

Package ‘CellaRepertorium’

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Type Package

Title Data structures, clustering and testing for single
cell immune receptor repertoires (scRNAseq RepSeq/AIRR-seq)

Version 1.8.0

Description Methods to cluster and analyze high-throughput
single cell immune cell repertoires, especially from the 10X Genomics
VDJ solution. Contains an R interface to CD-HIT (Li and Godzik 2006).
Methods to visualize and analyze paired heavy-light chain data.
Tests for specific expansion, as well as omnibus oligoclonality under
hypergeometric models.

License GPL-3

Depends R (>= 4.0)

Imports dplyr,
tibble,
stringr,
Biostrings,
Rcpp,
reshape2,
methods,
rlang (>= 0.3),
purrr,
Matrix,
S4Vectors,
BiocGenerics,
tidyr,
forcats,
progress,
stats,
utils,
generics,
glue

Suggests testthat,
readr,
knitr,

```

rmarkdown,
ggplot2,
BiocStyle,
ggdendro,
broom,
lme4,
RColorBrewer,
SingleCellExperiment,
scater,
broom.mixed,
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igraph,
ggraph

```

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'RcppExports.R'

'auxiliary.R'

'ccdb_join.R'

'cdhit-methods.R'

'cluster-testing.R'

'clustering-methods.R'

'data.R'

'ggplot2-utils.R'

'pairing-methods.R'

'reexports.R'

'permutation-testing.R'

'plot_clustering.R'

'plotting.R'

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R topics documented:

.cluster_permute_test	4
canonicalize_cell	5
canonicalize_cluster	6
ccdb_ex	7
ccdb_join	8
cdhit	9
cdhit_ccdb	10
cland	11
cluster_filterset	12
cluster_germline	13
cluster_permute_test	13
cluster_plot	15
cluster_test_by	16
ContigCellDB	17
contigs_qc	19
crosstab_by_celltype	20
cross_tab_tbl	20
entropy	21
equalize_ccdb	22
fancy_name_contigs	23
filter_cdb	24
fine_clustering	25
fine_cluster_seqs	26
generate_pseudobulk	27
guess_celltype	28
hushWarning	28
ig_chain_recode	29
map_axis_labels	30
pairing_tables	31
plot_cluster_factors	33
plot_permute_test	34
purity	35
rank_prevalence_ccdb	35
rbind,ContigCellDB-method	37
reexports	37
right_join_warn	38
split_cdb	39
[[,ContigCellDB,character,missing-method	39
\$/,ContigCellDB-method	41

`.cluster_permute_test` *Cell permutation tests (internal)*

Description

Cell permutation tests (internal)

Usage

```
.cluster_permute_test(
  labels,
  covariates,
  strata,
  statistic,
  contrasts,
  n_perm,
  alternative,
  ...
)
```

Arguments

<code>labels</code>	factor of length <code>n</code>
<code>covariates</code>	<code>data.frame</code> of length <code>n</code>
<code>strata</code>	factor
<code>statistic</code>	function of label (vector) and covariate (<code>data.frame</code>). If this returns a vector, then by default each level will be compared against each other, pairwise, but see the next section.
<code>contrasts</code>	an optional list of numeric vectors. Each will be dotted with the statistic, or optionally a matrix provided in which case each row would be tested one-by-one.
<code>n_perm</code>	number of permutations to run
<code>alternative</code>	character naming the direction statistic should be fall under the alternative hypothesis
<code>...</code>	passed along to <code>statistic</code>

Value

a list containing the observed value of the statistic, the permuted values of the statistic, its expectation (under independence), a p-value, and the Monte Carlo standard error (of the expected value).

canonicalize_cell	<i>Find a canonical contig to represent a cell</i>
-------------------	--

Description

Using filtering in `contig_filter_args` and sorting in `tie_break_keys` and order find a single, canonical contig to represent each cell. Fields in `contig_fields` will be copied over to the `cell_tbl`.

Usage

```
canonicalize_cell(
  ccdb,
  contig_filter_args = TRUE,
  tie_break_keys = c("umis", "reads"),
  contig_fields = tie_break_keys,
  order = 1,
  overwrite = TRUE
)
```

Arguments

<code>ccdb</code>	<code>ContigCellDB()</code>
<code>contig_filter_args</code>	an expression passed to <code>dplyr::filter()</code> . Unlike <code>filter</code> , multiple criteria must be & together, rather than using commas to separate. These act on <code>ccdb\$contig_tbl</code>
<code>tie_break_keys</code>	(optional) character naming fields in <code>contig_tbl</code> that are used sort the contig table in descending order. Used to break ties if <code>contig_filter_args</code> does not return a unique contig for each cluster
<code>contig_fields</code>	Optional fields from <code>contig_tbl</code> that will be copied into the <code>cluster_tbl</code> from the canonical contig.
<code>order</code>	The rank order of the contig, based on <code>tie_break_keys</code> to return. If <code>tie_break_keys</code> included an ordered factor (such as <code>chain</code>) this could be used to return the second chain.
<code>overwrite</code>	logical – should non-key fields in <code>y</code> be overwritten using <code>x</code> , or should a suffix (<code>".y"</code>) be added

Value

`ContigCellDB()` with some number of clusters/contigs/cells but with "canonical" values copied into `cell_tbl`

See Also

`canonicalize_cluster()`

Examples

```
# Report beta chain with highest umi-count, breaking ties with reads
data(ccdb_ex)
beta = canonicalize_cell(ccdb_ex, chain == 'TRB',
  tie_break_keys = c('umis', 'reads'),
  contig_fields = c('umis', 'reads', 'chain', 'v_gene', 'd_gene', 'j_gene'))
head(beta$cell_tbl)

# Stable: only adds fields to `cell_tbl`
stopifnot(dplyr::all_equal(beta$cell_tbl[ccdb_ex$cell_pk],
  ccdb_ex$cell_tbl[ccdb_ex$cell_pk], ignore_row_order = TRUE))

#Report cdr3 with highest UMI count, but only when > 5 UMIs support it
umi5 = canonicalize_cell(ccdb_ex, umis > 5,
  tie_break_keys = c('umis', 'reads'), contig_fields = c('umis', 'cdr3'))
stopifnot(all(umi5$cell_tbl$umis > 5, na.rm = TRUE))
```

canonicalize_cluster *Find a canonical contig to represent a cluster*

Description

Find a canonical contig to represent a cluster

Usage

```
canonicalize_cluster(
  ccdb,
  contig_filter_args,
  tie_break_keys = character(),
  order = 1,
  representative = ccdb$cluster_pk[1],
  contig_fields = c("cdr3", "cdr3_nt", "chain", "v_gene", "d_gene", "j_gene"),
  overwrite = TRUE
)
```

Arguments

ccdb [ContigCellDB\(\)](#)

contig_filter_args an expression passed to [dplyr::filter\(\)](#). Unlike filter, multiple criteria must be & together, rather than using commas to separate. These act on `ccdb$contig_tbl`

tie_break_keys (optional) character naming fields in `contig_tbl` that are used sort the contig table in descending order. Used to break ties if `contig_filter_args` does not return a unique contig for each cluster

order	The rank order of the contig, based on tie_break_keys to return. If tie_break_keys included an ordered factor (such as chain) this could be used to return the second chain.
representative	an optional field from contig_tbl that will be made unique. Serve as a surrogate cluster_pk.
contig_fields	Optional fields from contig_tbl that will be copied into the cluster_tbl from the canonical contig.
overwrite	logical – should non-key fields in y be overwritten using x, or should a suffix (".y") be added

Value

`ContigCellDB()` with some number of clusters/contigs/cells but with "canonical" values copied into cluster_tbl

