRUE),
   format = c("long", "wide")
)
```

Arguments

idata An ImmunData object.

autojoin Logical. If TRUE, join repertoire metadata by the schema repertoire id. Change

the default behaviour by calling options(immunarch.autojoin = FALSE).

format String. One of "long" ("long" tibble with imd_repertoire_id, facet columns,

and value; useful for visualizations) or "wide" (wide/unmelted table of features, with each row corresponding to a specific repertoire / pair of repertoires;

useful for Machine Learning).

Value

airr_public_intersection:

A **symmetric numeric matrix** where rows/columns are repertoire_id and each cell is the count of shared unique receptors. The diagonal contains per-repertoire richness (total unique receptors). Row/column names are repertoire IDs.

```
airr_public_jaccard:
```

A **symmetric numeric matrix** where rows/columns are repertoire_id and each cell is the Jaccard similarity in [0, 1]. The diagonal is 1. Row/column names are repertoire IDs.

See Also

immundata::ImmunData

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Examples

```
# Limit the number of threads used by the underlying DB for this session.
# Change this only if you know what you're doing (e.g., multi-user machines, shared CI/servers).
db_exec("SET threads TO 1")
# Load data
immdata <- get_test_idata() |> agg_repertoires("Therapy")

# 
# airr_public_intersection
# 
## Not run:
m_pub <- airr_public_intersection(immdata)

## End(Not run)

# 
# airr_public_jaccard
# 
## Not run:
m_jac <- airr_public_jaccard(immdata)

## End(Not run)</pre>
```

airr_stats

Compute key immune repertoire statistics

Description

[Experimental]

A family of functions that extract core descriptive statistics from an ImmunData object.

Available functions:

Supported methods are the following.

airr_stats_chains — count V(D)J *chains* per repertoire (optionally split by locus). Quickly gauges capture depth per repertoire and, when split by locus, reveals TRA/TRB/IGH balance. Use it for QC, library-size checks, and to spot locus-specific dropouts or over-representation.

airr_stats_lengths — count the number of sequence lengths per repertoire. Summarizes the CDR3 length distribution, a sensitive QC fingerprint of repertoire prep and selection. Helpful for detecting primer/UMI biases, comparing cohorts, and deriving length-based features for models.

airr_stats_genes - count V(D)J gene segments per repertoire, optionally split by locus and using either receptor counts or barcode/UMI counts as the measure. Profiles V/D/J gene usage to characterize repertoire composition and germline biases, with optional locus split. Useful for cohort comparisons, flagging clonal expansions, and producing ML-ready features for repertoire-level ML tasks.

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Usage

```
airr_stats_chains(
 idata,
  locus_col = NA,
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
airr_stats_lengths(
  idata,
  seq_col = "cdr3_aa",
 autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
airr_stats_genes(
  idata,
  gene_col = "v_call",
 level = c("receptor", "barcode"),
 by = c(NA, "locus"),
 autojoin = getOption("immundata.autojoin", TRUE),
 format = c("long", "wide")
)
```

Arguments

idata	An ImmunData object.
locus_col	Column in idata\$annotations that stores the locus (e.g. "locus"). If NULL or missing, the result is not split by locus.
autojoin	Logical. If TRUE, join repertoire metadata by the schema repertoire id. Change the default behaviour by calling options(immunarch.autojoin = FALSE).
format	String. One of "long" ("long" tibble with imd_repertoire_id, facet columns, and value; useful for visualizations) or "wide" (wide/unmelted table of features, with each row corresponding to a specific repertoire / pair of repertoires; useful for Machine Learning).
seq_col	Character vector with names of the columns containing sequences.
gene_col	A single column name in idata\$annotations with gene segment calls (e.g., "v_call", "d_call", "j_call", "c_call"). Default is "v_call".
level	One of "receptor" or "barcode". If "receptor" (default), the function counts unique receptors (one per receptor ID) that carry a given gene segment. If "barcode", the function sums counts (e.g., cells/UMIs) per gene segment using the column defined by immundata::imd_schema("count").
by	Either NULL (no split) or "locus". When "locus", the result is further split by the locus column if present (as given by immundata::imd_schema("locus"));

otherwise a warning is emitted and the split is ignored.

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Value

```
airr_stats_chains Returns a tibble with columns::
```

- repertoire_id repertoire identifier
- locus TRA, TRB, IGH, ... (present only if locus_col is supplied)
- n_chains number of chains

airr_stats_lengths Returns a tibble with columns::

- repertoire_id repertoire identifier
- seq_len lengths of sequences
- n number of receptors

airr_stats_genes A tibble with columns::

- repertoire_id repertoire identifier
- (optional) locus TRA, TRB, IGH, ... (present only when by = "locus" and the locus column exists)
- <gene_col> the gene segment value (e.g., V gene)
- n the measure:
 - if level = "receptor": number of receptors carrying the gene segment
 - if level = "barcode": sum of counts across receptors for the segment

See Also

immundata::ImmunData

```
# Limit the number of threads used by the underlying DB for this session.
# Change this only if you know what you're doing (e.g., multi-user machines, shared CI/servers).
db_exec("SET threads TO 2")

# Load data
## Not run:
immdata <- get_test_idata() |> agg_repertoires("Therapy")

## End(Not run)

# # airr_stats_chains
#
## Not run:
airr_stats_chains(immdata)

## End(Not run)

# # airr_stats_lengths
# airr_stats_lengths
# airr_stats_lengths
```

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```
## Not run:
airr_stats_lengths(immdata)

## End(Not run)

# 
# airr_stats_genes

## Not run:

# V gene usage by receptor count
airr_stats_genes(immdata, gene_col = "v_call", level = "receptor")

# V gene usage by summed cell/UMI counts (if a count column is present)
airr_stats_genes(immdata, gene_col = "v_call", level = "barcode")

# Split by locus (TRA/TRB/... if locus column exists)
airr_stats_genes(immdata, gene_col = "v_call", level = "receptor", by = "locus")

## End(Not run)
```

annotate_clonality

Annotate clonality - per-receptor labels for overabundance

Description

[Experimental]

A small family of helpers that **add clonality labels to each receptor** in an immundata::ImmunData object.

Available functions:

- annotate_clonality_rank() label by rank bins within each repertoire.
- annotate_clonality_prop() label by **proportion bins** (named thresholds).

annotate_clonality_rank() - for each repertoire, receptors are ordered by within-repertoire abundance (proportion) and assigned a **rank bin** label.

annotate_clonality_prop() - label each receptor by **proportion bin** using named thresholds (matched in descending order; else "Ultra-rare").

Usage

```
annotate_clonality_rank(
  idata,
  bins = c(10, 30, 100, 300, 1000, 10000, 1e+05),
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
```

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```
annotate_clonality_prop(
  idata,
bins = c(Hyperexpanded = 0.01, Large = 0.001, Medium = 1e-04, Small = 1e-05, Rare =
     1e-06),
  autojoin = getOption("immundata.autojoin", TRUE),
  format = c("long", "wide")
)
```

Arguments

idata An immundata::ImmunData object.

bins A **named** numeric vector of thresholds (e.g., c(Hyperexpanded = 1e-2, Large

= 1e-3, ...)). Names become bin labels and must be non-empty. Internally

sorted in descending order.

autojoin Logical. If TRUE, join repertoire metadata by the schema repertoire id. Change

the default behaviour by calling options(immunarch.autojoin = FALSE).

format String. One of "long" ("long" tibble with imd_repertoire_id, facet columns,

and value; useful for visualizations) or "wide" (wide/unmelted table of features, with each row corresponding to a specific repertoire / pair of repertoires;

useful for Machine Learning).

Value

An immundata::ImmunData whose \$annotations gains:

clonal_rank_bin - integer-like label with the applied rank threshold (outside all thresholds -> NA).

An immundata::ImmunData whose \$annotations gains:

• clonal_prop_bin - label from names(bins) or "Ultra-rare".

See Also

• Per-repertoire summaries: airr_clonality

• Data container: immundata::ImmunData

```
## Not run:
idata <- get_test_idata() |> agg_repertoires("Therapy")
idata_rank <- annotate_clonality_rank(idata)
idata_prop <- annotate_clonality_prop(idata)
## End(Not run)</pre>
```

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apply_symm

Apply function to each pair of data frames from a list.

Description

Apply the given function to every pair in the given datalist. Function either symmetrical (i.e. fun(x,y) == fun(y,x)) or assymmetrical (i.e. fun(x,y) != fun(y,x)).

Usage

```
apply_symm(.datalist, .fun, ..., .diag = NA, .verbose = TRUE)
apply_asymm(.datalist, .fun, ..., .diag = NA, .verbose = TRUE)
```

Arguments

.datalist List with some data.frames.

. fun Function to apply, which return basic class value.

.. Arguments passsed to .fun.

. diag Either NA for NA or something else != NULL for .fun(x,x).

. verbose if TRUE then output a progress bar.

Value

Matrix with values M(i,j) = fun(datalist(i), datalist(j))

Examples

```
data(immdata)
apply_symm(immdata$data, function(x, y) {
  nrow(x) + nrow(y)
})
```

bcrdata

BCR dataset

Description

A dataset with BCR data for testing and examplatory purposes.

Usage

bcrdata

20 bunch_translate

Format

A list of two elements. The first element ("data") is a list of 1 element named "full_clones" that contains immune repertoire data frame. The second element ("meta") is empty metadata table.

data List of immune repertoire data frames.

meta Metadata ...

bunch_translate

Nucleotide to amino acid sequence translation

Description

Nucleotide to amino acid sequence translation

Usage

```
bunch_translate(.seq, .two.way = TRUE, .ignore.n = FALSE)
```

Arguments

. seq Vector or list of strings.

. two.way Logical. If TRUE (default) then translate from the both ends (like MIXCR).

.ignore.n Logical. If FALSE (default) then return NA for sequences that have N, else

parse triplets with N as ~

Value

Character vector of translated input sequences.

```
data(immdata)
head(bunch_translate(immdata$data[[1]]$CDR3.nt))
```

check_distribution 21

check_distribution

Check and normalise distributions

Description

Check if the given .data is a distribution and normalise it if necessary with an optional Laplace correction.

Usage

```
check_distribution(
   .data,
   .do.norm = NA,
   .laplace = 1,
   .na.val = 0,
   .warn.zero = FALSE,
   .warn.sum = TRUE
)
```

Arguments

.data	Numeric vector of values.
.do.norm	One of the three values - NA, TRUE or FALSE. If NA then checks for distrubution (sum(.data) == 1) and normalises if needed with the given laplace correction value. if TRUE then does the normalisation and laplace correction. If FALSE then doesn't do either normalisation or laplace correction.
.laplace	Value for the laplace correction.
.na.val	Replace all NAs with this value.
.warn.zero	if TRUE then the function checks if in the resulted vector (after normalisation) are any zeros, and prints a warning message if there are some.
.warn.sum	if TRUE then the function checks if the sum of resulted vector (after normalisation) is equal to one, and prints a warning message if not.

Value

Numeric vector.

Developer Examples

 $immunarch:::check_distribution(c(1, 2, 3)) \ immunarch:::check_distribution(c(1, 2, 3), TRUE) \ immunarch:::check_distribution(c(1, 2, 3), FALSE)$

22 coding

coding

Filter out coding and non-coding clonotype sequences

Description

Filter out clonotypes with non-coding, coding, in-frame or out-of-frame CDR3 sequences:

coding() - remove all non-coding sequences (i.e., remove all sequences with stop codons and frame shifts);

noncoding() - remove all coding sequences (i.e., leave sequences with stop codons and frame shifts only);

inframes() - remove all out-of-frame sequences (i.e., remove all sequences with frame shifts); outofframes() - remove all in-frame sequences (i.e., leave sequences with frame shifts only).

Note: the function will remove all clonotypes sequences with NAs in the CDR3 amino acid column.

Usage

```
coding(.data)
noncoding(.data)
inframes(.data)
outofframes(.data)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch data format

Competent users may provide advanced data representations: DBI database connections, Apache Spark DataFrame from "copy_to" or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

Value

Filtered data frame.

```
data(immdata)
immdata_cod <- coding(immdata$data)
immdata_cod1 <- coding(immdata$data[[1]])</pre>
```

dbAnnotate 23

dbAnnotate Annotate clonotypes in immune repertoires using clonotype databases (e.g., VDJDB, McPAS)	dbAnnotate	
--	------------	--

Description

[Deprecated]

Annotate clonotypes by matching them to known condition-associated immune receptors in a database. Before using this function, you must download or load the relevant database files. For more information, see the online tutorial.

Usage

```
dbAnnotate(.data, .db, .data.col, .db.col)
```

Arguments

.data	The data to process. It can be a data.frame, a data.table::data.table, or a list of these objects. Every object must have columns in the immunarch compatible format. immunarch_data_format Competent users may provide advanced data representations: DBI database con-
	nections, or a list of these objects. They are supported with the same limitations as basic objects.
	Note: each connection must represent a separate repertoire.
. db	A data frame or a data table with an immune receptor database. See dbLoad on how to load databases into R.
.data.col	Character vector. Vector of columns in the input repertoires to use for clonotype search. E.g., "CDR3.aa" or c("CDR3.aa", "V.name").
.db.col	Character vector. Vector of columns in the database to use for clonotype search. The order must match the order of ".data.col". E.g., if ".data.col" is c("CDR3.aa", "V.name"), then ".db.col" must have the exact order of columns. i.e., the first column must correspond to CDR3 amino acid sequences, and the second column must correspond to V gene segment names.

Value

Data frame with input sequences and counts or proportions for each of the input repertoire.

```
data(immdata)

#' # Example file path
file_path <- paste0(system.file(package = "immunarch"), "/extdata/db/vdjdb.example.txt")</pre>
```

24 dbLoad

```
# Load the database with human-only TRB-only receptors for all known antigens
db <- dbLoad(file_path, "vdjdb", "HomoSapiens", "TRB")
res <- dbAnnotate(immdata$data, db, "CDR3.aa", "cdr3")
res</pre>
```

dbLoad

Load clonotype databases such as VDJDB and McPAS into the R workspace

Description

[Deprecated]

The function automatically detects the database format and loads it into R. Additionally, the function provides a general query interface to databases that allows filtering by species, chain types (i.e., locus) and pathology (i.e., antigen species).

Currently we support three popular databases:

```
VDJDB - https://github.com/antigenomics/vdjdb-db
McPAS-TCR - https://friedmanlab.weizmann.ac.il/McPAS-TCR/
TBAdb from PIRD - https://db.cngb.org/pird/
```

Usage

```
dbLoad(.path, .db, .species = NA, .chain = NA, .pathology = NA)
```

Arguments

.path	Character. A path to the database file, e.g., "/Users/researcher/Downloads/McPAS-TCR.csv".
. db	Character. A database type: either "vdjdb", "vdjdb-search", "mcpas" or "tbadb". "vdjdb" for VDJDB; "vdjdb-search" for search table obtained from the web interface of VDJDB; "mcpas" for McPAS-TCR; "tbadb" for PIRD TBAdb.
.species	Character. A string or a vector of strings specifying which species need to be in the database, e.g., "HomoSapiens". Pass NA (by default) to load all available species.
.chain	Character. A string or a vector of strings specifying which chains need to be in the database, e.g., "TRB". Pass NA (by default) to load all available chains.
.pathology	Character. A string or a vector of strings specifying which disease, virus, bacteria or any condition needs to be in the database, e.g., "CMV". Pass NA (by default) to load all available conditions.

Value

Data frame with the input database records.

entropy 25

Examples

```
# Example file path
file_path <- paste0(system.file(package = "immunarch"), "/extdata/db/vdjdb.example.txt")
# Load the database with human-only TRB-only receptors for all known antigens
db <- dbLoad(file_path, "vdjdb", "HomoSapiens", "TRB")
db</pre>
```

entropy

Information measures

Description

[Deprecated]

Compute information-based estimates and distances.

