Package ‘Platypus’

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Type Package

Title Single-Cell Immune Repertoire and Gene Expression Analysis

Description We present 'Platypus', an open-source software platform providing a user-friendly interface to investigate B-cell receptor and T-cell receptor repertoires from scSeq experiments. 'Platypus' provides a framework to automate and ease the analysis of single-cell immune repertoires while also incorporating transcriptional information involving unsupervised clustering, gene expression and gene ontology (Yermanos A, et al (2021) <doi:10.1093/nargab/lqab023>).

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Maintainer Alexander Yermanos <ayermanos@gmail.com>

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**Author**  Alexander Yermanos [aut, cre],  
Andreas Agrafiotis [ctb],  
Raphael Kuhn [ctb],  
Danielle Shlesinger [ctb],  
Jiami Han [ctb],  
Tudor-Stefan Cotet [ctb],  
Victor Kreiner [ctb]

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Description

AntibodyForest takes the output of either ConvertStructure or CsvToDf or SubRepertoires or RemoveNets and outputs B cell phylogenetic networks in tree format. There is also the possibility to give the full-length list of clonal lineages, which contains both isotype and transcriptional cluster information, only when no prior data transformation is desired. Each network represents a clonal lineage, referring to the number of B cell receptor sequences originating from an independent V(D)J recombination event. Each vertex represents a unique recovered full-length variable heavy and light chain antibody sequence of a clonal family. Edges separating nodes are drawn given that clonal variants are similarly related according to their Levenshtein distance. Edge weights are extracted from the distance matrix apart from the special case of unmutated germline, in which the weights of outgoing edges from it are either set to 1 or to the difference between the corresponding distance from the matrix and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. At tree building, starting from the reference ancestral germline, each node is connected to nodes that can be reached via the minimum distance based on the distance matrix calculation. Therefore, potential edges that go back to previous tree layers along with bidirectional circles are eliminated. Polytomies, displayed by B cell clones producing multiple distinct offsprings, are resolved in case of reaching nodes with equal minimum distance. Indeed, the algorithm removes edges either randomly from the recipient nodes, based on the node closest or farthest from the germline, considering the number of intermediate nodes or edge path length, or the highest/lowest counting of cells on the present node. Additional ties are settled by random edge selection. Consequently, parsimony holds, meaning that each daughter node has only one parent. Distinct tree topologies enable to visually investigate the trade-off between balance and evolution, and further quantify the amount of diversification of the subsequent detected clonal abundant clones during somatic hypermutation and class switching. The minimum decision-based criterion determines the amount of balance presented in the tree, while the maximum decision-based method the amount of evolution presented in the tree. Single color or color distribution on each node demonstrates the proportion of B cells with the specific isotype(s) or transcriptional cluster(s), while setting the size of vertices can be performed based on the number of unique sequences per clone, vertex betweenness and vertex closeness. Scaling of nodes by their relative clonal expansion assists in pinpointing identical antibody sequences across a multitude of B cells. Node labeling can depict clonal frequency.

Usage

AbForests_AntibodyForest(
    full_list,
    csv,
    files,
    distance_mat,
    clonal_frequency,
    scaleByClonalFreq,
    weight,
AbForests_AntibodyForest

tie_flag,  
scaleBybetweenness,  
scaleByclocloseness_metr,  
opt,  
random.seed,  
alg_opt,  
cdr3
)

Arguments

full_list  
a list of clone lineages, represented as data.frames

csv  
an indicator variable. TRUE if full_list argument is a list of csv files, FALSE otherwise

files  
a list of data.frames. Each data.frame contains 2 columns, one that describes the sequences and the other which type of information (isotype or cluster) is considered in the analysis. All these cases are determined by the user.

distance_mat  
a custom integer distance matrix, or NULL for using the default distance matrix (calucated based on the levenshtein distance, which counts the number of mutations between sequences).

clonal_frequency  
a logical variable, TRUE if labeling of vertices is based on clonal frequency and FALSE otherwise.

scaleByClonalFreq  
logical variable with TRUE if vertex size is scaled by the number of unique sequences per clone and FALSE otherwise.

weight  
logical variable. When its value is FALSE, then the weights of outgoing edges from Germline node are set to 1. When its value is TRUE, the weights are set to the difference between the number of mutations among sequences in germline and connected nodes(value in the corresponding distance matrix) and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. In both cases, weights of remaining edges are extracted from the distance matrix. Outgoing edges from Germline represent the number of mutations of sequences having as common ancestor the Germline.