See Also

`canonicalize_cell()` `left_join_warn()`

Examples

```
library(dplyr)
data(ccdb_ex)
ccdb_ex_small = ccdb_ex
ccdb_ex_small$cell_tbl = ccdb_ex_small$cell_tbl[1:200,]
ccdb_ex_small = cdhit_ccdb(ccdb_ex_small,
  sequence_key = 'cdr3_nt', type = 'DNA', cluster_name = 'DNA97',
  identity = .965, min_length = 12, G = 1)
ccdb_ex_small = fine_clustering(ccdb_ex_small, sequence_key = 'cdr3_nt', type = 'DNA')

# Canonicalize with the medoid contig is probably what is most common
ccdb_medoid = canonicalize_cluster(ccdb_ex_small)

# But there are other possibilities.
# To pass multiple "AND" filter arguments must use &
ccdb_umi = canonicalize_cluster(ccdb_ex_small,
  contig_filter_args = chain == 'TRA' & length > 500, tie_break_keys = 'umis',
  contig_fields = c('chain', 'length'))
ccdb_umi$cluster_tbl %>% dplyr::select(chain, length) %>% summary()
```

ccdb_ex

A preconstructed ContigClusterDB from the contigs_qc data

Description

A preconstructed ContigClusterDB from the contigs_qc data

Usage

```
data(ccdb_ex)
```

Format

```
ccdb_ex = ContigCellDB_10XVDJ(contigs_qc, contig_pk = c('pop', 'sample', 'barcode', 'contig_id'), cell_pk = c('pop', 'sample', 'barcode'))
```

See Also

```
contigs\_qc
```

ccdb_join	<i>Join dataframe or SingleCellExperiment object with ContigCellDB object</i>
-----------	---

Description

Join dataframe or SingleCellExperiment object with ContigCellDB object

Usage

```
ccdb_join(template, ccdb, join_fun = dplyr::left_join, by = ccdb$cell_pk)
```

Arguments

- template data.frame or SingleCellExperiment object to be joined with ccdb.
- ccdb A ContigCellDB object.
- join_fun Function used for the join operation.
- by A character vector of variables to join by.

Value

```
ContigCellDB\(\)
```

Examples

```
data(ccdb_ex)
to_join = dplyr::bind_rows(ccdb_ex$cell_tbl[1:10,],
dplyr::tibble(barcode = c('extra1', 'extra2'), sample = LETTERS[1:2],
pop = LETTERS[1:2]))
ccdb_join(to_join, ccdb_ex)
```


Description

CDHIT is a greedy algorithm to cluster amino acid or DNA sequences based on a minimum identity. By default, in this package it is configured perform ungapped, global alignments with no clipping at start or end. The `identity` is the number of identical characters in alignment divided by the full length of the shorter sequence. Set `s < 1` to change the minimum coverage of the shorter sequence, which will allow clipping at start or end. Changing `G = 0` changes the meaning of the `identity` to be the number of identical characters in the alignment divided by the length of the alignment. In this case, you must also set the alignment coverage controls `aL`, `AL`, `aS`, `AS`.

Usage

```
cdhit(
  seqs,
  identity = NULL,
  kmerSize = NULL,
  min_length = 6,
  s = 1,
  G = 1,
  only_index = FALSE,
  showProgress = interactive(),
  ...
)
```

Arguments

<code>seqs</code>	AAseq or DNaseq
<code>identity</code>	minimum proportion identity
<code>kmerSize</code>	word size. If <code>NULL</code> , it will be chosen automatically based on the identity. You may need to lower it below 5 for AAseq with identity less than .7.
<code>min_length</code>	Minimum length for sequences to be clustered. An error if something smaller is passed.
<code>s</code>	fraction of shorter sequence covered by alignment.
<code>G</code>	1 for global alignment, 0 for local. If doubt, pick global.
<code>only_index</code>	if <code>TRUE</code> only return the integer cluster indices, otherwise return a tibble.
<code>showProgress</code>	show a status bar
<code>...</code>	other arguments that can be passed to <code>cdhit</code> , see https://github.com/weizhongli/cdhit/wiki/3.-User's-Guide#CDHIT for details. These will override any default values.

Details

CDHit is by Fu, Niu, Zhu, Wu and Li (2012). The R interface is originally by Thomas Lin Pedersen and was transcribed here because it is not exported from the package FindMyFriends, which is orphaned.

Value

vector of integer of length seqs providing the cluster ID for each sequence, or a tibble. See details.

Examples

```
fasta_path = system.file('extdata', 'demo.fasta', package='CellaRepertorium')
aaseq = Biostrings::readAAStringSet(fasta_path)
# 100% identity, global alignment
cdhit(aaseq, identity = 1, only_index = TRUE)[1:10]
# 100% identity, local alignment with no padding of endpoints
cdhit(aaseq, identity = 1, G = 0, aL = 1, aS = 1, only_index = TRUE)[1:10]
# 100% identity, local alignment with .9 padding of endpoints
cdhit(aaseq, identity = 1, G = 0, aL = .9, aS = .9, only_index = TRUE)[1:10]
# a tibble
tbl = cdhit(aaseq, identity = 1, G = 0, aL = .9, aS = .9, only_index = FALSE)
```

cdhit_ccdb

Use `cdhit()` to cluster a `ContigCellDB()`

Description

See <https://github.com/weizhongli/cdhit/wiki/3.-User's-Guide#CDHIT> for details on other potential arguments to These will override any default values.

Usage

```
cdhit_ccdb(
  cddb,
  sequence_key,
  type = c("DNA", "AA"),
  cluster_pk = "cluster_idx",
  ...
)
```

Arguments

cddb	An object of class <code>ContigCellDB()</code>
sequence_key	character naming the column in the <code>contig_tbl</code> containing the sequence to be clustered
type	one of 'DNA' or 'AA'

cluster_pk character specifying key, and name for the clustering.
... Arguments passed on to [cdhit](#)
identity minimum proportion identity
kmerSize word size. If NULL, it will be chosen automatically based on the identity. You may need to lower it below 5 for AAseq with identity less than .7.
min_length Minimum length for sequences to be clustered. An error if something smaller is passed.
s fraction of shorter sequence covered by alignment.
showProgress show a status bar
G 1 for global alignment, 0 for local. If doubt, pick global.

Value

[ContigCellDB\(\)](#)

See Also

[cdhit\(\)](#)

Examples

```
data(ccdb_ex)
res = cdhit_ccdb(ccdb_ex, 'cdr3_nt', type = 'DNA',
cluster_name = 'DNA97', identity = .965, min_length = 12, G = 1)
res$cluster_tbl
res$contig_tbl
res$cluster_pk
```

cland	<i>Cluster "And" intersection</i>
-------	-----------------------------------

Description

For each contig present in both X and Y, a new cluster is defined that combines cluster identities in both X and Y. In the resulting ContigCellDB, two contigs are in the same cluster if they are in the same cluster in X and the same cluster in Y. X and Y must have matching contig_pk. The contig_tbl has fields from X for contigs present in both X and Y. The cell_tbl from X is carried forward unchanged, while the cluster_tbl in the result contains the mapping between the ancestral clustering, and the derived.

Usage

cland(X, Y, new_pk)

Arguments

X	ContigCellDB
Y	ContigCellDB
new_pk	optional character naming the new pk.