Usage

```
entropy(.data, .base = 2, .norm = FALSE, .do.norm = NA, .laplace = 1e-12)
kl_div(.alpha, .beta, .base = 2, .do.norm = NA, .laplace = 1e-12)
js_div(.alpha, .beta, .base = 2, .do.norm = NA, .laplace = 1e-12, .norm.entropy = FALSE)
cross_entropy(.alpha, .beta, .base = 2, .do.norm = NA, .laplace = 1e-12, .norm.entropy = FALSE)
```

Arguments

.data	Numeric vector. Any distribution.
.base	Numeric. A base of logarithm.
.norm	Logical. If TRUE then normalises the entropy by the maximal value of the entropy.
.do.norm	If TRUE then normalises the input distributions to make them sum up to 1.
.laplace	Numeric. A value for the laplace correction.
.alpha	Numeric vector. A distribution of some random value.
.beta	Numeric vector. A distribution of some random value.
.norm.entropy	Logical. If TRUE then normalises the resulting value by the average entropy of input distributions.

Value

A numeric value.

26 fixVis

Examples

```
P <- abs(rnorm(10))
Q <- abs(rnorm(10))
entropy(P)
kl_div(P, Q)
js_div(P, Q)
cross_entropy(P, Q)</pre>
```

fixVis

Manipulate ggplot plots and create publication-ready plots

Description

[Deprecated]

The fixVis is a built-in software tool for the manipulation of plots, such as adjusting title text font and size, axes, and more. It is a powerful tool designed to produce publication-ready plots with minimal amount of coding.

Usage

```
fixVis(.plot = NA)
```

Arguments

.plot

A ggplot2 plot.

Value

No return value because it is an application.

```
if (interactive()) {
    # Compute gene usage, visualise it and tweak via fixVis
    data(immdata) # load test data
    gu <- geneUsage(immdata$data)
    p <- vis(gu)
    fixVis(p)
}</pre>
```

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geneUsage

Main function for estimation of V-gene and J-gene statistics

Description

[Deprecated]

An utility function to analyse the immune receptor gene usage (IGHD, IGHJ, IDHV, IGIJ, IGKJ, IGKV, IGLJ, IGLV, TRAJ, TRAV, TRBD, etc.) and statistics. For gene details run gene_stats().

Usage

```
geneUsage(
   .data,
   .gene = c("hs.trbv", "HomoSapiens.TRBJ", "macmul.IGHV"),
   .quant = c(NA, "count"),
   .ambig = c("inc", "exc", "maj"),
   .type = c("segment", "allele", "family"),
   .norm = FALSE
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.gene

A character vector of length one with the name of the gene you want to analyse of the specific species. If you provide a vector of different length, only the first element will be used. The string should also contain the species of interest, for example, valid ".gene" arguments are "hs.trbv", "HomoSapiens.TRBJ" or "macmul.IGHV". For details run gene_stats().

.quant

Selects the column with data to evaluate. Pass NA if you want to compute gene statistics at the clonotype level without re-weighting. Pass "count" to use the "Clones" column to weight genes by abundance of their corresponding clonotypes.

.ambig

An option to handle ambiguous gene assignments, e.g., "TRAV1,TRAV2".

- Pass "inc" to include all possible gene segments, so "TRAV1,TRAV2" is counted as a different gene segment.
- Pass "exc" to exclude all ambiguous gene assignments, so "TRAV1,TRAV2" is excluded from the resultant gene table.

28 geneUsageAnalysis

We recommend to turn it on by passing "inc" (turned on by default). You can exclude data for the cases where there is no clear match for gene, include it for every supplied gene, or pick only first from the set. Set it to "exc", "inc" or "maj", respectively.

. type Set the type of data to evaluate: "segment", "allele", or "family".

. norm If TRUE then return proportions of genes. If FALSE then return counts of genes.

Value

A data frame with rows corresponding to gene segments and columns corresponding to the input samples.

Examples

```
data(immdata)
gu <- geneUsage(immdata$data)
vis(gu)</pre>
```

geneUsageAnalysis

Post-analysis of V-gene and J-gene statistics: PCA, clustering, etc.

Description

[Deprecated]

The geneUsageAnalysis() function deploys several data analysis methods, including PCA, multidimensional scaling, Jensen-Shannon divergence, k-means, hierarchical clustering, DBscan, and different correlation coefficients.

Usage

```
geneUsageAnalysis(
    .data,
    .method = c("js+hclust", "pca+kmeans", "anova", "js+pca+kmeans"),
    .base = 2,
    .norm.entropy = FALSE,
    .cor = c("pearson", "kendall", "spearman"),
    .do.norm = TRUE,
    .laplace = 1e-12,
    .verbose = TRUE,
    .k = 2,
    .eps = 0.01,
    .perp = 1,
    .theta = 0.1
)
```

gene_segments 29

Arguments

.data	The geneUsageAnalysis() function runs on the output from geneUsage().
.method	A string that defines the type of analysis to perform. Can be "pca", "mds", "js", "kmeans", "hclust", "dbscan" or "cor" if you want to calculate correlation coefficient. In the latter case you have to provide .cor argument.
.base	A numerical value that defines the logarithm base for Jensen-Shannon divergence.
.norm.entropy	A logical value. Set TRUE to normalise your data if you haven't done it already.
.cor	A string that defines the correlation coefficient for analysis. Can be "pearson", "kendall" or "spearman".
.do.norm	A logical value. If TRUE it forces Laplace smoothing, if NA it checks if smoothing is necessary, if FALSE does nothing.
.laplace	The numeric value, which is used as a pseudocount for Laplace smoothing.
.verbose	A logical value.
.k	The number of clusters to create, passed as k to hcut or as centers to kmeans.
.eps	A numerical value, DBscan epsylon parameter, see immunr_dbscan().
.perp	A numerical value, t-SNE perplexity, see immunr_tsne().
.theta	A numerical value, t-SNE theta parameter, see immunr_tsne().

Value

Depends on the last element in the .method string. See immunr_tsne for more info.

Examples

```
data(immdata)
gu <- geneUsage(immdata$data, .norm = TRUE)
geneUsageAnalysis(gu, "js+hclust", .verbose = FALSE) %>% vis()
```

gene_segments Gene segments table

Description

Gene segments table

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gene_stats

WIP

Description

WIP

Usage

```
gene_stats()
```

Value

gene_stats returns all segment gene statistics

Examples

```
gene_stats()
get_genes("hs.trbv", "segment")
```

getKmers

Calculate the k-mer statistics of immune repertoires

Description

[Deprecated]

Usage

```
getKmers(.data, .k, .col = c("aa", "nt"), .coding = TRUE)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of

these objects.

Every object must have columns in the immunarch compatible format. immunarch data format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.k Integer. Length of k-mers.

.col Character. Which column to use, pass "aa" (by default) for CDR3 amino acid

sequence, pass "nt" for CDR3 nucleotide sequences.

.coding Logical. If TRUE (by default) then removes all non-coding sequences from

input data first.

get_immunarch_news 31

Value

Data frame with two columns (k-mers and their counts).

Examples

```
data(immdata)
kmers <- getKmers(immdata$data[[1]], 5)
kmers %>% vis()
```

get_immunarch_news

Get the Latest immunarch Update

Description

Retrieves an update message for immunarch.

Usage

```
get_immunarch_news(datepoint = "latest")
```

Arguments

datepoint

A string specifying the update date. Use "latest" for the most recent update or supply a valid date key (e.g., "Apr 2025").

Details

If datepoint is set to "latest", the function returns the most recent update. Otherwise, specify the update date key (e.g., "Apr 2025") to retrieve that particular update. If no matching update is found, a warning is issued along with available update keys.

Value

A character string with the update details or a warning if the key is not found.

See Also

```
list_immunarch_news()
```

has_class

group_from_metadata

Get a character vector of samples' groups from the input metadata file

Description

Get a character vector of samples' groups from the input metadata file

Usage

```
group_from_metadata(.by, .metadata, .sep = "; ")
```

Arguments

by Character vector. Specify a column or columns in the input metadata to group

by.

.metadata Metadata object.

. sep Character vector. Defines a separator between groups if more than one group

passed in .by.

Value

Character vector with group names.

Developer Examples

```
immunarch:::group\_from\_metadata("Status", data.frame(Status = c("A", "A", "B", "B", "C")))
```

has_class

Check for the specific class

Description

A function to check if an input object has a specific class name.

Usage

```
has_class(.data, .class)
```

Arguments

.data Any R object.

. class Character vector. Specifies a class name to check against.

Value

Logical value.

immdata 33

Developer Examples

tmp <- "abc" immunarch:::has_class(tmp, "new_class") tmp <- immunarch:::add_class(tmp, "new_class")
immunarch:::has_class(tmp, "new_class")</pre>

immdata

Single chain immune repertoire dataset

Description

A dataset with single chain TCR data for testing and examplatory purposes.

Usage

immdata

Format

A list of two elements. The first element ("data") is a list with data frames with clonotype tables. The second element ("meta") is a metadata table.

data List of immune repertoire data frames.

meta Metadata ...

Description

Get a list of package updates

Usage

immunarch_v1_updates

Format

An object of class list of length 1.

34 immunr_hclust

immunr_data_format

Specification of the data format used by immunarch dataframes

Description

- "Clones" number of barcodes (events, UMIs) or reads;
- "Proportion" proportion of barcodes (events, UMIs) or reads;
- "CDR3.nt" CDR3 nucleotide sequence;
- "CDR3.aa" CDR3 amino acid sequence;
- "V.name" names of aligned Variable gene segments;
- "D.name" names of aligned Diversity gene segments or NA;
- "J.name" names of aligned Joining gene segments;
- "V.end" last positions of aligned V gene segments (1-based);
- "D.start" positions of D'5 end of aligned D gene segments (1-based);
- "D.end" positions of D'3 end of aligned D gene segments (1-based);
- "J.start" first positions of aligned J gene segments (1-based);
- "VJ.ins" number of inserted nucleotides (N-nucleotides) at V-J junction (-1 for receptors with VDJ recombination);
- "VD.ins" number of inserted nucleotides (N-nucleotides) at V-D junction (-1 for receptors with VJ recombination);
- "DJ.ins" number of inserted nucleotides (N-nucleotides) at D-J junction (-1 for receptors with VJ recombination);
- "Sequence" full nucleotide sequence.

immunr_hclust

Clustering of objects or distance matrices

Description

[Deprecated]

Clusters the data with one of the following methods:

- immunr_hclust clusters the data using the hierarchical clustering from hcut;
- immunr_kmeans clusters the data using the K-means algorithm from kmeans;
- immunr_dbscan clusters the data using the DBSCAN algorithm from dbscan.

immunr_hclust 35

Usage

```
immunr_hclust(.data, .k = 2, .k.max = nrow(.data) - 1, .method = "complete", .dist = TRUE)
immunr_kmeans(.data, .k = 2, .k.max = as.integer(sqrt(nrow(.data))) + 1,
.method = c("silhouette", "gap_stat"))
immunr_dbscan(.data, .eps, .dist = TRUE)
```

Arguments

.data	Matrix or data frame with features, distance matrix or output from repOverlap-Analysis or geneUsageAnalysis functions.
.k	The number of clusters to create, defined as k to hout or as centers to kmeans.
.k.max	Limits the maximum number of clusters. It is passed as k.max to factoex-tra::fviz_nbclust for immunr_hclust and immunr_kmeans.
.method	Passed to factoextra::hcut or as factoextra::fviz_nbclust.
	In case of factoextra::hcut the agglomeration method is going to be used (argument hc_method).
	In case of factoextra::fviz_nbclust it is the method to be used for estimating the optimal number of clusters (argument method).
.dist	If TRUE then ".data" is expected to be a distance matrix. If FALSE then the euclidean distance is computed for the input objects.
.eps	Local radius for expanding clusters, minimal distance between points to expand clusters. Passed as eps to dbscan.

Value

immunr_hclust - list with two elements. The first element is an output from factoextra::hcut. The second element is an output from factoextra::fviz_nbclust

immunr_kmeans - list with three elements. The first element is an output from kmeans. The second element is an output from factoextra::fviz_nbclust. The third element is the input dataset .data.

immunr_dbscan - list with two elements. The first element is an output from fpc::dbscan. The second element is the input dataset .data.

```
data(immdata)
gu <- geneUsage(immdata$data, .norm = TRUE)
immunr_hclust(t(as.matrix(gu[, -1])), .dist = FALSE)
gu[is.na(gu)] <- 0
immunr_kmeans(t(as.matrix(gu[, -1])))</pre>
```

36 immunr_pca

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Dimensionality reduction

Description

[Deprecated]

Collects a set of principal variables, reducing the number of not important variables to analyse. Dimensionality reduction makes data analysis algorithms work faster and sometimes more accurate, since it also reduces noise in the data. Currently available methods are:

- immunr_pca performs PCA (Principal Component Analysis) using prcomp;
- immunr_mds performs MDS (Multi-Dimensional Scaling) using isoMODS from MASS package.
- immunr_tsne performs tSNE (t-Distributed Stochastic Neighbour Embedding) using Rtsne Rtsne package.

Usage

```
immunr_pca(.data, .scale = default_scale_fun, .raw = TRUE, .orig = FALSE, .dist = FALSE)
immunr_mds(.data, .scale = default_scale_fun, .raw = TRUE, .orig = FALSE, .dist = TRUE)
immunr_tsne(.data, .perp = 1, .dist = TRUE, ...)
```

Arguments

.data	A matrix or a data frame with features, distance matrix or output from repOver-lapAnalysis or geneUsageAnalysis functions.	
.scale	A function to apply to your data before passing it to any of dimensionality reduction algorithms. There is no scaling by default.	
.raw	If TRUE then returns the non-processed output from dimensionality reduction algorithms. Pass FALSE if you want to visualise results.	
.orig	If TRUE then returns the original result from algorithms. Pass FALSE if you want to visualise results.	
.dist	If TRUE then assumes that ".data" is a distance matrix.	
.perp	The perplexity parameter for Rtsne. Specifies the number of neighbors each data point must have in the resulting plot.	
	Other parameters passed to Rtsne.	

Value

```
immunr_pca - an output from prcomp.
immunr_mds - an output from isoMDS.
immunr_tsne - an output from Rtsne.
```

inc_overlap 37

See Also

vis.immunr_pca for visualisations.

Examples

```
data(immdata)
gu <- geneUsage(immdata$data)
gu[is.na(gu)] <- 0
gu <- t(as.matrix(gu[, -1]))
immunr_pca(gu)
immunr_mds(dist(gu))
immunr_tsne(dist(gu))</pre>
```

inc_overlap

Incremental counting of repertoire similarity

Description

[Deprecated]

For reference please look up https://www.pnas.org/content/111/16/5980 (Fig. 4).