tie_flag  
a string, with options 'rand', 'full', 'close_to_germ', 'far_from_germ', 'close_path_to_germ', 'far_path_from_germ', 'most_expanded' and 'least_expanded' for removing edges when equal distance (tie) in distance matrix. 'rand' means random pruning in one of nodes, 'full' means keeping all nodes, close_to_germ means pruning of node(s) farthest from germline (based on number of intermediate nodes), far_from_germ means pruning of node(s) closest to germline (based on number of intermediate nodes), close_path_to_germ means pruning of node(s) farthest from germline (based on edge path length), far_path_from_germ means pruning of node(s) closest to germline (based on edge path length), most_expanded means pruning of node(s) with the lowest B cell count(clonal frequency) and least_expanded, which means pruning of node(s) with the highest B cell count(clonal frequency). In cases of subsequent ties, a random node is selected.
scaleBybetweenness
    logical variable with TRUE if vertex size is scaled by the vertex betweenness centrality.

toolsBycloseness_metr
    logical variable with TRUE if vertex size is scaled by closeness centrality of vertices in graph.

opt
    a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.

random.seed
    a random seed, specified by the user, when random sampling of sequences happens in each of the cases described in tie_flag argument.

alg_opt
    a string denoting the version of the edge selection algorithm used in the construction of networks. Possible choices: "naive", "two-step".

cdr3
    variable with values 0 if the user desires to select full length sequences (only when the input is a list of csv files), 1 for sequences in the CDR3 only (only when the input is a list of csv files) and NULL otherwise.

Value

graphs. A list of lists. E.g graphs[[1]][[1]] network: an igraph object, containing the first network in tree format. graphs[[1]][[2]] legend: contains the legend parameters of the first network. graphs[[1]][[3]] count.rand: contains the number of randomly considered nodes for the first network. graphs[[1]][[4]] adj.matrix: contains the adjacency matrix for the first network. graphs[[1]][[5]] distance.matrix: contains the distance matrix for the first network. graphs[[1]][[6]] cells.per.network: contains the number of cells for the first network. graphs[[1]][[7]] variants.per.network: contains the number of variants for the first network. graphs[[1]][[8]] variant.sequences: contains the sequences of the variants for the first network. graphs[[1]][[9]] cells.per.variant: contains the number of cells per variant (clonal frequency) for the first network. graphs[[1]][[10]] cell.indices.per.variant: the indices of cells per variant for the first network. graphs[[1]][[11]] new.variant.names: contains the names of variants for the first network. graphs[[1]][[12]] germline.index: contains the index of germline sequence for the first network. graphs[[1]][[13]] isotype.per.variant: contains the isotypes corresponding to each variant for the first network. graphs[[1]][[14]] transcriptome.cluster.per.variant: contains the transcriptional clusters corresponding to each variant for the first network. graphs[[1]][[15]] isotype.per.cell: contains the isotype corresponding to each cell for the first network. graphs[[1]][[16]] transcriptome.cluster.per.cell: contains the transcriptional cluster corresponding to each cell for the first network.

See Also

ConvertStructure, CsvToDf, SubRepertoires, RemoveNets

Examples

```
## Not run:
AbForests_AntibodyForest(full_list = Platypus::new, csv=FALSE, files, clonal_frequency=TRUE, scaleByClonalFreq=TRUE, weight=TRUE, tie_flag='close_to_germ', scaleBybetweenness=FALSE, scaleBycloseness_metr=FALSE, opt="cluster", alg_opt="0", cdr3=NULL)
```
AbForests_CompareForests

Comparison of distinct B cell repertoires

Description

CompareForests takes the output of AntibodyForest for 2 distinct repertoires and performs a comparison of these 2 repertoires.

Usage

AbForests_CompareForests(
  list1,
  list2,
  DAG,
  clonal_frequency,
  scaleByClonalFreq,
  weight,
  tie_flag,
  opt
)

Arguments

list1
  a list of lists. Each sublist contains an igraph object with the networks of the evolved B clonal lineages in tree format, their legend and the number of randomly considered nodes per network for Repertoire of 1 (Output of AntibodyForest). E.g. list1[[1]][[1]] is an igraph object, containing the first network of the evolved B clonal lineage in tree format. list1[[1]][[2]] contains the legend parameters of the first network of the evolved B clonal lineage. list1[[1]][[3]] is the number of randomly considered nodes for the first network of the evolved B clonal lineage.

list2
  a list of lists. Each sublist contains an igraph object with the networks of the evolved B clonal lineages in tree format, their legend and the number of randomly considered nodes per network for Repertoire of 2 (Output of AntibodyForest). E.g. list2[[1]][[1]] is an igraph object, containing the first network of the evolved B clonal lineage in tree format. list2[[1]][[2]] contains the legend parameters of the first network of the evolved B clonal lineage. list2[[1]][[3]] is the number of randomly considered nodes for the first network of the evolved B clonal lineage.