Examples

```
data(ccdb_ex)
ccdb_germ = cluster_germline(ccdb_ex, cluster_pk = 'germline_idx')
ccdb_cdr3 = cdhit_ccdb(ccdb_ex, 'cdr3_nt', type = 'DNA',
cluster_name = 'DNA97', identity = .965, min_length = 12, G = 1)
ccdb_cdr3 = cland(ccdb_cdr3, ccdb_germ)
```

cluster_filterset	<i>A filtration of clusters</i>
-------------------	---------------------------------

Description

Return clusters that match all provided conditions

Usage

```
cluster_filterset(min_number = 0, min_freq = 0, white_list = NULL)
```

Arguments

min_number	integer At least this many cells
min_freq	numeric At least this frequency
white_list	data.frame keyed by cluster_pk that must match

Value

object representing the filtration (currently a list)

Examples

```
cluster_filterset(min_number = 1, min_freq = 0)
```

cluster_germline	<i>Cluster contigs by germline properties</i>
------------------	---

Description

Cluster contigs by germline properties

Usage

```
cluster_germline(
  ccdb,
  segment_keys = c("v_gene", "j_gene", "chain"),
  cluster_pk = "cluster_idx"
)
```

Arguments

ccdb	ContigCellDB()
segment_keys	fields in contig_tbl that identify a cluster
cluster_pk	name of cluster to be added to cluster_tbl

Value

[ContigCellDB\(\)](#)

Examples

```
data(ccdb_ex)
ccdb_ex = cluster_germline(ccdb_ex)
ccdb_ex$cluster_tbl
```

cluster_permute_test	<i>Tests for independence between labels and covariates using permutation of cells</i>
----------------------	--

Description

This tests a statistic for association between labels (for instance, cluster/clonal ID) and covariates (for instance, subject or treatment) by permuting the link between the two. Each observation represents a cell. statistic is any function of labels

Usage

```
cluster_permute_test(
  ccdb,
  cell_covariate_keys,
  cell_label_key = ccdb$cluster_pk,
  cell_stratify_keys,
  statistic,
  contrasts = NULL,
  n_perm,
  alternative = c("two.sided", "less", "greater"),
  sanity_check_strata = TRUE,
  ...
)
```

Arguments

<code>ccdb</code>	ContigCellDB
<code>cell_covariate_keys</code>	character naming fields in <code>ccdb\$cell_tbl</code>
<code>cell_label_key</code>	character naming a single field in <code>ccdb\$cell_tbl</code>
<code>cell_stratify_keys</code>	optional character naming fields in <code>ccdb\$cell_tbl</code> under which permutations of <code>cell_label_key</code> will occur. This means that the test will occur conditional on these covariates. Must be disjoint from <code>cell_covariate_keys</code> .
<code>statistic</code>	function of label (vector) and covariate (data.frame). If this returns a vector, then by default each level will be compared against each other, pairwise, but see the next section.
<code>contrasts</code>	an optional list of numeric vectors. Each will be dotted with the statistic, or optionally a matrix provided in which case each row would be tested one-by-one.
<code>n_perm</code>	number of permutations to run
<code>alternative</code>	character naming the direction statistic should be fall under the alternative hypothesis
<code>sanity_check_strata</code>	logical, should <code>cell_stratify_keys</code> be checked for sanity?
<code>...</code>	passed to <code>statistic</code>

Value

a list containing the observed value of the statistic, the permuted values of the statistic, its expectation (under independence), a p-value, and the Monte Carlo standard error (of the expected value).

See Also

[purity\(\)](#)

Examples

```
library(dplyr)
# covariate should name one or more columns in `cell_tbl`

cluster_idx = c(1, 1, 1, 2, 2, 3, 3)
subject = c('A', 'A', 'B', 'B', 'B', 'C', 'C')
contig_tbl = tibble(contig_pk = seq_along(cluster_idx), cluster_idx, subject)
ccdb_test = ContigCellDB(contig_tbl = contig_tbl, contig_pk = 'contig_pk',
  cell_pk = c('contig_pk', 'subject', 'cluster_idx'), cluster_pk = 'cluster_idx')
ccdb_test$cell_tbl

clust_test = cluster_permute_test(ccdb_test, 'subject', 'cluster_idx',
  statistic = purity, n_perm = 50)
library(ggplot2)
plot_permute_test(perm_test = clust_test)
tidy.PermuteTest(clust_test)
```

cluster_plot

*Make a plot showing properties of the clustering***Description**

The number of elements per cluster and the average distance between the medoid and other elements are plotted.

Usage

```
cluster_plot(cdb, return_plotlist = FALSE)
```

Arguments

cdb A fine_clustering ContigCellDB object

return_plotlist should a list of ggplot2 plots be returned. If FALSE, a cowplot composite is returned.

Value

a cowplot composite or a list of plots.

Examples

```
library(dplyr)
data(ccdb_ex)
ccdb_ex_small = ccdb_ex
ccdb_ex_small$cell_tbl = ccdb_ex_small$cell_tbl[1:200,]
ccdb_ex_small = cdhit_ccdb(ccdb_ex_small,
  sequence_key = 'cdr3_nt', type = 'DNA', cluster_name = 'DNA97',
  identity = .965, min_length = 12, G = 1)
```

```
ccdb_ex_small = fine_clustering(ccdb_ex_small, sequence_key = 'cdr3_nt', type = 'DNA')

# Canonicalize with the medoid contig is probably what is most common
ccdb_medoid = canonicalize_cluster(ccdb_ex_small)

# But there are other possibilities.
# To pass multiple "AND" filter arguments must use &
ccdb_umi = canonicalize_cluster(ccdb_ex_small,
  contig_filter_args = chain == 'TRA' & length > 500, tie_break_keys = 'umis',
  contig_fields = c('chain', 'length'))
ccdb_umi$cluster_tbl %>% dplyr::select(chain, length) %>% summary()
cluster_plot(ccdb_ex_small)
```

cluster_test_by	<i>Test clusters for differential usage</i>
-----------------	---

Description

Typically one will want to stratify by chain by calling `cluster_test_by`, as this will calculate the number of cell "trials" separately depending on the chain recovered.

Usage

```
cluster_test_by(ccdb, fields = "chain", tbl = "cluster_tbl", ...)

cluster_logistic_test(
  formula,
  ccdb,
  filterset = cluster_filterset(),
  contig_filter_args = TRUE,
  tie_break_keys = c("umis", "reads"),
  add_cluster_tbl = FALSE,
  keep_fit = FALSE,
  fitter = glm_glmr,
  silent = FALSE
)
```

Arguments

<code>ccdb</code>	ContigCellDB()
<code>fields</code>	character naming fields in <code>tbl</code>
<code>tbl</code>	one of <code>contig_tbl</code> , <code>cell_tbl</code> or <code>cluster_tbl</code>
<code>...</code>	passed to <code>cluster_logistic_test</code>
<code>formula</code>	the right-hand side of a glmer or glm-style formula.
<code>filterset</code>	a call to cluster_filterset() that will be used to subset clusters.

<code>contig_filter_args</code>	an expression passed to <code>dplyr::filter()</code> . Unlike <code>filter</code> , multiple criteria must be & together, rather than using commas to separate. These act on <code>ccdb\$contig_tbl</code>
<code>tie_break_keys</code>	(optional) character naming fields in <code>contig_tbl</code> that are used sort the contig table in descending order. Used to break ties if <code>contig_filter_args</code> does not return a unique contig for each cluster
<code>add_cluster_tbl</code>	logical should the output be joined to the <code>cluster_tbl</code> ?
<code>keep_fit</code>	logical as to whether the fit objects should be returned as a list column
<code>fitter</code>	a function taking arguments <code>formula</code> , <code>data</code> , <code>is_mixed</code> and <code>keep_fit</code> that is run on each cluster. Should return a <code>tibble</code> or <code>data.frame</code>
<code>silent</code>	logical. Should warnings from fitting functions should be suppressed?

Value

table with one row per cluster/term.