Usage

```
inc_overlap(
   .data,
   .fun,
   .step = 1000,
   .n.steps = 10,
   .downsample = FALSE,
   .bootstrap = NA,
   .verbose.inc = TRUE,
   ...
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

. fun

Function to compute overlaps. e.g., morisita_index.

38 list_immunarch_news

.step	Either an integer or a numeric vector. In the first case, the integer defines the step of incremental overlap. In the second case, the vector encodes all repertoire sampling depths.
.n.steps	Integer. Number of steps if .step is a single integer. Skipped if ".step" is a numeric vector.
.downsample	If TRUE then performs downsampling to N clonotypes at each step instead of choosing the top N clonotypes.
.bootstrap	Set NA to turn off any bootstrapping, set a number to perform bootstrapping with this number of tries.
.verbose.inc	Logical. If TRUE then shows the output from the computation process.
•••	Other arguments passed to . fun.

Value

List with overlap matrices.

Examples

Description

Returns the list of available update keys for immunarch v1.

Usage

```
list_immunarch_news()
```

Value

A character vector containing all the date keys for the available updates.

See Also

```
get_immunarch_news()
```

matrixdiagcopy 39

matrixdiagcopy

Copy the upper matrix triangle to the lower one

Description

Copy the upper matrix triangle to the lower one

Usage

```
matrixdiagcopy(.mat)
```

Arguments

.mat

Matrix.

Value

Matrix with its upper tri part copied to the lower tri part.

Developer Examples

```
mat \leftarrow matrix(0, 3, 3) mat mat(1, 3) \leftarrow 1 mat \leftarrow immunarch:::matrixdiagcopy(mat) mat
```

public_matrix

Get a matrix with public clonotype frequencies

Description

[Deprecated]

Usage

```
public_matrix(.data)
```

Arguments

.data

Public repertoire, an output from pubRep.

Value

Matrix with per-sample clonotype counts / proportions only.

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Examples

```
data(immdata)
immdata$data <- lapply(immdata$data, head, 2000)
pr <- pubRep(immdata$data, .verbose = FALSE)
pr.mat <- public_matrix(pr)
dim(pr.mat)
head(pr.mat)</pre>
```

pubRep

Create a repertoire of public clonotypes

Description

[Deprecated]

Usage

```
pubRep(
   .data,
   .col = "aa+v",
   .quant = c("count", "prop"),
   .coding = TRUE,
   .min.samples = 1,
   .max.samples = NA,
   .verbose = TRUE
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.col

A string that specifies the column(s) to be processed. Outputs one of the following strings, separated by the plus sign: "nt" for nucleotide sequences, "aa" for amino acid sequences, "v" for V gene segments, "j" for J gene segments. E.g., pass "aa+v" to compute overlaps on CDR3 amino acid sequences paired with V gene segments, i.e., in this case a unique clonotype is a pair of CDR3 amino acid and V gene segment.

.quant

A string that specifies the column to be processed. Set "count" to see public clonotype sharing with the number of clones, set "prop" to see proportions.

.coding

Logical. If TRUE then preprocesses the data to filter out non-coding sequences.

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.min.samples	Integer. A minimal number of samples a clonotype must have to be included in the public repertoire table.
.max.samples	Integer. A maxminal number of samples a clonotype must have to be included in the public repertoire table. Set NA (by default) to have the maximal amount of samples.
.verbose	Logical. If TRUE then outputs the progress.

Value

Data table with columns for:

- Clonotypes (e.g., CDR3 sequence, or two columns for CDR3 sequence and V gene)
- Incidence of clonotypes
- Per-sample proportions or counts

Examples

```
# Subset the data to make the example faster to run
immdata$data <- lapply(immdata$data, head, 2000)
pr <- pubRep(immdata$data, .verbose = FALSE)
vis(pr, "clonotypes", 1, 2)
```

pubRepApply

Apply transformations to public repertoires

Description

[Deprecated]

Usage

```
pubRepApply(.pr1, .pr2, .fun = function(x) log10(x[1])/log10(x[2]))
```

Arguments

.pr1	First public repertoire.
.pr2	Second public repertoire.
.fun	A function to apply to pairs of frequencies of same clonotypes from "pr1" and "pr2". By default - $log(X) / log(Y)$ where X, Y - frequencies of the same clonotype, found in both public repertoires.

Value

Work in progress.

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Examples

```
data(immdata)
immdata$data <- lapply(immdata$data, head, 2000)
pr <- pubRep(immdata$data, .verbose = FALSE)
pr1 <- pubRepFilter(pr, immdata$meta, .by = c(Status = "MS"))
pr2 <- pubRepFilter(pr, immdata$meta, .by = c(Status = "C"))
prapp <- pubRepApply(pr1, pr2)
head(prapp)</pre>
```

pubRepFilter

Filter out clonotypes from public repertoires

Description

[Deprecated]

Filter our clonotypes with low incidence in a specific group.

Usage

```
pubRepFilter(.pr, .meta, .by, .min.samples = 1)
```

Arguments

.pr Public repertoires, an output from pubRep.

.meta Metadata file.

. by Named character vector. Names of the group to filter by.

.min.samples Integer. Filters out clonotypes with the number of samples below than this num-

ber.

Value

Data frame with filtered clonotypes.

```
data(immdata)
immdata$data <- lapply(immdata$data, head, 2000)
pr <- pubRep(immdata$data, .verbose = FALSE)
pr1 <- pubRepFilter(pr, immdata$meta, .by = c(Status = "MS"))
head(pr1)</pre>
```

pubRepStatistics 43

pubRepStatistics	Statistics of number of public clonotypes for each possible combinations of repertoires

Description

[Deprecated]

Usage

```
pubRepStatistics(.data, .by = NA, .meta = NA)
```

Arguments

. data Public repertoire, an output from the pubRep function.

.by Work in Progress..meta Work in Progress.

Value

Data frame with incidence statistics per sample.

Examples

```
data(immdata)
immdata$data <- lapply(immdata$data, head, 2000)
pr <- pubRep(immdata$data, .verbose = FALSE)
pubRepStatistics(pr) %>% vis()
```

repAlignLineage

Aligns all sequences incliding germline within each clonal lineage within each cluster

Description

[Deprecated]

This function aligns all sequences (incliding germline) that belong to one clonal lineage and one cluster. After clustering and building the clonal lineage and germline, the next step is to analyze the degree of mutation and maturity of each clonal lineage. This allows for finding high mature cells and cells with a large number of offspring. The phylogenetic analysis will find mutations that increase the affinity of BCR. Making alignment of the sequence is the first step towards sequence analysis including BCR.

Usage

```
repAlignLineage(.data, .min_lineage_sequences, .prepare_threads, .align_threads, .nofail)
```

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Arguments

. data The data to be processed. Can be data.frame, data.table::data.table or a list of these objects.

.min_lineage_sequences

If number of sequences in the same clonal lineage and the same cluster (not including germline) is lower than this threshold, this group of sequences will be filtered out from the dataframe; so only large enough lineages will be included.

.prepare_threads

Number of threads to prepare results table. Please note that high number can cause heavy memory usage!

.align_threads Number of threads for lineage alignment.

It must have columns in the immunarch compatible format immunarch_data_format, and also must contain 'Cluster' column, which is added by seqCluster() function, and 'Germline.sequence' column, which is added by repGermline() function.

.nofail Will return NA instead of stopping if Clustal W is not installed. Used to avoid raising errors in examples on computers where Clustal W is not installed.

Value

Dataframe or list of dataframes (if input is a list with multiple samples). The dataframe has these columns:

• Cluster: cluster name

• Germline: germline sequence

• Alignment: DNAbin object with alignment

- Sequences: nested dataframe containing all sequences for this combination of cluster and germline; it has columns
 - Sequence, CDR1.nt, CDR2.nt, CDR3.nt, FR1.nt, FR2.nt, FR3.nt, FR4.nt, V.allele, J.allele, V.aa, J.aa: all values taken from the input dataframe
 - Clone.ID: taken from the input dataframe, or created (filled with row numbers) if missing
 - Clones: taken from the input dataframe, or created (filled with '1' values) if missing

```
data(bcrdata)
bcr_data <- bcrdata$data

bcr_data %>%
   seqCluster(seqDist(bcr_data), .fixed_threshold = 3) %>%
   repGermline(.threads = 1) %>%
   repAlignLineage(.min_lineage_sequences = 2, .align_threads = 2, .nofail = TRUE)
```

repClonalFamily 45

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Builds a phylogenetic tree using the sequences of a clonal lineage

Description

[Deprecated]

This function uses the PHYLIP package to make phylogenetic analysis. For making trees it uses maximum parsimony methods.

Usage

```
repClonalFamily(.data, .vis_groups, .threads, .nofail)
```

Arguments

.vis_groups Groups for visualization, used to annotate specific clones on chart and display

them in different colors. This is a named list, where names are for the chart legend, and list items are clone IDs that belong to the groups. It's not necessary to assign groups to all clonotypes; unassigned ones will be displayed on the chart as "Clonotype" category. It's also possible to assign multiple clonotypes to the same group by providing nested lists or vectors of clone IDs instead of single clone IDs. Example: .vis_groups = list(A = 817, B = 201, C = list(303,

42))

. threads Number of threads to use.

. nofail Returns NA instead of stopping if PHYLIP is not installed. Used to avoid raising

errors in examples on computers where PHYLIP is not installed.

Value

Dataframe or list of dataframes (if input is a list with multiple samples). The dataframe has these columns:

- · Cluster: cluster name
- Germline.Input: germline sequence, like it was in the input; not aligned
- Germline.Output: germline sequence, parsed from PHYLIP dnapars function output; it contains difference of germline from the common ancestor; "." characters mean matching letters
- Common.Ancestor: common ancestor sequence, parsed from PHYLIP dnapars function output
- Trunk.Length: mean trunk length, representing the distance between the most recent common ancestor and germline sequence as a measure of the maturity of a lineage
- Tree: output tree in "phylo" format, loaded from by PHYLIP dnapars function output
- TreeStats: nested dataframe containing data about tree nodes, needed for visualization
- Sequences: nested dataframe containing all sequences for this combination of cluster and germline; it contains regions from original sequences, saved for repSomaticHypermutation() calculation, and also data needed for visualizations

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Examples

```
data(bcrdata)
bcr_data <- bcrdata$data

bcr_data %>%
   seqCluster(seqDist(bcr_data), .fixed_threshold = 3) %>%
   repGermline(.threads = 1) %>%
   repAlignLineage(.min_lineage_sequences = 2, .align_threads = 2, .nofail = TRUE) %>%
   repClonalFamily(.threads = 1, .nofail = TRUE)
```

repClonality

Clonality analysis of immune repertoires

Description

[Deprecated]

repClonality function encompasses several methods to measure clonal proportions in a given repertoire.

Usage

```
repClonality(
   .data,
   .method = c("clonal.prop", "homeo", "top", "rare"),
   .perc = 10,
   .clone.types = c(Rare = 1e-05, Small = 1e-04, Medium = 0.001, Large = 0.01,
        Hyperexpanded = 1),
   .head = c(10, 100, 1000, 3000, 10000, 30000, 1e+05),
   .bound = c(1, 3, 10, 30, 100)
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.method

A String with one of the following options: "clonal.prop", "homeo", "top" or "rare".

Set "clonal.prop" to compute clonal proportions or in other words percentage of clonotypes required to occupy specified by .perc percent of the total immune repertoire.

repClonality 47

Set "homeo" to analyse relative abundance (also known as clonal space homeostasis), which is defined as the proportion of repertoire occupied by clonal groups with specific abundances..

Set "top" to estimate relative abundance for the groups of top clonotypes in repertoire, e.g., ten most abundant clonotypes. Use ".head" to define index intervals, such as 10, 100 and so on.

Set "rare" to estimate relative abundance for the groups of rare clonotypes with low counts. Use ".bound" to define the threshold of clonotype groups.

perc A single numerical value ranging from 0 to 100.

.clone.types A named numerical vector with the threshold of the half-closed intervals that

mark off clonal groups.

. head A numerical vector with ranges of the top clonotypes.

bound A numerical vector with ranges of abundance for the rare clonotypes in the

dataset.

Details

Clonal proportion assessment is a different approach to estimate repertoire diversity. When visualised, it allows for thorough examination of immune repertoire structure and composition.

In its core this type of analysis is similar to the relative species abundance concept in ecology. Relative abundance is the percent composition of an organism of a particular kind relative to the total number of organisms in the area.

A stacked barplot of relative clonotype abundances can be therefore viewed as a non-parametric approach to comparing their underlying distributions.

Value

If input data is a single immune repertoire, then the function returns a numeric vector with clonality statistics.

Otherwise, it returns a numeric matrix with clonality statistics for all input repertoires.

See Also

```
repDiversity
```

```
# Load the data
data(immdata)
imm_pr <- repClonality(immdata$data, .method = "clonal.prop")
vis(imm_pr)
imm_top <- repClonality(immdata$data, .method = "top", .head = c(10, 100, 1000, 3000, 10000))
vis(imm_top)
imm_rare <- repClonality(immdata$data, .method = "rare")
vis(imm_rare)</pre>
```

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```
imm_hom <- repClonality(immdata$data, .method = "homeo")
vis(imm_hom)</pre>
```

repDiversity

The main function for immune repertoire diversity estimation

Description

[Deprecated]

This is a utility function to estimate the diversity of species or objects in the given distribution.

Note: functions will check if .data is a distribution of a random variable (sum == 1) or not. To force normalisation and / or to prevent this, set .do.norm to TRUE (do normalisation) or FALSE (don't do normalisation), respectively.

Usage

```
repDiversity(
  .data,
  .method = "chao1",
  .col = "aa",
  .max.q = 6,
  .min.q = 1,
  .q = 5,
  .step = NA,
  .quantile = c(0.025, 0.975),
  .extrapolation = NA,
  .perc = 50,
  .norm = TRUE,
  .verbose = TRUE,
  .do.norm = NA,
  .laplace = 0
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.method

Picks a method used for estimation out of a following list: chao1, hill, div, gini.simp, inv.simp, gini, raref, d50, dxx.

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.col A string that specifies the column(s) to be processed. Pass one of the following

strings, separated by the plus sign: "nt" for nucleotide sequences, "aa" for amino acid sequences, "v" for V gene segments, "j" for J gene segments. E.g., pass "aa+v" to compute diversity estimations on CDR3 amino acid sequences paired with V gene segments, i.e., in this case a unique clonotype is a pair of CDR3 amino acid and V gene segment. Clonal counts of equal clonotypes will be

summed up.

.max.q The max hill number to calculate (default: 5).

.min.q Function calculates several hill numbers. Set the min (default: 1).

. q q-parameter for the Diversity index.

. step Rarefaction step's size.

. quantile Numeric vector with quantiles for confidence intervals.

.extrapolation $\mbox{ An integer.}$ An upper limit for the number of clones to extrapolate to. Pass $\mbox{ 0}$

(zero) to turn extrapolation subroutines off.

. perc Set the percent to dXX index measurement.

.norm Normalises rarefaction curves..verbose If TRUE then outputs progress.

.do.norm One of the three values - NA, TRUE or FALSE. If NA then checks for distrubu-

tion (sum(.data) == 1) and normalises if needed with the given laplace correction value. if TRUE then does normalisation and laplace correction. If FALSE then

doesn't do neither normalisaton nor laplace correction.

. laplace A numeric value, which is used as a pseudocount for Laplace smoothing.

Details

- True diversity, or the effective number of types, refers to the number of equally-abundant types needed for the average proportional abundance of the types to equal that observed in the dataset of interest where all types may not be equally abundant.
- Inverse Simpson index is the effective number of types that is obtained when the weighted
 arithmetic mean is used to quantify average proportional abundance of types in the dataset of
 interest.
- The Gini coefficient measures the inequality among values of a frequency distribution (for example levels of income). A Gini coefficient of zero expresses perfect equality, where all values are the same (for example, where everyone has the same income). A Gini coefficient of one (or 100 percents) expresses maximal inequality among values (for example where only one person has all the income).
- The Gini-Simpson index is the probability of interspecific encounter, i.e., probability that two entities represent different types.
- Chao1 estimator is a nonparameteric asymptotic estimator of species richness (number of species in a population).
- Rarefaction is a technique to assess species richness from the results of sampling through extrapolation.
- Hill numbers are a mathematically unified family of diversity indices (differing among themselves only by an exponent q).

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 d50 is a recently developed immune diversity estimate. It calculates the minimum number of distinct clonotypes amounting to greater than or equal to 50 percent of a total of sequencing reads obtained following amplification and sequencing

 dXX is a similar to d50 index where XX corresponds to desirable percent of total sequencing reads.

Value

div, gini, gini.simp, inv.simp, raref return numeric vector of length 1 with value.

chao1 returns 4 values: estimated number of species, standart deviation of this number and two 95% confidence intervals for the species number.

hill returns a vector of specified length .max.q - .min.q

For most methods, if input data is a single immune repertoire, then the function returns a numeric vector with diversity statistics.

Otherwise, it returns a numeric matrix with diversity statistics for all input repertoires.

For Chao1 the function returns a matrix with diversity estimations.

For rarefaction the function returns either a matrix with diversity estimatinos on different step of the simulaiton process or a list with such matrices.

See Also

```
repOverlap, entropy, repClonality Rarefaction wiki https://en.wikipedia.org/wiki/Rarefaction_ (ecology) Hill numbers paper https://www.uvm.edu/~ngotelli/manuscriptpdfs/ChaoHill. pdf Diversity wiki https://en.wikipedia.org/wiki/Measurement_of_biodiversity
```

```
data(immdata)
# Make data smaller for testing purposes
immdata$data <- top(immdata$data, 4000)
# chao1
repDiversity(.data = immdata$data, .method = "chao1") %>% vis()
# Hill numbers
repDiversity(
   .data = immdata$data, .method = "hill", .max.q = 6,
   .min.q = 1, .do.norm = NA, .laplace = 0
) %>% vis()
# diversity
repDiversity(.data = immdata$data, .method = "div", .q = 5, .do.norm = NA, .laplace = 0) %>%
   vis()
# Gini-Simpson
repDiversity(.data = immdata$data, .method = "gini.simp", .q = 5, .do.norm = NA, .laplace = 0) %>%
   vis()
```

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```
# inverse Simpson
repDiversity(.data = immdata$data, .method = "inv.simp", .do.norm = NA, .laplace = 0) %>% vis()
# Gini coefficient
repDiversity(.data = immdata$data, .method = "gini", .do.norm = NA, .laplace = 0)
# d50
repDiversity(.data = immdata$data, .method = "d50") %>% vis()
```

repExplore

Main function for exploratory data analysis: compute the distribution of lengths, clones, etc.