Functions

- `cluster_test_by`: split `ccdb` and conduct tests within strata

Examples

```
library(dplyr)
data(ccdb_ex)
ccdb_ex = cluster_germline(ccdb_ex)
trav1 = filter(ccdb_ex$cluster_tbl, v_gene == 'TRAV1')
cluster_logistic_test(~pop + (1|sample), ccdb_ex,
  filterset = cluster_filterset(white_list= trav1))
# Fixed effect analysis of each cluster, by chain
prev4 = ccdb_ex$contig_tbl %>% group_by(cluster_idx) %>%
  summarize(n()) %>% filter(`n()`>= 4)
cluster_test_by(ccdb = ccdb_ex, fields = 'chain',
  tbl = 'cluster_tbl', formula = ~ pop, filterset = cluster_filterset(white_list= prev4))
```

Description

Construct a ContigCellDB

Usage

```
ContigCellDB(
  contig_tbl,
  contig_pk,
  cell_tbl,
  cell_pk,
  cluster_tbl,
  cluster_pk = character(),
  equalize = TRUE
)

ContigCellDB_10XVDJ(
  contig_tbl,
  contig_pk = c("barcode", "contig_id"),
  cell_pk = "barcode",
  ...
)
```

Arguments

<code>contig_tbl</code>	a data frame of contigs, and additional fields describing their properties
<code>contig_pk</code>	character vector naming fields in <code>contig_tbl</code> that uniquely identify a row/contig
<code>cell_tbl</code>	a data frame of cell barcodes, and (optional) additional fields describing their properties
<code>cell_pk</code>	character vector naming fields in <code>cell_tbl</code> that uniquely identify a cell barcode
<code>cluster_tbl</code>	A data frame that provide cluster assignments for each contig
<code>cluster_pk</code>	If <code>cluster_tbl</code> was provided, a character vector naming fields in <code>cluster_tbl</code> that uniquely identify a cluster
<code>equalize</code>	logical. Should the contig, cells and clusters be equalized by taking the intersection of their common keys?
<code>...</code>	passed to ContigCellDB()

Value

ContigCellDB

Functions

- `ContigCellDB_10XVDJ`: provide defaults that correspond to identifiers in 10X VDJ data

Accessors/mutators

See [\\$,ContigCellDB-method](#) for more on how to access and mutate slots. See [mutate_cdb\(\)](#) and [filter_cdb\(\)](#) for endomorphic filtering/mutation methods See [split_cdb\(\)](#) to split into a list, and [rbind.ContigCellDB\(\)](#) for the inverse operation.

See Also

[\\$,ContigCellDB-method](#)

Examples

```
data(contigs_qc)
contigs_qc

cdb = ContigCellDB(contigs_qc, contig_pk = c('barcode', 'pop', 'sample', 'contig_id'),
  cell_pk = c('barcode', 'pop', 'sample'))
cdb

# everything that was in contigs_qc
cdb$contig_tbl

# Only the cell_pk are included by default (until clustering/canonicalization)
cdb$cell_tbl

# Empty, since no cluster_pk was specified
cdb$cluster_tbl

# Keys
cdb$contig_pk
cdb$cell_pk
cdb$cluster_pk
```

contigs_qc

Filtered and annotated contigs of TCR from mice

Description

Data for c57bl6 and balbc mice TCR were downloaded from 10x Genomics website as shown in `system.file('script/10XMouseTCR_v3_chem.R', package = 'CellaRepertorium')`. Additional processing of these data is done in the vignette `mouse_tcell_qc` and are serialized to serve as an examples for other vignettes and documentation.

Usage

```
data(contigs_qc)
```

Format

A data frame of 3399 contigs and 22 fields, all except 4 are originally defined in <https://support.10xgenomics.com/single-cell-vdj/software/pipelines/latest/output/annotation#contig>. The following fields were defined ex post facto.

1. anno_file: Path to original csv file
2. pop: Mouse strain.

- 3. sample: An artificial "replicate" from the original data defined by subsampling with replacement
- 4. celltype: The putative cell type of the contig.

crosstab_by_celltype	<i>Count contig UMIs by celltype</i>
----------------------	--------------------------------------

Description

Count contig UMIs by celltype

Usage

crosstab_by_celltype(ccdb)

Arguments

ccdb A ContigCellDB object

Value

a table, keyed by cell_pk counting UMIs per celltype

See Also

[guess_celltype\(\)](#)

Examples

```
data(ccdb_ex)
nrow(ccdb_ex$cell_tbl)
total_umi = crosstab_by_celltype(ccdb_ex)
nrow(total_umi)
```

cross_tab_tbl	<i>Generate a 2d cross tab using arbitrary numbers of columns as factors</i>
---------------	--

Description

As many rows as unique combs of x_fields As many columns as unique combs of y_fields No NA.

Usage

cross_tab_tbl(tbl, x_fields, y_fields)

Arguments

<code>tbl</code>	<code>data.frame</code>
<code>x_fields</code>	character fields in <code>tbl</code>
<code>y_fields</code>	character fields in <code>tbl</code>

Value

tibble

Examples

```
cross_tab_tbl(mtcars, c('cyl', 'gear'), 'carb')
```

entropy

Calculate the entropy of a vector

Description

Calculate the entropy of a vector

Usage

```
entropy(v, pseudo_count = length(v)/1000, na.action = na.fail)
```

```
np(v, p = 0.05, pseudo_count = p/5, na.action = na.fail)
```

```
modal_category(v, na.action = na.fail)
```

Arguments

<code>v</code>	categorical vector
<code>pseudo_count</code>	number of pseudo counts to add on, to stabilize empty categories
<code>na.action</code>	how to handle NA values
<code>p</code>	proportion threshold

Value

the sample entropy

Functions

- `np`: The number of categories exceeding `p` proportion of the total
- `modal_category`: The modal category of `v`. Ties are broken by lexicographic order of the factor levels.

Examples

```

v2 = gl(2, 4)
v4 = gl(4, 4)
stopifnot(entropy(v2) < entropy(v4))
v_empty = v2[1:4] #empty level 2
stopifnot(is.finite(entropy(v_empty))) # pseudo_count

np(v4, p = .2, pseudo_count = 0)
np(v4, p = .25, pseudo_count = 0)
np(v4, p = .25, pseudo_count = .0001)

modal_category(v4)
modal_category(v4[-1])

```

equalize_ccdb

Take the intersection of keys in tables in x

Description

The cells in `cell_tbl`, and clusters in `cluster_tbl` can potentially be a superset of the `contig_tbl`.

Usage

```
equalize_ccdb(x, cell = TRUE, contig = TRUE, cluster = TRUE, sort = FALSE)
```

Arguments

<code>x</code>	<code>ContigCellDB()</code>
<code>cell</code>	logical equalize cells
<code>contig</code>	logical equalize contigs
<code>cluster</code>	logical equalize clusters
<code>sort</code>	logical should equalized fields also be <code>order()</code> ed by their primary keys?

Details

- `equalize_ccdb(x, cell = TRUE)` trims cells that aren't in `contig_tbl` or `cluster_tbl`.
- `equalize_ccdb(x, cluster = TRUE)` trims clusters that aren't in `contig_tbl`.
- `equalize_ccdb(x, contig = TRUE)` trims contigs that aren't `cell_tbl` or `cluster_tbl`.

Value

`ContigCellDB()`

Default equalization

Modification to `contig_tbl` (with `$`) always equalizes contigs and clusters. Modification to `cell_tbl` equalizes only contigs. Modification to `cluster_tbl` equalizes contigs and clusters.

Examples

```
library(dplyr)
tbl = tibble(clust_idx = gl(3, 2), cell_idx = rep(1:3, times = 2), contig_idx = 1:6)
ccdb = ContigCellDB(tbl, contig_pk = c('cell_idx', 'contig_idx'),
  cell_pk = 'cell_idx', cluster_pk = 'clust_idx')
# 3 cells
ccdb
ccdb$cell_tbl = bind_rows(ccdb$cell_tbl, tibble(cell_idx = 0))
# 4 cells now
ccdb
# 3 cells again
equalize_ccdb(ccdb)
# remove all contigs from cell 1, and one contig from cell 2
ccdb$contig_tbl = ccdb$contig_tbl[-c(1, 2, 4),]
# no changes to cell_tbl yet
ccdb
# trim cell_tbl to 2 cells, keep all clusters
equalize_ccdb(ccdb, cluster = FALSE)
# trim both cells and clusters
equalize_ccdb(ccdb, cluster = TRUE)
```

fancy_name_contigs	<i>Generate a legible name for a series of contigs</i>
--------------------	--

Description

Generate a legible name for a series of contigs

Usage

```
fancy_name_contigs(contig_tbl, prefix)
```

Arguments

contig_tbl	An all_contig_annotations.csv file, output from VDJ Cell ranger. Importantly, this should contain columns chain, v_gene, d_gene, j_gene
prefix	an optional prefix added to each contig, eg, possibly a sample id.

Value

character

Examples

```
library(dplyr)
contig_anno_path = system.file('extdata', 'all_contig_annotations_balbc_1.csv.xz',
  package = 'CellaRepertorium')
contig_anno = readr::read_csv(contig_anno_path)
contig_anno = contig_anno %>% mutate(fancy_name =
```

```
fancy_name_contigs(., prefix = 'b6_1'))
stopifnot(!any(duplicated(contig_anno$fancy_name)))
```

filter_cdb

Create new or update existing columns of ContigCellDB tables

Description

Create new or update existing columns of ContigCellDB tables

Usage

```
filter_cdb(ccdb, ..., tbl = "contig_tbl")
```

```
mutate_cdb(ccdb, ..., tbl = "contig_tbl")
```

Arguments

ccdb	ContigCellDB()
...	name and value pair of column that will be updated
tbl	character. One of contig_tbl, cell_tbl or cluster_tbl, naming the table to be updated.

Value

ContigCellDB object with updated table

Functions

- filter_cdb: Filter rows of a table in a ContigCellDB object

See Also

[dplyr::mutate\(\)](#)

[dplyr::filter\(\)](#)

Examples

```
data(ccdb_ex)
subset_contig = filter_cdb(ccdb_ex,full_length, productive == 'True',
high_confidence, chain != 'Multi', nchar(cdr3) > 5)
subset_cell = filter_cdb(ccdb_ex, sample == 4, tbl = 'cell_tbl')
data(ccdb_ex)
new_contig = mutate_cdb(ccdb_ex, new_col = 1)
new_cell = mutate_cdb(ccdb_ex, new_col = 1, tbl = 'contig_tbl')
```

fine_clustering	<i>Perform additional clustering of sequences within groups</i>
-----------------	---

Description

Perform additional clustering of sequences within groups

Usage

```
fine_clustering(
  ccdb,
  sequence_key,
  type,
  max_affinity = NULL,
  keep_clustering_details = FALSE,
  ...
)
```

Arguments

ccdb	A ContigCellDB() object
sequence_key	character naming column in <code>contig_tbl</code> with sequence
type	'AA' or 'DNA'
max_affinity	numeric naming the maximal affinity for the sparse affinity matrix that is constructed. Not currently used.
keep_clustering_details	logical – should output of <code>fine_cluster_seqs</code> be kept as a list column
...	Arguments passed on to fine_cluster_seqs
	<code>big_memory_brute</code> attempt to cluster more than 4000 sequences? Clustering is quadratic, so this will take a long time and might exhaust memory
	<code>method</code> one of 'substitutionMatrix' or 'levenshtein'
	<code>substitution_matrix</code> a character vector naming a substitution matrix available in Biostrings, or a substitution matrix itself

Value

[ContigCellDB\(\)](#) object with updated `contig_tbl` and `cluster_tbl`

Examples

```
library(dplyr)
data(ccdb_ex)
ccdb_ex_small = ccdb_ex
ccdb_ex_small$cell_tbl = ccdb_ex_small$cell_tbl[1:200,]
ccdb_ex_small = cdhit_ccdb(ccdb_ex_small,
  sequence_key = 'cdr3_nt', type = 'DNA', cluster_name = 'DNA97',
```

```

identity = .965, min_length = 12, G = 1)
ccdb_ex_small = fine_clustering(ccdb_ex_small, sequence_key = 'cdr3_nt', type = 'DNA')

# Canonicalize with the medoid contig is probably what is most common
ccdb_medoid = canonicalize_cluster(ccdb_ex_small)

# But there are other possibilities.
# To pass multiple "AND" filter arguments must use &
ccdb_umi = canonicalize_cluster(ccdb_ex_small,
contig_filter_args = chain == 'TRA' & length > 500, tie_break_keys = 'umis',
contig_fields = c('chain', 'length'))
ccdb_umi$cluster_tbl %>% dplyr::select(chain, length) %>% summary()

```

fine_cluster_seqs	<i>Calculate distances and perform hierarchical clustering on a set of sequences</i>
-------------------	--

Description

The distances between AA sequences is defined to be $1 - \text{score} / \max(\text{score})$ times the median length of the input sequences. The distances between nucleotide sequences is defined to be $\text{edit_distance} / \max(\text{edit_distance})$ times the median length of input sequences.

Usage

```

fine_cluster_seqs(
  seqs,
  type = "AA",
  big_memory_brute = FALSE,
  method = "levenshtein",
  substitution_matrix = "BLOSUM100",
  cluster_fun = "none",
  cluster_method = "complete"
)

```

Arguments

seqs	character vector, DNASTringSet or AAStringSet
type	character either AA or DNA specifying type of seqs
big_memory_brute	attempt to cluster more than 4000 sequences? Clustering is quadratic, so this will take a long time and might exhaust memory
method	one of 'substitutionMatrix' or 'levenshtein'
substitution_matrix	a character vector naming a substitution matrix available in Biostrings, or a substitution matrix itself
cluster_fun	character, one of "hclust" or "none", determining if distance matrices should also be clustered with hclust
cluster_method	character passed to hclust

Value

list

See Also`hclust()`, `Biostrings::stringDist()`**Examples**

```

fasta_path = system.file('extdata', 'demo.fasta', package='CellaRepertorium')
aaseq = Biostrings::readAAStringSet(fasta_path)[1:100]
cls = fine_cluster_seqs(aaseq, cluster_fun = 'hclust')
plot(cls$cluster)

```

generate_pseudobulk	<i>Generate "pseudobulk" data from a ContigCellDB</i>
---------------------	---

Description

Tabulate contigs with a unique combination of class_keys per total_keys. For instance, total_keys might be a sample identifier, and class_keys might be the V- and J- gene identities. The idea is that this might mimic the data generated in a bulk experiment.

Usage

```
generate_pseudobulk(ccdb, class_keys, total_keys, type = c("cell", "umi"))
```

Arguments

ccdb	<code>ContigCellDB()</code>
class_keys	character naming fields in <code>contig_tbl</code> that define unique classes of the repertoire
total_keys	character naming fields to be conditioned upon when calculating the total.
type	one of "cell" or "umi"

Details

This function is currently rather 10x-specific, in that it is assumed that columns barcode and umis exist.

Value

tibble

Examples

```

data(ccdb_ex)
ccdb_ex = cluster_germline(ccdb_ex)
pseudo = generate_pseudobulk(ccdb_ex, c('v_gene', 'j_gene', 'chain'), c('pop', 'sample'))

```

guess_celltype	<i>Guess the cell type of a contig from the chain ID</i>
----------------	--

Description

This function is likely dependent on annotations from 10X and may change or break as their pipeline changes.