Description

[Deprecated]

The repExplore function calculates the basic statistics of repertoire: the number of unique immune receptor clonotypes, their relative abundances, and sequence length distribution across the input dataset.

Usage

```
repExplore(
   .data,
   .method = c("volume", "count", "len", "clones"),
   .col = c("nt", "aa"),
   .coding = TRUE
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.method

A string that specifies the method of analysis. It can be either "volume", "count", "len" or "clones".

When .method is set to "volume" the repExplore calculates the number of unique clonotypes in the input data.

When .method is set to "count" the repExplore calculates the distribution of clonotype abundances, i.e., how frequent receptors with different abundances are.

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When .method is set to "len" the repExplore calculates the distribution of CDR3 sequence lengths.

When .method is set to "clones" the repExplore returns the number of clones (i.e., cells) per input repertoire.

.col A string that specifies the column to be processed. Pass "nt" for nucleotide

sequence or "aa" for amino acid sequence.

. coding If TRUE, then only coding sequences will be analysed.

Value

If input data is a single immune repertoire, then the function returns a numeric vector with exploratory analysis statistics.

Otherwise, it returns a numeric matrix with exploratory analysis statistics for all input repertoires.

See Also

```
vis.immunr_exp_vol
```

Examples

```
data(immdata)
# Calculate statistics and generate a visual output with vis()
repExplore(immdata$data, .method = "volume") %>% vis()
repExplore(immdata$data, .method = "count") %>% vis()
repExplore(immdata$data, .method = "len") %>% vis()
```

repFilter

Main function for data filtering

Description

[Deprecated]

Usage

```
repFilter(
   .data,
   .method = "by.clonotype",
   .query = list(CDR3.aa = exclude("partial", "out_of_frame")),
   .match = "exact"
)
```

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Arguments

.data

The data to be processed. Must be the list of 2 elements: a data table and a metadata table.

.method

Method of filtering. Implemented methods: by.meta, by.repertoire (by.rep), by.clonotype (by.cl) Default value: 'by.clonotype'.

.query

Filtering query. It's a named list of filters that will be applied to data. Possible values for names in this list are dependent on filter methods:

- by.meta: filters by metadata. Names in the named list are metadata column headers.
- by.repertoire: filters by the number of clonotypes or total number of clones in sample. Possible names in the named list are "n_clonotypes" and "n_clones".
- by.clonotype: filters by data in all samples. Names in the named list are data column headers. Elements of the named list for each of the filters are filtering options. Possible values for filtering options:
- include("STR1", "STR2", ...): keeps only rows with matching values. Available for methods: "by.meta", "by.clonotype".
- exclude("STR1", "STR2", ...): removes rows with matching values. Available for methods: "by.meta", "by.clonotype".
- lessthan(value): keeps rows/samples with numeric values less than specified. Available for methods: "by.meta", "by.repertoire", "by.clonotype".
- morethan(value): keeps rows/samples with numeric values more than specified. Available for methods: "by.meta", "by.repertoire", "by.clonotype".
- interval(from, to): keeps rows/samples with numeric values that fits in this interval. from is inclusive, to is exclusive. Available for methods: "by.meta", "by.repertoire", "by.clonotype". Default value: 'list(CDR3.aa = exclude("partial", "out_of_frame"))'.

.match

Matching method for "include" and "exclude" options in query. Possible values:

- exact: matches only the exact specified string;
- startswith: matches all strings starting with the specified substring;
- substring: matches all strings containing the specified substring. Default value: 'exact'.

repGermline

Creates germlines for clonal lineages

Description

[Deprecated]

This function creates germlines for clonal lineages. B cell clonal lineage represents a set of B cells that presumably have a common origin (arising from the same VDJ rearrangement event) and a common ancestor. Each clonal lineage has its own germline sequence that represents the ancestral sequence for each BCR in clonal lineage. In other words, germline sequence is a sequence of Bcells immediately after VDJ recombination, before B-cell maturation and hypermutation process. Germline sequence is useful for assessing the degree of mutation and maturity of the repertoire.

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Usage

```
repGermline(.data, .species, .min_nuc_outside_cdr3, .threads)
```

Arguments

.data The data to be processed. Can be data.frame, data.table::data.table or a list of

these objects.

It must have columns in the immunarch compatible format immunarch_data_format.

. species Species from which the data was acquired. Available options: "HomoSapi-

ens" (default), "MusMusculus", "BosTaurus", "CamelusDromedarius", "CanisLupusFamiliaris", "DanioRerio", "MacacaMulatta", "MusMusculusDomesticus", "MusMusculusCastaneus", "MusMusculusMolossinus", "MusMusculusMusculus", "MusSpretus", "OncorhynchusMykiss", "OrnithorhynchusAnatinus",

"OryctolagusCuniculus", "RattusNorvegicus", "SusScrofa".

.min_nuc_outside_cdr3

This parameter sets how many nucleotides should have V or J chain outside of

CDR3 to be considered good for further alignment.

. threads Number of threads to use.

Value

Data with added columns:

- Sequence (FR1+CDR1+FR2+CDR2+FR3+CDR3+FR4 in nucleotides; the column will be replaced if exists)
- V.allele, J.allele (chosen alleles of V and J genes),
- V.aa, J.aa (V and J sequences from original clonotype, outside CDR3, converted to amino acids)
- Germline.sequence (combined germline nucleotide sequence)

Examples

```
data(bcrdata)
bcrdata$data %>%
  top(5) %>%
  repGermline()
```

repLoad

Load immune repertoire files into the R workspace

Description

[Deprecated]

The repLoad function loads repertoire files into R workspace in the immunarch format where you can immediately use them for the analysis. repLoad automatically detects the right format for your files, so all you need is simply provide the path to your files.

See "Details" for more information on supported formats. See "Examples" for diving right into it.

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Usage

```
repLoad(.path, .mode = "paired", .coding = TRUE, ...)
```

Arguments

.path

A character string specifying the path to the input data. Input data can be one of the following:

- a single repertoire file. In this case repLoad returns an R data.frame;
- a vector of paths to repertoire files. Same as in the case with no metadata file presented in the next section below;
- a path to the folder with repertoire files and, if available, metadata file "metadata.txt". If the metadata file if presented, then the repLoad returns a list with two elements "data" and "meta". "data" is an another list with repertoire R data.frames. "meta" is a data frame with the metadata. If the metadata file "metadata.txt" is not presented, then the repLoad creates a dummy metadata file with sample names and returns a list with two elements "data" and "meta". If input data has multiple chains or cell types stored in the same file (for example, like in 10xGenomics repertoire files), such repertoire files will be splitted to different R data frames with only one type of chain and cell presented. The metadata file will have additional columns specifying cell and chain types for different samples.

.mode

Either "single" for single chain data or "paired" for paired chain data.

Currently "single" works for every format, and "paired" works only for 10X Genomics data.

By default, 10X Genomics data will be loaded as paired chain data, and other files will be loaded as single chain data.

.coding

A logical value. Set TRUE to get coding-only clonotypes (by defaul). Set FALSE to get all clonotypes.

. . Extra arguments for parsing functions

Details

The metadata has to be a tab delimited file with first column named "Sample". It can have any number of additional columns with arbitrary names. The first column should contain base names of files without extensions in your folder. Example:

Sample	Sex	Age	Status
immunoseq_1	M	1	C
immunoseq_2	M	2	C
immunoseq_3	FALSE	3	A

Currently, Immunarch support the following formats:

- "immunoseq" ImmunoSEQ of any version. http://www.adaptivebiotech.com/immunoseq
- "mitcr" MiTCR. https://github.com/milaboratory/mitcr
- "mixcr" MiXCR (the "all" files) of any version. https://github.com/milaboratory/mixcr

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- "migec" MiGEC. http://migec.readthedocs.io/en/latest/
- "migmap" For parsing IgBLAST results postprocessed with MigMap. https://github.com/mikessh/migmap
- "tcr" tcR, our previous package. https://imminfo.github.io/tcr/
- "vdjtools" VDJtools of any version. http://vdjtools-doc.readthedocs.io/en/latest/
- "imgt" IMGT HighV-QUEST. http://www.imgt.org/HighV-QUEST/
- "airr" adaptive immune receptor repertoire (AIRR) data format. http://docs.airr-community.org/en/latest/datarep/overv
- "10x" 10XGenomics clonotype annotations tables. https://support.10xgenomics.com/single-cell-vdj/software/pipelines/latest/output/annotation
- "archer" ArcherDX clonotype tables. https://archerdx.com/

Value

A list with two named elements:

- "data" is a list of input samples;
- "meta" is a data frame with sample metadata.

See Also

immunr_data_format for immunarch data format; repSave for file saving; repOverlap, geneUsage and repDiversity for starting with immune repertoires basic statistics.

```
# To load the data from a single file (note that you don't need to specify the data format):
file_path <- paste0(system.file(package = "immunarch"), "/extdata/io/Sample1.tsv.gz")</pre>
immdata <- repLoad(file_path)</pre>
# Suppose you have a following structure in your folder:
# >_ ls
# immunoseq1.txt
# immunoseq2.txt
# immunoseq3.txt
# metadata.txt
# To load the whole folder with every file in it type:
file_path <- paste0(system.file(package = "immunarch"), "/extdata/io/")</pre>
immdata <- repLoad(file_path)</pre>
print(names(immdata))
# We recommend creating a metadata file named "metadata.txt" in the folder.
# In that case, when you load your data you will see:
# > immdata <- repLoad("path/to/your/folder/")</pre>
# > names(immdata)
# [1] "data" "meta"
# If you do not have "metadata.txt", you will see the same output,
# but your metadata will be almost empty:
```

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```
# > immdata <- repLoad("path/to/your/folder/")
# > names(immdata)
# [1] "data" "meta"
```

rep0verlap

Main function for public clonotype statistics calculations

Description

[Deprecated]

The rep0verlap function is designed to analyse the overlap between two or more repertoires. It contains a number of methods to compare immune receptor sequences that are shared between individuals.

Usage

```
repOverlap(
   .data,
   .method = c("public", "overlap", "jaccard", "tversky", "cosine", "morisita",
        "inc+public", "inc+morisita"),
   .col = "aa",
   .a = 0.5,
   .b = 0.5,
   .verbose = TRUE,
   .step = 1000,
   .n.steps = 10,
   .downsample = FALSE,
   .bootstrap = NA,
   .verbose.inc = NA,
   .force.matrix = FALSE
)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.method

A string that specifies the method of analysis or a combination of methods. The repOverlap function supports following basic methods: "public", "overlap", "jaccard", "tversky", "cosine", "morisita". If vector of multiple methods is given for this parameter, the first method will be used.

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.col	A string that specifies the column(s) to be processed. Pass one of the following strings, separated by the plus sign: "nt" for nucleotide sequences, "aa" for amino acid sequences, "v" for V gene segments, "j" for J gene segments. E.g., pass "aa+v" to compute overlaps on CDR3 amino acid sequences paired with V gene segments, i.e., in this case a unique clonotype is a pair of CDR3 amino acid and V gene segment. Clonal counts of equal clonotypes will be summed up.
.a, .b	Alpha and beta parameters for Tversky Index. Default values give the Jaccard index measure.
.verbose	if TRUE then output the progress.
.step	Either an integer or a numeric vector.
	In the first case, the integer defines the step of incremental overlap.
	In the second case, the vector encodes all repertoire sampling depths.
.n.steps	Skipped if ".step" is a numeric vector.
.downsample	If TRUE then performs downsampling to N clonotypes at each step instead of choosing the top N clonotypes in incremental overlaps. Change nothing of you are using conventional methods.
.bootstrap	Set NA to turn off any bootstrapping, set a number to perform bootstrapping with this number of tries.
.verbose.inc	Logical. If TRUE then shows output from the computation process.
.force.matrix	Logical. If TRUE then always forces the matrix output even in case of two input repertoires.

Details

"public" and "shared" are synonyms that exist for the convenience of researchers.

The "overlap" coefficient is a similarity measure that measures the overlap between two finite sets.

The "jaccard" index is conceptually a percentage of how many objects two sets have in common out of how many objects they have total.

The "tversky" index is an asymmetric similarity measure on sets that compares a variant to a prototype.

The "cosine" index is a measure of similarity between two non-zero vectors of an inner product space that measures the cosine of the angle between them.

The "morisita" index measures how many times it is more likely to randomly select two sampled points from the same quadrat (the dataset is covered by a regular grid of changing size) then it would be in the case of a random distribution generated from a Poisson process. Duplicate objects are merged with their counts are summed up.

Value

In most cases the return value is a matrix with overlap values for each pair of repertoires.

If only two repertoires were provided, return value is single numeric value.

If one of the incremental method is chosen, return list of overlap matrix.

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See Also

```
inc_overlap, vis
```

Examples

```
data(immdata)

# Make data smaller for testing purposes
immdata$data <- top(immdata$data, 4000)

ov <- repOverlap(immdata$data, .verbose = FALSE)
vis(ov)

ov <- repOverlap(immdata$data, "jaccard", .verbose = FALSE)
vis(ov, "heatmap2")</pre>
```

repOverlapAnalysis

Post-analysis of public clonotype statistics: PCA, clustering, etc.

Description

[Deprecated]

The rep0verlapAnalysis() function contains advanced data analysis methods. You can use several clustering and dimensionality reduction techniques in order to investigate further the difference between repertoires provided.

To cluster a subset of similar data with repOverlapAnalysis() you can perform hierarchical clustering, k-means or dbscan ('hclust', 'kmeans', 'dbscan' respectively).

To reduce dimensions, for example, to select features for subsequent analysis, you can execute the multidimensional scaling or t-sne algorithms ('mds' and 'tsne' respectively).

Usage