Usage

```
guess_celltype(chain)
```

Arguments

chain	character which will be parsed to try to infer celltype
-------	---

Value

contig table with celltype column

See Also

[crosstab_by_celltype\(\)](#)

Examples

```
data(ccdb_ex)
table(guess_celltype(ccdb_ex$contig_tbl$chain))
```

hushWarning	<i>Selectively muffle warnings based on output</i>
-------------	--

Description

Selectively muffle warnings based on output

Usage

```
hushWarning(expr, regexp)
```

Arguments

expr	an expression
regexp	a regexp to be matched (with str_detect)

Value

the result of expr

Examples

```
CellaRepertorium::hushWarning(warning('Beware the rabbit'), 'rabbit')
CellaRepertorium::hushWarning(warning('Beware the rabbit'), 'hedgehog')
```

ig_chain_recode	<i>Categorize the pairing present in a cell</i>
-----------------	---

Description

For each cell (defined by ccdb\$cell_pk) count the number of each level of chain_key occurs, and cross tabulate. Also for each cell, paste together all values chain_key. Return a tibble, keyed by cells that includes the counts of the chains, the raw_chain_type and any additional output from running chain_recode_fun.

Usage

```
ig_chain_recode(tbl)

tcr_chain_recode(tbl)

enumerate_pairing(ccdb, chain_key = "chain", chain_recode_fun = NULL)
```

Arguments

tbl	output from enumerate_pairing containing TRA/TRB or IGH/IHK/IHL columns
ccdb	ContigCellDB
chain_key	character naming the field in the contig_tbl identifying chain
chain_recode_fun	a function that operates on the output of this function that further reduces the chain combinations to some other summary. Set to 'guess' to apply functions that may work for 10X data or NULL to skip. See CellaRepertorium::tcr_chain_recode for an example.

Value

a tibble keyed by cells.

Functions

- ig_chain_recode: Recode a table with IG chains
- tcr_chain_recode: Recode a table with TCR chains

Examples

```
data(ccdb_ex)
enumerate_pairing(ccdb_ex)
enumerate_pairing(ccdb_ex, chain_recode_fun = 'guess')
```

map_axis_labels	<i>Color axis labels</i>
-----------------	--------------------------

Description

Color axis labels

Usage

```
map_axis_labels(
  plt,
  label_data_x = NULL,
  label_data_y = NULL,
  aes_label,
  scale = ggplot2::scale_color_hue(aesthetics = "axis_color")
)
```

Arguments

plt	<code>ggplot2::ggplot()</code> object
label_data_x	<code>data.frame()</code> containing the mapping between x-axis labels and aes_label
label_data_y	<code>data.frame()</code> containing the mapping between y-axis labels and aes_label
aes_label	character or bare symbol giving the column in label_data to be mapped
scale	ggplot2 discrete color

Value

plt with axis text modified

Examples

```
require(ggplot2)
require(dplyr)
plt = ggplot(mpg, aes(x = manufacturer, y = drv)) + geom_jitter()
label_data = mpg %>% select(manufacturer) %>% unique() %>%
mutate(euro = manufacturer %in% c('audi', 'volkswagen'))
map_axis_labels(plt, label_data_x = label_data, aes_label = euro)
```

pairing_tables	<i>Generate a list of tables representing clusters paired in cells</i>
----------------	--

Description

A contingency table of every combination of `cluster_idx` up to `table_order` is generated. Combinations that are found in at least `min_expansion` number of cells are reported. All cells that have these combinations are returned, as well as cells that only have `orphan_level` of matching `cluster_idx`.

Usage

```
pairing_tables(
  ccdb,
  ranking_key = "grp_rank",
  table_order = 2,
  min_expansion = 2,
  orphan_level = 1,
  cluster_keys = character(),
  cluster_whitelist = NULL,
  cluster_blacklist = NULL
)
```

Arguments

<code>ccdb</code>	ContigCellDB
<code>ranking_key</code>	field in <code>ccdb\$contig_tbl</code> giving the ranking of each contig per cell. Probably generated by a call to rank_prevalence_ccdb() or rank_chain_ccdb() .
<code>table_order</code>	Integer larger than 1. What order of <code>cluster_idx</code> will be paired, eg, <code>order = 2</code> means that the first and second highest ranked contigs will be sought and paired in each cell
<code>min_expansion</code>	the minimal number of times a pairing needs to occur for it to be reported
<code>orphan_level</code>	Integer in interval <code>[1, table_order]</code> . Given that at least <code>min_expansion</code> cells are found that have <code>table_order</code> chains identical, how many <code>cluster_idx</code> pairs will we match on to select other cells. Example: <code>orphan_level=1</code> means that cells that share just a single chain with an expanded pair will be reported.
<code>cluster_keys</code>	optional character naming additional columns in <code>ccdb\$cluster_tbl</code> to be reported in the pairing
<code>cluster_whitelist</code>	a table of pairings or clusters that should always be reported. Here the clusters must be named "cluster_idx.1", "cluster_idx.2" (if order-2 pairs are being selected) rather than with "ccdb\$cluster_pk"
<code>cluster_blacklist</code>	a table of pairings or clusters that will never be reported. Must be named as per <code>cluster_whitelist</code> .

Details

For example, if `table_order=2` and `min_expansion=2` then heavy/light or alpha/beta pairs found two or more times will be returned (as well as alpha-alpha pairs, etc, if those are present). If `orphan_level=1` then all cells that share just a single chain with an expanded clone will be returned.

The `cluster_idx.1_fct` and `cluster_idx.2_fct` fields in `cell_tbl`, `idx1_tbl`, `idx2_tbl` are cast to factors and ordered such that pairings will tend to occur along the diagonal when they are cross-tabulated. This facilitates plotting.

Value

list of tables. The `cell_tbl` is keyed by the `cell_identifiers`, with fields "cluster_idx.1", "cluster_idx.2", etc, IDing the contigs present in each cell. "cluster_idx.1_fct" and "cluster_idx.2_fct" cast these fields to factors and are reordered to maximize the number of pairs along the diagonal. The `idx1_tbl` and `idx2_tbl` report information (passed in about the `cluster_idx` by `feature_tbl`.) The `cluster_pair_tbl` reports all pairings found of contigs, and the number of times observed.

See Also

[rank_prevalence_ccdb\(\)](#)

Examples

```
library(dplyr)
tbl = tibble(clust_idx = gl(3, 2), cell_idx = rep(1:3, times = 2), contig_idx = 1:6)
ccdb = ContigCellDB(tbl, contig_pk = c('cell_idx', 'contig_idx'),
  cell_pk = 'cell_idx', cluster_pk = 'clust_idx')
# add `grp_rank` to ccdb$contig_tbl indicating how frequent a cluster is
ccdb = rank_prevalence_ccdb(ccdb, tie_break_keys = character())
# using `grp_rank` to determine pairing
# no pairs found twice
pt1 = pairing_tables(ccdb)
# all pairs found, found once.
pt2 = pairing_tables(ccdb, min_expansion = 1)
pt2$cell_tbl
tbl2 = bind_rows(tbl, tbl %>% mutate(cell_idx = rep(4:6, times = 2)))
ccdb2 = ContigCellDB(tbl2, contig_pk = c('cell_idx', 'contig_idx'), cell_pk = 'cell_idx',
  cluster_pk = 'clust_idx') %>% rank_prevalence_ccdb(tie_break_keys = character())
#all pairs found twice
pt3 = pairing_tables(ccdb2, min_expansion = 1)
pt3$cell_tbl
ccdb2$contig_tbl = ccdb2$contig_tbl %>%
  mutate(umis = 1, reads = 1, chain = rep(c('TRA', 'TRB'), times = 6))
ccdb2 = rank_chain_ccdb(ccdb2, tie_break_keys = character())
pt4 = pairing_tables(ccdb2, min_expansion = 1, table_order = 2)
```

plot_cluster_factors *Visualization of pairs of cluster factor*

Description

With factors, a pair of variables present in the `contig_tbl` and the `cluster_tbl`, generate and plot cross-tabs of the number of contigs, or its pearson residual.

Usage

```
plot_cluster_factors(
  ccdb,
  factors,
  type = c("heatmap", "network"),
  statistic = c("pearson", "contigs"),
  ncluster = 0,
  chaintype
)
```

Arguments

<code>ccdb</code>	A ContigCellDB object.
<code>factors</code>	character length 2 of fields present
<code>type</code>	Type of visualization, a heatmap or a node-edge network plot
<code>statistic</code>	Cluster characteristics visualized by pearson residuals or raw contig counts
<code>ncluster</code>	integer. Omit factors that occur less than nclusters. For clarity of visualization.
<code>chaintype</code>	Character in <code>ccdb\$contig_tbl\$chain</code> . If passed will subset contigs belonging to specified chain (IGH,IGK,IGL,TRA,TRB)

Value

A `ggraph` object if `type == 'network'`, and a `ggplot` object if `type == 'heatmap'`

See Also

`canonicalize_cluster` to "roll-up" additional contig variables into the `'cluster_tbl'`

Examples

```
library(ggraph)
data(ccdb_ex)
ccdb_germline_ex = cluster_germline(ccdb_ex, segment_keys = c('v_gene', 'j_gene', 'chain'),
  cluster_pk = 'segment_idx')
ccdb_germline_ex = fine_clustering(ccdb_germline_ex, sequence_key = 'cdr3_nt', type = 'DNA')
plot_cluster_factors(ccdb_germline_ex, factors = c('v_gene', 'j_gene'),
  statistic = 'pearson', type = 'network' ,ncluster = 10, chaintype = 'TRB')
```

```
plot_cluster_factors(ccdb_germline_ex, factors = c('v_gene', 'j_gene'),
  statistic = 'contigs', type = 'heatmap')
plot_cluster_factors(ccdb_germline_ex, factors = c('v_gene', 'j_gene'),
  statistic = 'contigs', type = 'network', ncluster = 10)
```

plot_permute_test	<i>Plot a histogram of permuted vs observed test statistic</i>
-------------------	--

Description

Plot a histogram of permuted vs observed test statistic

Usage

```
plot_permute_test(perm_test)

## S3 method for class 'PermuteTestList'
tidy(x, ...)

## S3 method for class 'PermuteTest'
tidy(x, ...)

## S3 method for class 'PermuteTest'
print(x, ...)

## S3 method for class 'PermuteTestList'
print(x, max = 3, ...)
```

Arguments

perm_test	PermuteTest or PermuteTestList output from cluster_permute_test()
x	PermuteTestList
...	ignored
max	maximum number of components to print

Methods (by generic)

- tidy: return permutations run using a sequence of contrasts as a tibble
- tidy: return permutations as a tibble
- print: pretty-print
- print: pretty-print

See Also

cluster_permute_test

purity	<i>Calculate number of cluster-subject singletons for the purposes of permutation testing</i>
--------	---

Description

Calculate number of cluster-subject singletons for the purposes of permutation testing

Usage

```
purity(cluster_idx, subject)
```

Arguments

cluster_idx	factor-like cluster variable
subject	factor-like subject

Value

average number of singletons

See Also

[cluster_permute_test\(\)](#)

Examples

```
message("see example(cluster_permute_test)")
```

rank_prevalence_ccdb	<i>Rank contigs, per cell, by experiment-wide prevalence of cluster_pk, which is added as the prevalence field</i>
----------------------	--

Description

Rank contigs, per cell, by experiment-wide prevalence of cluster_pk, which is added as the prevalence field

Usage

```
rank_prevalence_ccdb(
  ccdb,
  contig_filter_args = TRUE,
  tie_break_keys = c("umis", "reads")
)

rank_chain_ccdb(
  ccdb,
  contig_filter_args = TRUE,
  tie_break_keys = c("umis", "reads"),
  chain_key = "chain",
  contig_fields = tie_break_keys,
  chain_levels = c("IGL", "IGK", "TRA", "TRB", "IGH")
)
```

Arguments

ccdb	ContigCellDB()
contig_filter_args	an expression passed to dplyr::filter() . Unlike filter, multiple criteria must be & together, rather than using commas to separate. These act on ccdb\$contig_tbl
tie_break_keys	(optional) character naming fields in contig_tbl that are used sort the contig table in descending order. Used to break ties if contig_filter_args does not return a unique contig for each cluster
chain_key	character naming the field in contig_tbl to be sorted on.
contig_fields	Optional fields from contig_tbl that will be copied into the cluster_tbl from the canonical contig.
chain_levels	an optional character vector providing the sort order of the chain column in tbl. If set to length zero, then the the ordering will be alphabetical

Value

ContigCellDB with modified contig_tbl

Functions

- rank_chain_ccdb: return a canonical contig by chain type, with TRB/IGH returned first. By default, ties are broken by umis and reads.

Examples

```
data(ccdb_ex)
ccdb_ex = cluster_germline(ccdb_ex)
rank_prev = rank_prevalence_ccdb(ccdb_ex)
rank_prev$contig_tbl
rank_chain = rank_chain_ccdb(ccdb_ex)
rank_chain$contig_tbl
```

rbind,ContigCellDB-method

Combine ContigCellDB along rows (contigs, cells or clusters).

Description

The union of the rows in each of the objects is taken, thus removing any rows that has an exact duplicate. This includes all fields, not just the primary key for that table. The union of the various primary keys is taken.

Usage

```
## S4 method for signature 'ContigCellDB'
rbind(..., deparse.level = 1)
```

Arguments

```
...          ContigCellDB()
deparse.level ignored
```

Value

ContigCellDB()

Examples

```
data(ccdb_ex)
splat = split_cdb(ccdb_ex, 'chain', 'contig_tbl')
unite = equalize_ccdb(rbind(splat$TRA, splat$TRB), sort = TRUE)
stopifnot(all.equal(unite, ccdb_ex))
```

reexports

Turn an object into a tidy tibble

Description

Turn an object into a tidy tibble

Usage

```
tidy(x, ...)
```

Arguments

```
x          An object to be converted into a tidy tibble::tibble().
...        Additional arguments to tidying method.
```

Value

A `tibble::tibble()` with information about model components.

Methods

No methods found in currently loaded packages.

right_join_warn	<i>Perform a <code>dplyr::left_join()</code> but check for non-key overlapping fields</i>
-----------------	---

Description

Perform a `dplyr` join, but either warn if the two tables share non-key fields. If `overwrite = TRUE`, then shared columns will pull from `x` otherwise a suffix will be added to `y`. To perform this check, `by` must be specified, and it is an error if it is not.

Usage

```
right_join_warn(...)
```

```
left_join_warn(x, y, by, overwrite = FALSE, join = left_join, ...)
```

Arguments

<code>...</code>	passed to joining function
<code>x</code>	A pair of data frames, data frame extensions (e.g. a tibble), or lazy data frames (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>y</code>	A pair of data frames, data frame extensions (e.g. a tibble), or lazy data frames (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>by</code>	character specifying columns in <code>x</code> and <code>y</code> to key on.
<code>overwrite</code>	logical – should non-key fields in <code>y</code> be overwritten using <code>x</code> , or should a suffix (" <code>.y</code> ") be added
<code>join</code>	function giving the type of join to perform, eg, left, right, inner, outer.

Value

data.frame or tibble

Functions

- `right_join_warn`: perform a `dplyr::right_join()`

Examples

```
left_join_warn(mtcars, mtcars, by = 'mpg')
left_join_warn(mtcars, mtcars, by = 'mpg', overwrite = TRUE)
```

split_cdb	<i>Split into a list of ContigCellDB() by named fields</i>
-----------	--

Description

Split into a list of [ContigCellDB\(\)](#) by named fields

Usage

```
split_cdb(ccdb, fields, tbl = "contig_tbl", drop = FALSE, equalize = TRUE)
```

Arguments

ccdb	ContigCellDB()
fields	character naming fields in tbl
tbl	one of contig_tbl, cell_tbl or cluster_tbl
drop	logical indicating if levels that do not occur should be dropped (if f is a factor or a list).
equalize	logical. Should the contig, cells and clusters be equalized by taking the intersection of their common keys?