```
repOverlapAnalysis(
   .data,
   .method = ("hclust"),
   .scale = default_scale_fun,
   .raw = TRUE,
   .perp = 1,
   .theta = 0.1,
   .eps = 0.01,
   .k = 2
)
```

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Arguments

.data	Any distance matrix between pairs of repertoires. You can also pass your output from rep0verlap().
.method	A string that defines the type of analysis to perform.
.scale	A function to scale the data before passing it to the MDS algorithm.
.raw	A logical value. Set TRUE if you want to receive raw output of clustering or dimensionality reduction function of choice. Set FALSE if you want to receive processed output that can be subjected to visualisation with vis() function.
.perp	A numerical value, t-SNE parameter, see immunr_tsne().
.theta	A numerical value, t-SNE parameter, see immunr_tsne().
.eps	A numerical value, DBscan epsylon parameter, see immunr_dbscan().
.k	The number of clusters to create, passed as k to hout or as centers to kmeans.

Value

Depends on the last element in the .method string. See immunr_tsne for more info.

Examples

```
data(immdata)
ov <- repOverlap(immdata$data)
repOverlapAnalysis(ov, "mds+hclust") %>% vis()
```

repSample

Downsampling and resampling of immune repertoires

Description

[Deprecated]

Sample (downsample) repertoires using different approches.

Usage

```
repSample(
   .data,
   .method = c("downsample", "resample", "sample"),
   .n = NA,
   .prob = TRUE
)
```

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Arguments

. data The data to be processed. Can be data.frame, data.table::data.table, or a list of

these objects.

Every object must have columns in the immunarch compatible format. immu-

 $narch_data_format$

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations

as basic objects.

Note: each connection must represent a separate repertoire.

.method Character. Name of a sampling method. See "Details" for more details. Default

value is "downsample" that downsamples the repertoires to the number of clones (i.e., reads / UMIs) that the smallest repertoire has, if user doesn't set any value

to the ".n" argument.

.n Integer. Number of clones / clonotypes / reads / UMIs to choose, depending on

the method. Set NA to sample repertoires to the size of the smallest repertoire

in the ".data".

.prob Logical. If TRUE then samples the clonotypes with probability weights equal

to their number of clones. Used only if ".method" is "sample".

Details

If .method is "downsample" then repSample chooses .n clones (not clonotypes!) from the input repertoires without any probabilistic simulation, but exactly computing each choosed clones. Such approach is is more consistent and biologically pleasant than an output from the function if .method is "resample".

If .method is "resample" then repSample uses multinomial distribution to compute the number of occurences for each cloneset. then it removes zero-number clonotypes and return the resulting data frame. Probabilities for rmultinom for each cloneset is a percentage of this cloneset in the "Proportion" column. It's a some sort of simulation of how clonotypes are chosen from the organisms.

if .method is "sample" then repSample chooses .n clonotypes (not clones!) randomly. Depending on the .prob argument, the function chooses clonotypes either according to their size (if .prob is TRUE, by default), or each clonotype has an equal chance to be choosed (if .prob is FALSE). Note that sampling is done without replacing.

Value

Subsampled immune repertoire or a list of subsampled immune repertoires.

See Also

rmultinom, clonal_proportion

```
data(immdata)
# Downsampling to 1000 clones (not clonotypes!)
tmp <- repSample(immdata$data[[1]], .n = 1000)</pre>
```

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```
sum(tmp$Clones)

# Downsampling to 1000 clonotypes
tmp <- repSample(immdata$data[[1]], "sample", .n = 1000)
nrow(tmp)

# Downsampling to the smallest repertoire by clones (not clonotypes!)
tmp <- repSample(immdata$data[c(1, 2)])
sum(tmp[[1]]$Clones)
sum(tmp[[2]]$Clones)

# Downsampling to the smallest repertoire by clonotypes
tmp <- repSample(immdata$data[c(1, 2)], "sample")
nrow(tmp[[1]]$Clones)
nrow(tmp[[2]]$Clones)</pre>
```

repSave

Save immune repertoires to the disk

Description

[Deprecated]

The repSave function is deigned to save your data to the disk in desirable format. Currently supports "immunarch" and "vdjtools" file formats.

Usage

```
repSave(.data, .path, .format = c("immunarch", "vdjtools"), .compress = TRUE)
```

Arguments

.data	An R dataframe, a list of R dataframes or a list with data and meta where first element is a list of dataframes and the latter is a dataframe with metadata.
.path	A string with the path to the output directory. It should include file name if a single dataframe is provided to .data argument.
.format	A string with desirable format specification. Current options are "immunarch" and "vdjtools".
.compress	A boolean value. Defines whether the output will be compressed or not.

Details

It is not necessary to create directories beforehand. If the provided directory does not exist it will be created automatically.

Value

No return value.

Examples

```
## Not run:
data(immdata)
# Reduce data to save time on examples
immdata$data <- map(immdata$data, ~ .x %>% head(10))
dirpath <- tempdir()
# Save the list of repertoires
repSave(immdata, dirpath)
# Load it and check if it is the same
new_immdata <- repLoad(dirpath)
# sum(immdata$data[[1]] != new_immdata$data[[1]], na.rm = TRUE)
# sum(immdata$data[2]] != new_immdata$data[2]], na.rm = TRUE)
# sum(immdata$meta != new_immdata$meta, na.rm = TRUE)</pre>
## End(Not run)
```

repSomaticHypermutation

Calculates number of mutations against the germline for each clonotype

Description

[Deprecated]

This function aligns V and J genes from the germline in each cluster with corresponding genes in each clonotype, saves the alignments for purpose of visualization, and calculates number of mutations for each clonotype.

Usage

```
repSomaticHypermutation(.data, .threads, .nofail)
```

Arguments

.data The data to be processed: an output of repClonalFamily(); variants with one

sample and list of samples are both supported.

. threads Number of threads to use.

.nofail Will return NA instead of stopping if Clustal W is not installed. Used to avoid

raising errors in examples on computers where Clustal W is not installed.

Value

Dataframe or list of dataframes (if input is a list with multiple samples). The dataframe has all the columns from repClonalFamily() output dataframe, with Sequence column unnested: the resulting dataframe has one line per clonotype. Clone.ID column contains original IDs for clonotypes, and can be used as dataframe key. New columns are added:

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- Germline.Alignment.V: contains V gene alignment of current clonotype with the germline
- Germline.Alignment.J: contains J gene alignment of current clonotype with the germline
- Substitutions: contains number of substitutions in the alignment (summary for V and J)
- Insertions: contains number of insertions in the clonotype relative to germline (summary for V and J)
- Deletions: contains number of deletions in the clonotype relative to germline (summary for V and J)
- Mutations: contains total number of mutations in the alignment (summary for V and J)

Examples

```
data(bcrdata)
bcr_data <- bcrdata$data

bcr_data %>%
  seqCluster(seqDist(bcr_data), .fixed_threshold = 3) %>%
  repGermline(.threads = 1) %>%
  repAlignLineage(.min_lineage_sequences = 2, .align_threads = 2, .nofail = TRUE) %>%
  repClonalFamily(.threads = 1, .nofail = TRUE) %>%
  repSomaticHypermutation(.threads = 1, .nofail = TRUE)
```

select_barcodes

Select specific clonotypes using barcodes from single-cell metadata

Description

[Deprecated]

Subsets the input immune repertoire by barcodes. Creates a vector of barcodes to subset or a vector cluster IDs and corresponding barcodes to get a list of immune repertoires corresponding to cluster IDs. Columns with clonotype counts and proportions are changed accordingly to the filtered barcodes.

Usage

```
select_barcodes(.data, .barcodes, .force.list = FALSE)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch data format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

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. barcodes Either a character vector with barcodes or a named character/factor vector with

barcodes as names and cluster IDs a vector elements. The output of Seurat's

Idents function works.

. force.list Logical. If TRUE then always returns a list, even if the result is one data frame.

Value

An immune repertoire (if ".barcodes" is a barcode vector) or a list of immune repertoires (if ".barcodes" is named vector or an output from Seurat::Idents()). Each element is an immune repertoire with clonotype barcodes corresponding to the input barcodes. The output list names are cluster names in the ".barcode" argument (Seurat::Idents() case only).

See Also

```
select clusters
```

Examples

```
## Not run:
data(immdata)
# Create a fake single-cell data
df <- immdata$data[[1]]
df$Barcode <- "AAAAACCCCC"
df$Barcode[51:nrow(df)] <- "GGGGGCCCCC"
barcodes <- "AAAAACCCCC"
df <- select_barcodes(df, barcodes)
nrow(df)
## End(Not run)</pre>
```

select_clusters

Split the immune repertoire data to clusters from single-cell barcodes

Description

[Deprecated]

Given the vector of barcodes from Seurat, splits the input repertoires to separate subsets following the barcodes' assigned IDs. Useful in case you want to split immune repertoires by patients or clusters.

Usage

```
select_clusters(.data, .clusters, .field = "Cluster")
```

seqCluster

Arguments

.data	List of two elements "data" and "meta", with "data" being a list of immune
	repertoires, and "meta" being a metadata table.
clusters	Factor vector with harcodes as vector names and cluster IDs as vector elements

Factor vector with barcodes as vector names and cluster IDs as vector elements.

The output of the Seurat Idents function works.

. field A string specifying the name of the field in the input metadata. New immune

repertoire subsets will have cluster IDs in this field.

Value

A list with two elements "data" and "meta" with updated immune repertoire tables and metadata.

See Also

```
select_barcodes
```

Examples

```
## Not run:
library(Seurat)
Idents(pbmc_small)
new_cluster_ids <- c("A", "B", "C")
new_cluster_ids <- levels(pbmc_small)
new_cluster_ids
pbmc_small <- RenameIdents(pbmc_small, new_cluster_ids)
## End(Not run)</pre>
```

seqCluster

Function for assigning clusters based on sequences similarity

Description

[Deprecated]

Graph clustering based on distances between sequences

Usage

```
seqCluster(.data, .dist, .perc_similarity, .nt_similarity, .fixed_threshold)
```

Arguments

. data The data which was used to caluculate .dist object. Can be data.frame, data.table::data.table,

or a list of these objects.

Every object must have columns in the immunarch compatible format immu-

narch_data_format

. dist List of distance objects produced with seqDist function.

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```
.perc_similarity
```

Numeric value between 0 and 1 specifying the maximum acceptable weight of an edge in a graph. This threshold depends on the length of sequences.

 $. \ nt_similarity \quad Numeric \ between \ 0-sequence \ length \ specifying \ the \ threshold \ of \ allowing \ a \ 1 \ in \ nucleotides \ mismatch \ in \ sequencies.$

```
.fixed_threshold
```

Numeric specifying the threshold on the maximum weight of an edge in a graph.

Value

Immdata data format object. Same as .data, but with extra 'Cluster' column with clusters assigned.

Examples

```
## Not run:
data(immdata)
# In this example, we will use only 2 samples with 500 clonotypes in each for time saving
input_data <- lapply(immdata$data[1:2], head, 500)
dist_result <- seqDist(input_data)
cluster_result <- seqCluster(input_data, dist_result, .fixed_threshold = 1)
## End(Not run)</pre>
```

seqDist

Function for computing distance for sequences

Description

[Deprecated]

Computing sequential distances between clonotypes from two repertoires:

Usage

```
seqDist(.data, .col = 'CDR3.nt', .method = 'hamming',
    .group_by = c("V.name", "J.name"), .group_by_seqLength = TRUE, .trim_genes = TRUE, ...)
```

Arguments

.data	The data to be processed. Can be data.frame, data.table::data.table, or a list of
	these objects.
	Every object must have columns in the immunarch compatible format immunarch_data_format
.col	A string that specifies the column name to be processed. The default value is 'CDR3.nt'.
.method	Character value or user-defined function.

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.group_by

Character vector of column names to group sequence by. The default value is c("V.first", "J.first"). Columns "V.first" and "J.first" containing first genes without allele suffixes are calculated automatically from "V.name" and "J.name" if absent in the data. Pass NA for no grouping options.

.group_by_seqLength

If TRUE - adds grouping by sequence length of .col argument

.trim_genes

If TRUE - use only general gene values (e.g. "IGHV1-18") of .group_by columns for clustering; if FALSE - can cause very small clusters in case of high resolution genotyping

. . . Extra arguments for user-defined function.

The default value is 'hamming' for Hamming distance which counts the number of character substitutions that turns b into a. If a and b have different number of characters the distance is Inf.

Other possible values are:

'1v' for Levenshtein distance which counts the number of deletions, insertions and substitutions necessary to turn b into a.

'lcs' for longest common substring is defined as the longest string can be obtained by pairing characters from a and b while keeping the order of characters intact.

In case of user-defined function, it should take x and y parameters as input and return dist object.

Value

Named list of list with dist objects for given repertoires for each combination of .group_by variable(s) and/or sequence length of .col.

```
## Not run:
data(immdata)
# Reducing data to save time on examples
immdata$data <- map(immdata$data, ~ .x %>% head(10))
# Computing hamming distance for the first two repertoires in `'immdata'`
seqDist(immdata$data[1:2])
# Here we define a custom distance function
# that will count the difference in number of characters in sequences.
f <- function(x, y) {</pre>
 res <- matrix(nrow = length(x), ncol = length(y))</pre>
 for (i in 1:length(x)) {
    res[i, ] <- abs(nchar(x[i]) - nchar(y))</pre>
 dimnames(res) \leftarrow list(x, y)
 return(as.dist(res))
}
seqDist(immdata$data[1:2], .method = f, .group_by_seqLength = FALSE)
```

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```
## End(Not run)
```

set_pb

Set and update progress bars

Description

Set and update progress bars

Usage

```
set_pb(.max)
add_pb(.pb, .value = 1)
```

Arguments

.max Integer. Maximal value of the progress bar.

.pb Progress bar object from set_pb.

. value Numeric. Value to add to the progress bar at each step.

Value

An updated progress bar.

Developer Examples

pb <- immunarch:::set_pb(100) immunarch:::add_pb(pb, 25) immunarch:::add_pb

spectratype

Immune repertoire spectratyping

Description

[Deprecated]

Usage

```
spectratype(.data, .quant = c("id", "count"), .col = "nt")
```

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Arguments

.data The data to be processed. Can be data.frame, data.table::data.table, or a list of

these objects.

Every object must have columns in the immunarch compatible format. immu-

narch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations

as basic objects.

Note: each connection must represent a separate repertoire.

.quant Select the column with clonal counts to evaluate. Set to "id" to count every

clonotype once. Set to "count" to take into the account number of clones per

clonotype.

.col A string that specifies the column(s) to be processed. The output is one of the

following strings, separated by the plus sign: "nt" for nucleotide sequences, "aa" for amino acid sequences, "v" for V gene segments, "j" for J gene segments. E.g., pass "aa+v" for spectratyping on CDR3 amino acid sequences paired with V gene segments, i.e., in this case a unique clonotype is a pair of CDR3 amino acid and V gene segment. Clonal counts of equal clonotypes will be summed

up.

Value

Data frame with distributions of clonotypes per CDR3 length.

Examples

```
# Load the data
data(immdata)
sp <- spectratype(immdata$data[[1]], .col = "aa+v")
vis(sp)</pre>
```

split_to_kmers

Analysis immune repertoire kmer statistics: sequence profiles, etc.

Description

[Deprecated]

Usage

```
split_to_kmers(.data, .k)
kmer_profile(.data, .method = c("freq", "prob", "wei", "self"), .remove.stop = TRUE)
```

switch_type 71

Arguments

. data Character vector or the output from getKmers.

.k Integer. Size of k-mers.

.method Character vector of length one. If "freq" then returns a position frequency matrix

(PFM) - a matrix with occurences of each amino acid in each position.

If "prob" then returns a position probability matrix (PPM) - a matrix with probabilities of occurences of each amino acid in each position. This is a traditional

representation of sequence motifs.

If "wei" then returns a position weight matrix (PWM) - a matrix with log likeli-

hoods of PPM elements.

If "self" then returns a matrix with self-information of elements in PWM.

For more information see https://en.wikipedia.org/wiki/Position_weight_matrix.

. remove.stop Logical. If TRUE (by default) remove stop codons.

Value

```
split_to_kmers - Data frame with two columns (k-mers and their counts). kmer_profile - a matrix with per-position amino acid statistics.
```

Examples

```
data(immdata)
kmers <- getKmers(immdata$data[[1]], 5)
kmer_profile(kmers) %>% vis()
```

switch_type

Return a column's name

Description

Return a column's name

Usage

```
switch_type(type)
process_col_argument(.col)
```

Arguments

type Character. Specifies the column to choose: "nt" chooses the CDR3 nucleotide

column, "aa" chooses the CDR3 amino acid column, "v" chooses the V gene

segment column, "j" chooses the J gene segment column.

. col A string that specifies the column(s) to be processed. Select one of the following

strings, separated by the plus sign: "nt" for nucleotide sequences, "aa" for amino

acid sequences, "v" for V gene segments, "j" for J gene segments.

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Value

A column's name.

Developer Examples

```
immunarch:::switch_type("nuc") immunarch:::switch_type("v")
```

top

Get the N most abundant clonotypes

Description

Get the N most abundant clonotypes

Usage

```
top(.data, .n = 10)
```

Arguments

.data

The data to be processed. Can be data.frame, data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

. n Numeric. Number of the most abundant clonotypes to return.

Value

Data frame with the .n most abundant clonotypes only.

```
data(immdata)
top(immdata$data)
top(immdata$data[[1]])
```

trackClonotypes 73

Description

[Deprecated]

Tracks the temporal dynamics of clonotypes in repertoires. For example, tracking across multiple time points after vaccination.

Note: duplicated clonotypes are merged and their counts are summed up.

Usage

```
trackClonotypes(.data, .which = list(1, 15), .col = "aa", .norm = TRUE)
```

Arguments

.data

The data to process. It can be a data.frame, a data.table::data.table, or a list of these objects.

Every object must have columns in the immunarch compatible format. immunarch_data_format

Competent users may provide advanced data representations: DBI database connections, or a list of these objects. They are supported with the same limitations as basic objects.

Note: each connection must represent a separate repertoire.

.which

An argument that regulates which clonotypes to choose for tracking. There are three options for this argument:

- 1. passes a list with two elements list(X, Y), where X is the name or the index of a target repertoire from ".data", and Y is the number of the most abundant clonotypes to take from X.
- 2. passes a character vector of sequences to take from all data frames;
- 3. passes a data frame (data table, database) with one or more columns first for sequences, and other for gene segments (if applicable).

See the "Examples" below with examples for each option.

.col

A character vector of length 1. Specifies an identifier for a column, from which the function chooses clonotype sequences. Specify "nt" for nucleotide sequences, "aa" for amino acid sequences, "aa+v" for amino acid sequences and Variable genes, "nt+j" for nucleotide sequences with Joining genes, or any combination of the above. Used only if ".which" has option 1) or option 2).

.norm

Logical. If TRUE then uses Proportion instead of the number of Clones per clonotype to store in the function output.

Value

Data frame with input sequences and counts or proportions for each of the input repertoire.

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Examples

```
# Load an example data that comes with immunarch
data(immdata)
# Make the data smaller in order to speed up the examples
immdata$data <- immdata$data[c(1, 2, 3, 7, 8, 9)]
immdata\$meta \leftarrow immdata\$meta[c(1, 2, 3, 7, 8, 9), ]
# Option 1
# Choose the first 10 amino acid clonotype sequences
# from the first repertoire to track
tc <- trackClonotypes(immdata$data, list(1, 10), .col = "aa")</pre>
# Choose the first 20 nucleotide clonotype sequences
# and their V genes from the "MS1" repertoire to track
tc <- trackClonotypes(immdata$data, list("MS1", 20), .col = "nt+v")</pre>
# Option 2
# Choose clonotypes with amino acid sequences "CASRGLITDTQYF" or "CSASRGSPNEQYF"
tc <- trackClonotypes(immdata$data, c("CASRGLITDTQYF", "CSASRGSPNEQYF"), .col = "aa")</pre>
# Choose the first 10 clonotypes from the first repertoire
# with amino acid sequences and V segments
target <- immdata$data[[1]] %>%
  select(CDR3.aa, V.name) %>%
  head(10)
tc <- trackClonotypes(immdata$data, target)</pre>
# Visualise the output regardless of the chosen option
# Therea are three way to visualise it, regulated by the .plot argument
vis(tc, .plot = "smooth")
vis(tc, .plot = "area")
vis(tc, .plot = "line")
# Visualising timepoints
# First, we create an additional column in the metadata with randomly choosen timepoints:
immdata$meta$Timepoint <- sample(1:length(immdata$data))</pre>
# Next, we create a vector with samples in the right order,
# according to the "Timepoint" column (from smallest to greatest):
sample_order <- order(immdata$meta$Timepoint)</pre>
# Sanity check: timepoints are following the right order:
immdata$meta$Timepoint[sample_order]
# Samples, sorted by the timepoints:
immdata$meta$Sample[sample_order]
# And finally, we visualise the data:
vis(tc, .order = sample_order)
```

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Description

[Deprecated]

Output from every function in immunarch can be visualised with a single function - vis. The vis automatically detects the type of the data and draws a proper visualisation. For example, output from the repOverlap function will be identified as repertoire overlap values and respective visualisation will be chosen without any additional arguments. See "Details" for the list of available visualisations.

Usage

```
vis(.data, ...)
```

Arguments

.data Pass the output from any immunarch analysis tool to vis().... Any other arguments, see the "Details" section for specific visualisation functions.

Details

List of available visualisations for different kinds of data.

Basic analysis:

- Exploratory analysis results (from repExplore) see vis.immunr_exp_vol;
- Clonality statistics (from repClonality) see vis.immunr_homeo.

Overlaps and public clonotypes:

- Overlaps (from repOverlap) using heatmaps, circos plots, polar area plots see vis.immunr_ov_matrix;
- Overlap clustering (from repOverlapAnalysis) see vis.immunr_hclust;
- Repertoire incremental overlaps (from repOverlap) see vis.immunr_inc_overlap;
- Public repertoire abundance (from pubRep) vis vis.immunr_public_repertoire.

Gene usage:

- Gene usage statistics (from geneUsage) using bar plots, box plots see vis.immunr_gene_usage;
- Gene usage distances (from geneUsageAnalysis) using heatmaps, circos plots, polar area plots
 see vis.immunr_ov_matrix;
- Gene usage clustering (from geneUsageAnalysis) see vis.immunr_hclust.

Diversity estimation:

• Diversity estimations (from repDiversity) - see vis.immunr_chao1.

BCR analysis:

• Clonal tree (from repClonalFamily) - see vis.clonal_family and vis.clonal_family_tree.

Advanced analysis:

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- Repertoire dynamics (from trackClonotypes) see vis.immunr_dynamics;
- Sequence logo plots of amino acid distributions (from kmer_profile) see vis_seqlogo;
- Kmers distributions (from getKmers) see vis.immunr_kmer_table;
- Mutation networks (from mutationNetwork) Work In Progress on vis.immunr_mutation_network;
- CDR3 amino acid properties, e.g., biophysical (from cdrProp) Work In Progress on vis.immunr_cdr_prop.

Additionaly, we provide a wrapper functions for visualisations of common data types:

- Any data frames or matrices using heatmaps see vis_heatmap and vis_heatmap2;
- Any data frames or matrices using circos plots see vis_circos.

Value

A ggplot2, pheatmap or circlize object.

See Also

fixVis for precise manipulation of plots.

Examples

```
## Not run:
# Load the test data
data(immdata)

# Compute and visualise:
ov <- repOverlap(immdata$data)
vis(ov)

gu <- geneUsage(immdata$data)
vis(gu)

dv <- repDiversity(immdata$data)
vis(dv)

## End(Not run)</pre>
```

vis.clonal_family

Visualise clonal family tree: wrapper for calling on the entire repClonalFamily output

Description

[Deprecated]

Usage

```
## S3 method for class 'clonal_family'
vis(.data, ...)
```

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Arguments

 $. \, data \qquad \qquad Clonal \, families \, from \, 1 \, or \, multiple \, samples: \, \verb"repClonalFamily" () \, output.$

... Not used here.

Value

A ggraph object.

Examples

```
## Not run:
data(bcrdata)
bcr_data <- bcrdata$data

clonal_family <- bcr_data %>%
    seqCluster(seqDist(bcr_data), .fixed_threshold = 3) %>%
    repGermline(.threads = 1) %>%
    repAlignLineage(.min_lineage_sequences = 2, .align_threads = 2, .nofail = TRUE) %>%
    repClonalFamily(.threads = 1, .nofail = TRUE) %>%
    vis()

## End(Not run)
```

vis.clonal_family_tree

Visualise clonal family tree

Description

[Deprecated]

Usage

```
## S3 method for class 'clonal_family_tree'
vis(.data, ...)
```

Arguments

.data Single clonal family tree data from 1 cluster: 1 element from TreeStats column from repClonalFamily() output.

... Not used here.

Value

A ggraph object.

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Examples

```
## Not run:
data(bcrdata)
bcr_data <- bcrdata$data

clonal_family <- bcr_data %>%
    seqCluster(seqDist(bcr_data), .fixed_threshold = 3) %>%
    repGermline(.threads = 1) %>%
    repAlignLineage(.min_lineage_sequences = 2, .align_threads = 2, .nofail = TRUE) %>%
    repClonalFamily(.threads = 1, .nofail = TRUE)

# This condition can be omitted; it prevents the example from crashing
# when ClustalW or PHYLIP are not installed
if (!("step_failure_ignored" %in% class(clonal_family))) {
    vis(clonal_family[["full_clones"]][["TreeStats"]][[2]])
}

## End(Not run)
```

vis.immunr_chao1

Visualise diversity.

Description

[Deprecated]

An utility function to visualise the output from repDiversity().

Usage

```
## S3 method for class 'immunr_chao1'
vis(
   .data,
   .by = NA,
   .meta = NA,
   .errorbars = c(0.025, 0.975),
   .errorbars.off = FALSE,
   .points = TRUE,
   .test = TRUE,
   .signif.label.size = 3.5,
   ...
)
```

Arguments

.data Output from repDiversity().

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by Pass NA if you want to plot samples without grouping.

You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".

You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should

pass NA to ".meta".

.meta A metadata object. An R dataframe with sample names and their properties,

such as age, serostatus or hla.

. errorbars A numeric vector of length two with quantiles for error bars on sectors. Disabled

if ".errorbars.off" is TRUE.

.errorbars.off If TRUE then plot CI bars for distances between each group. Disabled if no

group passed to the ".by" argument.

. points A logical value defining whether points will be visualised or not.

. test A logical vector whether statistical tests should be applied. See "Details" for

more information.

.signif.label.size

An integer value defining the size of text for p-value.

... Not used here.

Details

If data is grouped, then statistical tests for comparing means of groups will be performed, unless .test = FALSE is supplied. In case there are only two groups, the Wilcoxon rank sum test (https://en.wikipedia.org/wiki/Wilcoxon_signed-rank_test) is performed (R function wilcox.test() with an argument exact = FALSE) for testing if there is a difference in mean rank values between two groups. In case there more than two groups, the Kruskal-Wallis test (https://en.wikipedia.org/wiki/Kruskal%E2%80%93Walliway_analysis_of_variance) is performed (R function kruskal.test()), that is equivalent to ANOVA for ranks and it tests whether samples from different groups originated from the same distribution. A significant Kruskal-Wallis test indicates that at least one sample stochastically dominates one other sample. Adjusted for multiple comparisons P-values are plotted on the top of groups. P-value adjusting is done using the Holm method (https://en.wikipedia.org/wiki/Holm%E2%80%93Bonferroni_method) (also known as Holm-Bonferroni correction). You can execute the command ?p.adjust in the R console to see more.

Value

A ggplot2 object.

See Also

repDiversity vis

Examples

```
## Not run:
data(immdata)
```

```
dv <- repDiversity(immdata$data, "chao1")
vis(dv)
## End(Not run)</pre>
```

vis.immunr_clonal_prop

Visualise results of the clonality analysis

Description

[Deprecated]

An utility function to visualise the output from repClonality().

Usage

```
## S3 method for class 'immunr_clonal_prop'
vis(
   .data,
   .by = NA,
   .meta = NA,
   .errorbars = c(0.025, 0.975),
   .errorbars.off = FALSE,
   .points = TRUE,
   .test = TRUE,
   .signif.label.size = 3.5,
   ...
)
```

Arguments

.data Output from repClonality().

by Pass NA if you want to plot samples without grouping.

You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".

You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should

pass NA to ".meta".

.meta A metadata object. An R dataframe with sample names and their properties,

such as age, serostatus or hla.

. errorbars A numeric vector of length two with quantiles for error bars on sectors. Disabled

if ".errorbars.off" is TRUE.

.errorbars.off If TRUE then plot CI bars for distances between each group. Disabled if no

group passed to the ".by" argument.

```
    .points A logical value defining whether points will be visualised or not.
    .test A logical vector whether statistical tests should be applied. See "Details" for more information.
    .signif.label.size

            An integer value defining the size of text for p-value.