Value

list of ContigCellDB

Examples

```
data(ccdb_ex)
splat = split_cdb(ccdb_ex, 'chain', 'contig_tbl')
stopifnot(all(splat$TRA$contig_tbl$chain == 'TRA'))
stopifnot(all(splat$TRB$contig_tbl$chain == 'TRB'))
```

[[,ContigCellDB,character,missing-method	
	<i>data.frame-like mutation/accessor generics for ContigCellDB objects</i>

Description

A ContigCellDB pretend to be a cell_tbl data.frame in several regards. This is to enable nesting ContigCellDB objects in the colData of a SingleCellExperiment and so that various plotting functionality in scater can do something sensible.

Usage

```
## S4 method for signature 'ContigCellDB,character,missing'
x[[i, j, ...]]

## S4 method for signature 'ContigCellDB,ANY,missing,ANY'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'ContigCellDB'
dim(x)

## S4 method for signature 'ContigCellDB'
dimnames(x)

## S4 method for signature 'ContigCellDB'
nrow(x)

## S4 method for signature 'ContigCellDB'
ncol(x)
```

Arguments

<code>x</code>	ContigCellDB
<code>i</code>	integer or character index
<code>j</code>	ignored
<code>...</code>	ignored
<code>drop</code>	ignored

Details

If `x` a ContigCellDB, then `dim(x)` and `dimnames(x)` return `dim(x$cell_tbl)` and `dimnames(x$cell_tbl)`, respectively, and `x[[col]]` returns `x$cell_tbl[[col]]`. Likewise indexing with `x[i,]` returns cells indexed by `i`. Finally `as.data.frame(x)` returns `x$cell_tbl`.

Value

See details.

Examples

```
data(ccdb_ex)
ccdb_ex[1:10,]
head(ccdb_ex[['barcode']])
dim(ccdb_ex)
dimnames(ccdb_ex)
```

\$.ContigCellDB-method *Access public members of ContigCellDB object.*

Description

Modification to members will trigger various forms of equalization. See [equalize_ccdb\(\)](#) for details.

Usage

```
## S4 method for signature 'ContigCellDB'
x$name

## S4 replacement method for signature 'ContigCellDB'
x$name <- value
```

Arguments

x	A ContigCellDB object
name	a slot of a ContigCellDB object (one of c('contig_tbl', 'cell_tbl', 'contig_pk', 'cell_pk', 'cluster_tbl', 'cluster_pk'))
value	The value assigned to a slot of ContigCellDB object

Value

Update or return a slot of [ContigCellDB\(\)](#)

See Also

[equalize_ccdb\(\)](#)

Examples

```
data(ccdb_ex)
ccdb_ex$contig_tbl
ccdb_ex$cell_tbl
ccdb_ex$cluster_tbl
data(ccdb_ex)
ccdb_ex$contig_pk = c("sample", "barcode", "contig_id") # 'pop' is technically redundant with 'sample'
# Take a subset of ccdb_ex
ccdb_ex
ccdb_ex$contig_tbl = dplyr::filter(ccdb_ex$contig_tbl, pop == 'b6')
ccdb_ex
```

Index

- * **datasets**
 - ccdb_ex, [7](#)
 - contigs_qc, [19](#)
 - .cluster_permute_test, [4](#)
 - [,ContigCellDB,ANY,missing,ANY-method
 - ([[,ContigCellDB,character,missing-method), [39](#)
 - [,ContigCellDB,ANY,missing-method
 - ([[,ContigCellDB,character,missing-method), [39](#)
 - [[,ContigCellDB,character,missing-method, [39](#)
 - \$,ContigCellDB-method, [41](#)
 - \$<-,ContigCellDB-method
 - (\$,ContigCellDB-method), [41](#)
- Biostrings::stringDist(), [27](#)
- canonicalize_cell, [5](#)
- canonicalize_cell(), [7](#)
- canonicalize_cluster, [6](#)
- canonicalize_cluster(), [5](#)
- ccdb_ex, [7](#)
- ccdb_join, [8](#)
- cdhit, [9](#), [11](#)
- cdhit(), [10](#), [11](#)
- cdhit_ccdb, [10](#)
- cland, [11](#)
- cluster_filterset, [12](#)
- cluster_filterset(), [16](#)
- cluster_germline, [13](#)
- cluster_logistic_test
 - (cluster_test_by), [16](#)
- cluster_permute_test, [13](#)
- cluster_permute_test(), [35](#)
- cluster_plot, [15](#)
- cluster_test_by, [16](#)
- ContigCellDB, [17](#)
- ContigCellDB(), [5–8](#), [10](#), [11](#), [13](#), [16](#), [18](#), [22](#), [24](#), [25](#), [27](#), [36](#), [37](#), [39](#), [41](#)
- ContigCellDB-mutate
 - (\$,ContigCellDB-method), [41](#)
- ContigCellDB_10XVDJ (ContigCellDB), [17](#)
- contigs_qc, [8](#), [19](#)
- cross_tab_tbl, [20](#)
- cross_tab_by_celltype, [20](#)
- crosstab_by_celltype(), [28](#)
- data, frame(), [30](#)
- dim,ContigCellDB-method
 - ([[,ContigCellDB,character,missing-method), [39](#)
- dimnames,ContigCellDB-method
 - ([[,ContigCellDB,character,missing-method), [39](#)
- dplyr::filter(), [5](#), [6](#), [17](#), [24](#), [36](#)
- dplyr::mutate(), [24](#)
- entropy, [21](#)
- enumerate_pairing (ig_chain_recode), [29](#)
- equalize_ccdb, [22](#)
- equalize_ccdb(), [41](#)
- fancy_name_contigs, [23](#)
- filter_cdb, [24](#)
- filter_cdb(), [18](#)
- fine_cluster_seqs, [25](#), [26](#)
- fine_clustering, [25](#)
- generate_pseudobulk, [27](#)
- ggplot2::ggplot(), [30](#)
- guess_celltype, [28](#)
- guess_celltype(), [20](#)
- hclust(), [27](#)
- hushWarning, [28](#)
- ig_chain_recode, [29](#)
- left_join_warn (right_join_warn), [38](#)
- left_join_warn(), [7](#)

map_axis_labels, [30](#)
modal_category (entropy), [21](#)
mutate_cdb (filter_cdb), [24](#)
mutate_cdb(), [18](#)

ncol, ContigCellDB-method
 ([[, ContigCellDB, character, missing-method),
 [39](#)
np (entropy), [21](#)
nrow, ContigCellDB-method
 ([[, ContigCellDB, character, missing-method),
 [39](#)

order(), [22](#)

pairing_tables, [31](#)
plot_cluster_factors, [33](#)
plot_permute_test, [34](#)
print.PermuteTest (plot_permute_test),
 [34](#)
print.PermuteTestList
 (plot_permute_test), [34](#)
purity, [35](#)
purity(), [14](#)

rank_chain_ccdb (rank_prevalence_ccdb),
 [35](#)
rank_chain_ccdb(), [31](#)
rank_prevalence_ccdb, [35](#)
rank_prevalence_ccdb(), [31](#), [32](#)
rbind, ContigCellDB-method, [37](#)
rbind.ContigCellDB
 (rbind, ContigCellDB-method), [37](#)
rbind.ContigCellDB(), [18](#)
reexports, [37](#)
right_join_warn, [38](#)

split_cdb, [39](#)
split_cdb(), [18](#)

tcr_chain_recode (ig_chain_recode), [29](#)
tibble::tibble(), [37](#), [38](#)
tidy (reexports), [37](#)
tidy.PermuteTest (plot_permute_test), [34](#)
tidy.PermuteTestList
 (plot_permute_test), [34](#)