    ... Not used here.
```

Details

If data is grouped, then statistical tests for comparing means of groups will be performed, unless .test = FALSE is supplied. In case there are only two groups, the Wilcoxon rank sum test (https://en.wikipedia.org/wiki/Wilcoxon_signed-rank_test) is performed (R function wilcox.test() with an argument exact = FALSE) for testing if there is a difference in mean rank values between two groups. In case there more than two groups, the Kruskal-Wallis test (https://en.wikipedia.org/wiki/Kruskal%E2%80%93Walliway_analysis_of_variance) is performed (R function kruskal.test()), that is equivalent to ANOVA for ranks and it tests whether samples from different groups originated from the same distribution. A significant Kruskal-Wallis test indicates that at least one sample stochastically dominates one other sample. Adjusted for multiple comparisons P-values are plotted on the top of groups. P-value adjusting is done using the Holm method (https://en.wikipedia.org/wiki/Holm%E2%80%93Bonferroni_method) (also known as Holm-Bonferroni correction). You can execute the command ?p.adjust in the R console to see more.

Value

A ggplot2 object.

See Also

repClonality vis

Examples

```
## Not run:
data(immdata)
clp <- repClonality(immdata$data, "clonal.prop")
vis(clp)

hom <- repClonality(immdata$data, "homeo")
# Remove p values and points from the plot
vis(hom, .by = "Status", .meta = immdata$meta, .test = FALSE, .points = FALSE)
## End(Not run)</pre>
```

Description

[Deprecated]

Usage

```
## S3 method for class 'immunr_dynamics'
vis(.data, .plot = c("smooth", "area", "line"), .order = NA, .log = FALSE, ...)
```

Arguments

.data	Output from the trackClonotypes function.
.plot	Character. Either "smooth", "area" or "line". Each specifies a type of plot for visualisation of clonotype dynamics.
.order	Numeric or character vector. Specifies the order to samples, e.g., it used for ordering samples by timepoints. Either See "Examples" below for more details.
.log	Logical. If TRUE then use log-scale for the frequency axis.
	Not used here.

Value

A ggplot2 object.

Examples

```
## Not run:
# Load an example data that comes with immunarch
data(immdata)
# Make the data smaller in order to speed up the examples
immdata$data <- immdata$data[c(1, 2, 3, 7, 8, 9)]
immdata\$meta \leftarrow immdata\$meta[c(1, 2, 3, 7, 8, 9), ]
# Option 1
# Choose the first 10 amino acid clonotype sequences
# from the first repertoire to track
tc <- trackClonotypes(immdata$data, list(1, 10), .col = "aa")</pre>
# Choose the first 20 nucleotide clonotype sequences
# and their V genes from the "MS1" repertoire to track
tc <- trackClonotypes(immdata$data, list("MS1", 20), .col = "nt+v")</pre>
# Option 2
# Choose clonotypes with amino acid sequences "CASRGLITDTQYF" or "CSASRGSPNEQYF"
tc <- trackClonotypes(immdata$data, c("CASRGLITDTQYF", "CSASRGSPNEQYF"), .col = "aa")</pre>
```

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```
# Option 3
# Choose the first 10 clonotypes from the first repertoire
# with amino acid sequences and V segments
target <- immdata$data[[1]] %>%
  select(CDR3.aa, V.name) %>%
  head(10)
tc <- trackClonotypes(immdata$data, target)</pre>
# Visualise the output regardless of the chosen option
# Therea are three way to visualise it, regulated by the .plot argument
vis(tc, .plot = "smooth")
vis(tc, .plot = "area")
vis(tc, .plot = "line")
# Visualising timepoints
# First, we create an additional column in the metadata with randomly choosen timepoints:
immdata$meta$Timepoint <- sample(1:length(immdata$data))</pre>
immdata$meta
# Next, we create a vector with samples in the right order,
# according to the "Timepoint" column (from smallest to greatest):
sample_order <- order(immdata$meta$Timepoint)</pre>
# Sanity check: timepoints are following the right order:
immdata$meta$Timepoint[sample_order]
# Samples, sorted by the timepoints:
immdata$meta$Sample[sample_order]
# And finally, we visualise the data:
vis(tc, .order = sample_order)
## End(Not run)
```

vis.immunr_exp_vol

Visualise results of the exploratory analysis

Description

[Deprecated]

An utility function to visualise the output from repExplore().

Usage

```
## S3 method for class 'immunr_exp_vol'
vis(
   .data,
   .by = NA,
   .meta = NA,
   .errorbars = c(0.025, 0.975),
   .errorbars.off = FALSE,
   .points = TRUE,
   .test = TRUE,
```

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```
.signif.label.size = 3.5,
...
)
```

Arguments

.data Output from repExplore().

by Pass NA if you want to plot samples without grouping.

You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".

You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should

pass NA to ".meta".

meta A metadata object. An R dataframe with sample names and their properties,

such as age, serostatus or hla.

errorbars A numeric vector of length two with quantiles for error bars on sectors. Disabled

if ".errorbars.off" is TRUE.

.errorbars.off If TRUE then plot CI bars for distances between each group. Disabled if no

group passed to the ".by" argument.

. points A logical value defining whether points will be visualised or not.

. test A logical vector whether statistical tests should be applied. See "Details" for

more information.

.signif.label.size

An integer value defining the size of text for p-value.

... Not used here.

Details

If data is grouped, then statistical tests for comparing means of groups will be performed, unless .test = FALSE is supplied. In case there are only two groups, the Wilcoxon rank sum test (https://en.wikipedia.org/wiki/Wilcoxon_signed-rank_test) is performed (R function wilcox.test() with an argument exact = FALSE) for testing if there is a difference in mean rank values between two groups. In case there more than two groups, the Kruskal-Wallis test (https://en.wikipedia.org/wiki/Kruskal%E2%80%93Walliway_analysis_of_variance) is performed (R function kruskal.test()), that is equivalent to ANOVA for ranks and it tests whether samples from different groups originated from the same distribution. A significant Kruskal-Wallis test indicates that at least one sample stochastically dominates one other sample. Adjusted for multiple comparisons P-values are plotted on the top of groups. P-value adjusting is done using the Holm method (https://en.wikipedia.org/wiki/Holm%E2%80%93Bonferroni_method) (also known as Holm-Bonferroni correction). You can execute the command ?p.adjust in the R console to see more.

Value

A ggplot2 object.

See Also

```
repExplore vis
```

Examples

```
## Not run:
data(immdata)
repExplore(immdata$data, "volume") %>% vis()
repExplore(immdata$data, "count") %>% vis()
repExplore(immdata$data, "len") %>% vis()
repExplore(immdata$data, "clones") %>% vis()
## End(Not run)
```

vis.immunr_gene_usage Histograms and boxplots (general case / gene usage)

Description

[Deprecated]

Visualise distributions of genes using heatmaps or other plots.

Usage

```
## S3 method for class 'immunr_gene_usage'
vis(.data, .plot = c("hist", "box", "heatmap", "heatmap2", "circos"), ...)
```

Arguments

.data Output from the geneUsage function.

.plot String specifying the plot type:

- "hist" for histograms using vis hist;
- "heatmap" for heatmaps using vis_heatmap;
- "heatmap2" for heatmaps using vis_heatmap2;
- "circos" for circos plots using vis_circos.

. Other arguments passed to corresponding functions depending on the plot type:

- "hist" passes arguments to vis_hist;
- "box" passes arguments to vis_box;
- "heatmap" passes arguments to vis_heatmap;
- "heatmap2" passes arguments to vis_heatmap2 and heatmap from the "pheatmap" package;
- "circos" passes arguments to vis_circos and circlize::chordDiagram from the "circlize" package.

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Value

A ggplot2 object, pheatmap or circlize object.

See Also

```
geneUsage
```

Examples

```
## Not run:
data(immdata)
gu <- geneUsage(immdata$data[[1]])
vis(gu)
gu <- geneUsage(immdata$data)
vis(gu, .by = "Status", .meta = immdata$meta)
vis(gu, "box", .by = "Status", .meta = immdata$meta)
## End(Not run)</pre>
```

vis.immunr_hclust

Visualisation of hierarchical clustering

Description

[Deprecated]

Visualisation of the results of hierarchical clustering. For other clustering visualisations see vis.immunr_kmeans.

Usage

```
## S3 method for class 'immunr_hclust'
vis(.data, .rect = FALSE, .plot = c("clust", "best"), ...)
```

Arguments

.data	Clustering results from repOverlapAnalysis or geneUsageAnalysis.
.rect	Passed to factoextra::fviz_dend - whether to add a rectangle around groups.
.plot	A character vector of length one or two specifying which plots to visualise. If "clust" then plot only the clustering. If "best" then plot the number of optimal clusters. If both then plot both.
	Not used here.

Value

Ggplot2 objects inside the patchwork container.

vis.immunr_inc_overlap

See Also

vis, repOverlapAnalysis, geneUsageAnalysis

Examples

```
## Not run:
data(immdata)
ov <- repOverlap(immdata$data)
repOverlapAnalysis(ov, "mds+hclust") %>% vis()
## End(Not run)
```

vis.immunr_inc_overlap

Visualise incremental overlaps

Description

[Deprecated]

Usage

```
## S3 method for class 'immunr_inc_overlap'
vis(.data, .target = 1, .grid = FALSE, .ncol = 2, ...)
```

Arguments

.data	Output from the $\ensuremath{\text{repOverlap}}$ function that uses "top" methods.
.target	Index of a repertoire to plot. Omitted if .grid is TRUE.
.grid	Logical. If TRUE then plot all similarities in a grid.
.ncol	Numeric. Number of columns in the resulting grid.
	Not used here.

Value

A ggplot2 object.

See Also

repOverlap

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Examples

```
## Not run:
data(immdata)
tmp <- repOverlap(immdata$data[1:4], "inc+overlap", .verbose.inc = FALSE, .verbose = FALSE)
vis(tmp, .target = 1)
vis(tmp, .grid = TRUE)
## End(Not run)</pre>
```

vis.immunr_kmeans

Visualisation of K-means and DBSCAN clustering

Description

[Deprecated]

Visualisation of the results of K-means and DBSCAN clustering. For hierarhical clustering visualisations see vis.immunr_hclust.

Usage

```
## S3 method for class 'immunr_kmeans'
vis(
   .data,
   .point = TRUE,
   .text = TRUE,
   .ellipse = TRUE,
   .point.size = 2,
   .text.size = 10,
   .plot = c("clust", "best"),
   ...
)
```

Arguments

.data	Clustering results from repOverlapAnalysis or geneUsageAnalysis.
.point	If TRUE then plot sample points. Passed to factoextra::fviz_cluster.
.text	If TRUE then plot text labels. Passed to factoextra::fviz_cluster.
.ellipse	If TRUE then plot ellipses around all samples. Passed to "ellipse" from factoextra::fviz_cluster.
.point.size	Size of points, passed to "pointsize" from factoextra::fviz_cluster.
.text.size	Size of text labels, passed to labelsize from factoextra::fviz_cluster.
.plot	A character vector of length one or two specifying which plots to visualise. If "clust" then plot only the clustering. If "best" then plot the number of optimal clusters. If both then plot both.
	Not used here.

vis.immunr_kmer_table

Value

Ggplot2 objects inside the pathwork container.

See Also

```
vis, repOverlapAnalysis, geneUsageAnalysis
```

Examples

```
## Not run:
data(immdata)
ov <- repOverlap(immdata$data)
repOverlapAnalysis(ov, "mds+kmeans") %>% vis()
## End(Not run)
```

Description

[Deprecated]

Plot a distribution (bar plot) of the most frequent kmers in a data.

Usage

```
## S3 method for class 'immunr_kmer_table'
vis(
   .data,
   .head = 100,
   .position = c("stack", "dodge", "fill"),
   .log = FALSE,
   ...
)
```

Arguments

.data	Data frame with two columns "Kmers" and "Count" or a list with such data frames. See Examples.
. head	Number of the most frequent kmers to choose for plotting from each data frame.
.position	Character vector of length 1. Position of bars for each kmers. Value for the ggplot2 argument position.
.log	Logical. If TRUE then plot log-scaled plots.
	Not used here.

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Value

```
A ggplot2 object.
```

See Also

```
get.kmers
```

Examples

```
## Not run:
# Load necessary data and package.
data(immdata)
# Get 5-mers.
imm.km <- getKmers(immdata$data[[1]], 5)
# Plots for kmer proportions in each data frame in immdata.
p1 <- vis(imm.km, .position = "stack")
p2 <- vis(imm.km, .position = "fill")
p1 + p2
## End(Not run)</pre>
```

vis.immunr_mds

PCA / MDS / tSNE visualisation (mainly overlap / gene usage)

Description

[Deprecated]

Usage

```
## $3 method for class 'immunr_mds'
vis(
   .data,
   .by = NA,
   .meta = NA,
   .point = TRUE,
   .text = TRUE,
   .ellipse = TRUE,
   .point.size = 2,
   .text.size = 4,
   ...
)
```

Arguments

.data

Output from analysis functions such as geneUsageAnalysis or immunr_pca, immunr_mds or immunr_tsne.

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. by Pass NA if you want to plot samples without grouping.

You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".

You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should

pass NA to ".meta".

.meta A metadata object. An R dataframe with sample names and their properties,

such as age, serostatus or hla.

point Logical. If TRUE then plot points corresponding to objects.

. text Logical. If TRUE then plot sample names.

. ellipse Logical. If TRUE then plot ellipses around clusters of grouped samples.

.point.size Numeric. A size of points to plot.

.text.size Numeric. A size of sample names' labels.

... Not used here.

Details

Other visualisation methods:

```
• PCA - vis.immunr_pca
```

• MDS - vis.immunr_mds

• tSNE - vis.immunr_tsne

Value

A ggplot2 object.

Examples

```
## Not run:
data(immdata)
ov <- repOverlap(immdata$data)
repOverlapAnalysis(ov, "mds") %>% vis()
## End(Not run)
```

vis.immunr_ov_matrix Repertoire overlap and gene usage visualisations

Description

[Deprecated]

Visualises matrices with overlap values or gene usage distances among samples. For details see the links below.

Usage

```
## S3 method for class 'immunr_ov_matrix'
vis(.data, .plot = c("heatmap", "heatmap2", "circos"), ...)
```

Arguments

 $. \, data \qquad \qquad Output \ from \ repOverlap \ or \ geneUsageAnalysis.$

.plot A string specifying the plot type:

- "heatmap" for heatmaps using vis_heatmap;
- "heatmap2" for heatmaps using vis_heatmap2;
- "circos" for circos plots using vis_circos;

Other arguments are passed through to the underlying plotting function:

- "heatmap" passes arguments to vis_heatmap;
- "heatmap2" passes arguments to vis_heatmap2 and heatmap from the "pheatmap" package;
- "circos" passes arguments to vis_circos and circlize::chordDiagram from the "circlize" package;

Value

A ggplot2, pheatmap or circlize object.

Examples

```
## Not run:
data(immdata)
ov <- repOverlap(immdata$data)
vis(ov)
vis(ov, "heatmap")
vis(ov, "heatmap2")
vis(ov, "circos")
## End(Not run)</pre>
```

```
vis.immunr_public_repertoire
```

Public repertoire visualisation

Description

[Deprecated]

Usage

```
## S3 method for class 'immunr_public_repertoire'
vis(.data, .plot = c("freq", "clonotypes"), ...)
```

Arguments

. data Public repertoire, an output from pubRep.

.plot A string specifying the plot type:

- "freq" for visualisation of the distribution of occurrences of clonotypes and their frequencies using vis_public_frequencies.
- "clonotypes" for visualisation of public clonotype frequencity correlations between pairs of samples using vis_public_clonotypes

Further arguments passed vis_public_frequencies or vis_public_clonotypes, depending on the ".plot" argument.

Value

A ggplot2 object.

Examples

```
## Not run:
data(immdata)
immdata$data <- lapply(immdata$data, head, 300)
pr <- pubRep(immdata$data, .verbose = FALSE)
vis(pr, "freq")
vis(pr, "freq", .type = "none")

vis(pr, "clonotypes", 1, 2)
## End(Not run)</pre>
```

```
vis.immunr_public_statistics
```

Visualise sharing of clonotypes among samples

Description

[Deprecated]

Visualise public clonotype frequencies.

Usage

```
## S3 method for class 'immunr_public_statistics'
vis(.data, ...)
```

Arguments

```
.data Public repertoire - an output from the pubRep function.... Other arguments passed directly to UpSetR::upset.
```

Value

A ggplot2 object.

Examples

```
## Not run:
data(immdata)
immdata$data <- lapply(immdata$data, head, 2000)
pr <- pubRep(immdata$data, .verbose = FALSE)
pubRepStatistics(pr) %>% vis()
## End(Not run)
```

```
vis.step_failure_ignored
```

Handler for .nofail argument of pipeline steps that prevents examples from crashing on computers where certain dependencies are not installed

Description

[Deprecated]

Usage

```
## S3 method for class 'step_failure_ignored'
vis(.data, ...)
```

Arguments

```
... Not used here.
... Not used here.
```

Value

An empty object with "step_failure_ignored" class.

vis_bar 95

vis_bar

Bar plots

Description

[Deprecated]

Usage

```
vis_bar(
  .data,
  .by = NA,
  .meta = NA,
  .errorbars = c(0.025, 0.975),
  .errorbars.off = FALSE,
  .stack = FALSE,
  .points = TRUE,
  .test = TRUE,
  .signif.label.size = 3.5,
  .errorbar.width = 0.2,
  .defgroupby = "Sample",
  .grouping.var = "Group",
  .labs = c("X", "Y"),
  .title = "Barplot (.title argument)",
  .subtitle = "Subtitle (.subtitle argument)",
  .legend = NA,
  .leg.title = "Legend (.leg.title argument)",
  .legend.pos = "right",
  .rotate_x = 90
)
```

Arguments

.errorbars

. by Pass NA if you want to plot samples without grouping.

You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".

You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should pass NA to ".meta".

.meta A metadata object. An R dataframe with sample names and their properties, such as age, serostatus or hla.

A numeric vector of length two with quantiles for error bars on sectors. Disabled

if ".errorbars.off" is TRUE.

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.errorbars.off If TRUE then plot CI bars for distances between each group. Disabled if no

group passed to the ".by" argument.

. stack If TRUE and .errorbars.off is TRUE then plot stacked bar plots for each Group

or Sample

. points A logical value defining whether points will be visualised or not.

. test A logical vector whether statistical tests should be applied. See "Details" for

more information.

.signif.label.size

An integer value defining the size of text for p-value.

.errorbar.width

Numeric. Width for error bars.

. defgroupby A name for the column with sample names.

.grouping.var A name for the column to group by.

. labs A character vector of length two specifying names for x-axis and y-axis.

.title The text for the plot's title..subtitle The text for the plot's subtitle.

. legend If TRUE then displays a legend, otherwise removes legend from the plot.

. leg. title The text for the plots's legend. Provide NULL to remove the legend's title com-

pletely.

.legend.pos Positions of the legend: either "top", "bottom", "left" or "right".

.rotate_x How much the x tick text should be rotated? In angles.

Value

A ggplot2 object.

Examples

```
## Not run:
vis_bar(data.frame(Sample = c("A", "B", "C"), Value = c(1, 2, 3)))
## End(Not run)
```

vis_box

Flexible box-plots for visualisation of distributions

Description

[Deprecated]

Visualisation of distributions using ggplot2-based boxplots.

vis_box 97

Usage

```
vis_box(
  .data,
  .by = NA,
  .meta = NA,
  .melt = TRUE,
  .points = TRUE,
  .test = TRUE,
  .signif.label.size = 3.5,
  .defgroupby = "Sample",
  .grouping.var = "Group",
  .labs = c("X", "Y"),
  .title = "Boxplot (.title argument)",
  .subtitle = "Subtitle (.subtitle argument)",
  .legend = NA,
  .leg.title = "Legend (.leg.title argument)",
  .legend.pos = "right"
)
```

Arguments

. data Input matrix or data frame.

. by Pass NA if you want to plot samples without grouping.

You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".

You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should pass NA to ".meta".

pass NA to ".meta"

.meta A metadata object. An R dataframe with sample names and their properties,

such as age, serostatus or hla.

.melt If TRUE then apply reshape2::melt to the ".data" before plotting. In this case

".data" is supposed to be a data frame with the first character column reserved for names of genes and other numeric columns reserved to counts or frequencies of genes. Each numeric column should be associated with a specific repertoire

sample.

. points A logical value defining whether points will be visualised or not.

. test A logical vector whether statistical tests should be applied. See "Details" for

more information.

.signif.label.size

An integer value defining the size of text for p-value.

. defgroupby A name for the column with sample names.

.grouping.var A name for the column to group by.

. labs Character vector of length two with names for x-axis and y-axis, respectively.

.title The text for the title of the plot.

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. subtitle The The text for the plot's subtitle.

. legend If TRUE then displays a legend, otherwise removes legend from the plot.

.leg.title The The text for the plots's legend. Provide NULL to remove the legend's title

completely.

.legend.pos Positions of the legend: either "top", "bottom", "left" or "right".

Value

A ggplot2 object.

See Also

```
vis.immunr_gene_usage, geneUsage
```

Examples

```
## Not run:
vis_box(data.frame(Sample = sample(c("A", "B", "C"), 100, TRUE), Value = rnorm(100)), .melt = FALSE)
## End(Not run)
```

vis_circos

Visualisation of matrices using circos plots

Description

[Deprecated]

Visualise matrices with the circlize::chordDiagram function from the circlize package.

Usage

```
vis_circos(.data, .title = NULL, ...)
```

Arguments

.data Input matrix.

. title The The text for the title of the plot.

... Other arguments passed to circlize::chordDiagram from the 'circlize' package.

Value

A circlize object.

See Also

vis, repOverlap.

vis_heatmap 99

Examples

```
## Not run:
data(immdata)
ov <- repOverlap(immdata$data)
vis(ov, .plot = "circos")
## End(Not run)</pre>
```

vis_heatmap

Visualisation of matrices and data frames using ggplo2-based heatmaps

Description

[Deprecated]

Fast and easy visualisations of matrices or data frames with functions based on the ggplot2 package.

Usage

```
vis_heatmap(
   .data,
   .text = TRUE,
   .scientific = FALSE,
   .signif.digits = 2,
   .text.size = 4,
   .axis.text.size = NULL,
   .labs = c("Sample", "Sample"),
   .title = "Overlap",
   .leg.title = "Overlap values",
   .legend = TRUE,
   .na.value = NA,
   .transpose = FALSE,
   ...
)
```

Arguments

. data Input object: a matrix or a data frame.

If matrix: column names and row names (if presented) will be used as names

for labs.

If data frame: the first column will be used for row names and removed from the

data. Other columns will be used for values in the heatmap.

. text If TRUE then plots values in the heatmap cells. If FALSE does not plot values,

just plot coloured cells instead.

. scientific If TRUE then uses the scientific notation for numbers (e.g., "2.0e+2").

. signif.digits Number of significant digits to display on plot.

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. text.size Size of text in the cells of heatmap.

.axis.text.size
Size of text on the axis labels.

.labs A character vector of length two with names for x-axis and y-axis, respectively.

.title The The text for the plot's title.

.leg.title The The text for the plots's legend. Provide NULL to remove the legend's title completely.

.legend If TRUE then displays a legend, otherwise removes the legend from the plot.

.na.value Replace NA values with this value. By default they remain NA.

. transpose Logical. If TRUE then switch rows and columns.

... Other passed arguments.

Value

A ggplot2 object.

See Also

vis, repOverlap.

Examples

```
## Not run:
data(immdata)
ov <- rep0verlap(immdata$data)
vis_heatmap(ov)
gu <- geneUsage(immdata$data, "hs.trbj")
vis_heatmap(gu)
## End(Not run)</pre>
```

vis_heatmap2

Visualisation of matrices using pheatmap-based heatmaps

Description

[Deprecated]

Visualise matrices with the functions based on the pheatmap package with minimum amount of arguments.

vis_heatmap2

Usage

Arguments

.data	Input matrix. Column names and row names (if presented) will be used as names for labs.
.meta	A metadata object. An R dataframe with sample names and their properties, such as age, serostatus or hla.
.by	Set NA if you want to plot samples without grouping.
.title	The text for the plot's title (same as the "main" argument in pheatmap).
.color	A vector specifying the colors (same as the "color" argument in pheatmap). Pass NA to use the default pheatmap colors.
	Other arguments for the pheatmap function.

Value

A pheatmap object.

See Also

```
vis, repOverlap
```

Examples

```
## Not run:
data(immdata)
ov <- repOverlap(immdata$data)
vis_heatmap2(ov)
## End(Not run)</pre>
```

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vis_hist

Visualisation of distributions using histograms

Description

[Deprecated]

Visualisation of distributions using ggplot2-based histograms.

Usage

```
vis_hist(
   .data,
   .by = NA,
   .meta = NA,
   .title = "Gene usage",
   .ncol = NA,
   .points = TRUE,
   .test = TRUE,
   .coord.flip = FALSE,
   .grid = FALSE,
   .labs = c("Gene", NA),
   .melt = TRUE,
   .legend = NA,
   .add.layer = NULL,
   ...
)
```

Arguments

.data	Input matrix or data frame.
.by	Pass NA if you want to plot samples without grouping.
	You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".
	You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should pass NA to ".meta".
.meta	A metadata object. An R dataframe with sample names and their properties, such as age, serostatus or hla.
.title	The text for the title of the plot.
.ncol	A number of columns to display. Provide NA (by default) if you want the function to automatically detect the optimal number of columns.
.points	A logical value defining whether points will be visualised or not.
.test	A logical vector whether statistical tests should be applied. See "Details" for more information.

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.coord.flip If TRUE then swap x- and y-axes. If TRUE then plot separate visualisations for each sample. .grid .labs A character vector of length two with names for x-axis and y-axis, respectively. .melt If TRUE then apply reshape2::melt to the ".data" before plotting. In this case ".data" is supposed to be a data frame with the first character column reserved for names of genes and other numeric columns reserved to counts or frequencies of genes. Each numeric column should be associated with a specific repertoire sample. .legend If TRUE then plots the legend. If FALSE removes the legend from the plot. If NA automatically detects the best way to display legend. .add.layer Addditional ggplot2 layers, that added to each plot in the output plot or grid of plots. Is not used here.

Details

If data is grouped, then statistical tests for comparing means of groups will be performed, unless .test = FALSE is supplied. In case there are only two groups, the Wilcoxon rank sum test (https://en.wikipedia.org/wiki/Wilcoxon_signed-rank_test) is performed (R function wilcox.test() with an argument exact = FALSE) for testing if there is a difference in mean rank values between two groups. In case there more than two groups, the Kruskal-Wallis test (https://en.wikipedia.org/wiki/Kruskal%E2%80%93Walliway_analysis_of_variance) is performed (R function kruskal.test()), that is equivalent to ANOVA for ranks and it tests whether samples from different groups originated from the same distribution. A significant Kruskal-Wallis test indicates that at least one sample stochastically dominates one other sample. Adjusted for multiple comparisons P-values are plotted on the top of groups. P-value adjusting is done using the Holm method (https://en.wikipedia.org/wiki/Holm%E2%80%93Bonferroni_method) (also known as Holm-Bonferroni correction). You can execute the command ?p.adjust in the R console to see more.

Value

A ggplot2 object.

See Also

vis.immunr_gene_usage, geneUsage

Examples

```
## Not run:
data(immdata)
imm_gu <- geneUsage(immdata$data[[1]])
vis(imm_gu,
    .plot = "hist", .add.layer =
        theme(axis.text.x = element_text(angle = 75, vjust = 1))
)
imm_gu <- geneUsage(immdata$data[1:4])
vis(imm_gu,</pre>
```

```
.plot = "hist", .grid = TRUE, .add.layer =
    theme(axis.text.x = element_text(angle = 75, vjust = 1))

## End(Not run)
```

```
vis_immunr_kmer_profile_main

Visualise kmer profiles
```

Description

```
#' [Deprecated]
```

Usage

```
vis_immunr_kmer_profile_main(.data, .plot, ...)
```

Arguments

. data Kmer data, an output from kmer_profile.

.plot String specifying the plot type:

- "seqlogo" for traditional sequence logo plots using vis_seqlogo;
- "textlogo" for modified approach to sequence logo plots via text labels using vis_textlogo;

Other arguments passed to vis_textlogo or vis_seqlogo, depending on the ".plot" argument.

Value

A ggplot2 object.

Examples

```
## Not run:
data(immdata)
getKmers(immdata$data[[1]], 5) %>%
   kmer_profile() %>%
   vis("seqlogo")
## End(Not run)
```

vis_public_clonotypes 105

vis_public_clonotypes Visualisation of public clonotypes

Description

[Deprecated]

Visualise correlation of public clonotype frequencies in pairs of repertoires.

Usage

```
vis_public_clonotypes(
   .data,
   .x.rep = NA,
   .y.rep = NA,
   .title = NA,
   .ncol = 3,
   .point.size.modif = 1,
   .cut.axes = TRUE,
   .density = TRUE,
   .lm = TRUE,
   .radj.size = 3.5
)
```

Arguments

.data	Public repertoire data - an output from the pubRep function.	
.x.rep	Either indices of samples or character vector of sample names for the x-axis. Must be of the same length as ".y.rep".	
.y.rep	Either indices of samples or character vector of sample names for the y-axis. Must be of the same length as ".x.rep".	
.title	The text for the title of the plot.	
.ncol	An integer number of columns to print in the grid of pairs of repertoires.	
.point.size.modif		
	An integer value that is a modifier of the point size. The larger the number, the larger the points.	
.cut.axes	If TRUE then axes limits become shorter.	
.density	If TRUE then displays density plot for distributions of clonotypes for each sample. If FALSE then removes density plot from the visualisation.	
.lm	If TRUE then fit a linear model and displays an R adjusted coefficient that shows how similar samples are in terms of shared clonotypes.	
.radj.size	An integer value, that defines the size of the The text for the R adjusted coefficient.	

Value

A ggplot2 object.

See Also

pubRep, vis.immunr_public_repertoire

Examples

```
## Not run:
data(immdata)
pr <- pubRep(immdata$data, .verbose = FALSE)
vis(pr, "clonotypes", 1, 2)
## End(Not run)</pre>
```

vis_public_frequencies

Public repertoire visualisation

Description

[Deprecated]

Visualise public clonotype frequencies.

Usage

```
vis_public_frequencies(
  .data,
  .by = NA,
  .meta = NA,
  .type = c("boxplot", "none", "mean")
)
```

Arguments

.data	Public repertoire - an output from the pubRep function.
. by	Pass NA if you want to plot samples without grouping.
	You can pass a character vector with one or several column names from ".meta" to group your data before plotting. In this case you should provide ".meta".
	You can pass a character vector that exactly matches the number of samples in your data, each value should correspond to a sample's property. It will be used to group data based on the values provided. Note that in this case you should pass NA to ".meta".
.meta	A metadata object. An R dataframe with sample names and their properties, such as age, serostatus or hla.
.type	Character. Either "boxplot" for plotting distributions of frequencies, "none" for plotting everything, or "mean" for plotting average values only.

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Value

A ggplot2 object.

Examples

```
## Not run:
data(immdata)
immdata$data <- lapply(immdata$data, head, 500)</pre>
pr <- pubRep(immdata$data, .verbose = FALSE)</pre>
vis(pr, "freq", .type = "boxplot")
vis(pr, "freq", .type = "none")
vis(pr, "freq", .type = "mean")
vis(pr, "freq", .by = "Status", .meta = immdata$meta)
## End(Not run)
```

vis_textlogo

Sequence logo plots for amino acid profiles.

Description

[Deprecated]

Plot sequence logo plots for visualising of amino acid motif sequences / profiles.

vis_textlogo plots sequences in a text format - each letter has the same height. Useful when there are no big differences between occurences of amino acids in the motif.

vis_seqlogo is a traditional sequence logo plots. Useful when there are one or two amino acids with clear differences in their occurrences.

Usage

```
vis_textlogo(.data, .replace.zero.with.na = TRUE, .width = 0.1, ...)
vis_seqlogo(.data, .scheme = "chemistry", ...)
```

Arguments

.data Output from the kmer.profile function.

.replace.zero.with.na

if TRUE then replace all zeros with NAs, therefore letters with zero frequency

wont appear at the plot.

.width Width for jitter, i.e., how much points will scatter around the verical line. Pass

0 (zero) to plot points on the straight vertical line for each position.

Not used here.

.scheme Character. An argument passed to geom_logo from ggseqlogo package specify-

ing how to colour symbols.

108 vis_textlogo

Value

A ggplot2 object.

See Also

```
getKmers, kmer_profile
```

Examples

```
## Not run:
data(immdata)
kmers <- getKmers(immdata$data[[1]], 5)
ppm <- kmer_profile(kmers, "prob")
vis(ppm, .plot = "text")
vis(ppm, .plot = "seq")

d <- kmer_profile(c("CASLL", "CASSQ", "CASGL"))
vis_textlogo(d)
vis_seqlogo(d)
## End(Not run)</pre>
```

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