DAG
  a logical variable, when TRUE a directed acyclic graph is produced.

clonal_frequency
  a logical variable, TRUE if labeling of vertices is based on clonal frequency and FALSE otherwise.
scaleByClonalFreq
logical variable with TRUE if vertex size is scaled by the number of unique sequences per clone and FALSE otherwise.

weight
logical variable. When its value is FALSE, then the weights of outgoing edges from Germline node are set to 1. When its value is TRUE, the weights are set to the difference between the number of mutations among sequences in germline and connected nodes (value in the corresponding distance matrix) and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. In both cases, weights of remaining edges are extracted from the distance matrix. Outgoing edges from Germline represent the number of mutations of sequences having as common ancestor the Germline.

tie_flag
a string, with options 'rand', 'full', 'close_to_germ', 'far_from_germ', 'close_path_to_germ', 'far_path_from_germ', 'most_expanded' and 'least_expanded' for removing edges when equal distance (tie) in distance matrix. 'rand' means random pruning in one of nodes, 'full' means keeping all nodes, close_to_germ means pruning of node(s) farthest from germline (based on number of intermediate nodes), far_from_germ means pruning of node(s) closest to germline (based on number of intermediate nodes), close_path_to_germ means pruning of node(s) farthest from germline (based on edge path length), far_path_from_germ means pruning of node(s) closest to germline (based on edge path length), most_expanded means pruning of node(s) with the lowest B cell count (clonal frequency) and least_expanded, which means pruning of node(s) with the highest B cell count (clonal frequency). In cases of subsequent ties, a random node is selected.

opt
a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.

Value

isotype_info_rep1 A data.frame. It summarizes isotype/cluster info for repertoire 1.

isotype_info_rep2 A data.frame. It summarizes isotype/cluster info for repertoire 2.
AbForests_ConvertStructure

Extract transcriptome/isotype information and B cell receptor sequences from single cell immune repertoire formatted as list of data.frames

Description

ConvertStructure alters a list of clone lineages, represented as data.frames and recovers the type of isotypes or transcriptional clusters and antibody sequences from these clone lineages. It can receive as input the original data or the output of SubRepertoiresByUniqueSeq or SubRepertoiresByCells. Then, the output list is used as input to the RemoveNets or AntibodyForest function.

Usage

AbForests_ConvertStructure(list, opt, cdr3)

Arguments

list a list of data.frames. Each data.frame may contain information concerning full length heavy and light chain sequences, CDRH3 and CDRL3 sequences, the type of isotype and the transcriptional cluster that corresponds to each of these sequences.

opt a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.

cdr3 variable with values 0 if the user desires to select full length sequences (only when the input is a list of csv files), 1 for sequences in the CDR3 only (only when the input is a list of csv files) and NULL otherwise.

Value

list a list of data.frames. Each data.frame contains 2 columns, one that describes the sequences and the other which type of information (isotype or cluster) is considered in the analysis. All these cases are determined by the user.
AbForests_CsvToDf

See Also
RemoveNets, AntibodyForest

Examples

## Not run:
ConvertStructure(list, opt="cluster", cdr3=NULL)
ConvertStructure(list, opt="isotype", 1)

## End(Not run)

AbForests_CsvToDf

Convert list of csvs, to nested list of data.frames

Description

CsvToDf converts a list of csv files, which are clone lineages to a list of data.frames.

Usage

AbForests_CsvToDf(files)

Arguments

files a list of csv files. Each csv file may contain information concerning full length
heavy and light chain sequences, CDRH3 and CDRL3 sequences, the type of
isotype and the transcriptional cluster that corresponds to each of these se-
quences.

Value

a list of data.frames. Each data.frame contains 2 columns, one that describes the sequences
and the other the type of information (isotype or cluster) is considered in the analysis. All these
cases are determined by the user.

See Also
AntibodyForest

Examples

## Not run:
CsvToDf(files)

## End(Not run)
AbForests_ForestMetrics

Calculate metrics for networks

Description

ForestMetrics takes the output of AntibodyForest and calculates metrics for each of the networks.

Usage

AbForests_ForestMetrics(
  graphs,
  DAG,
  clonal_frequency,
  scaleByClonalFreq,
  weight,
  tie_flag,
  opt
)

Arguments

graphs A list of lists. Each sublist contains an igraph object with the networks of the evolved B clonal lineages in tree format, their legend and the number of randomly considered nodes per network (Output of AntibodyForest function). E.g. graphs[[1]][[1]] is an igraph object, containing the first network of the evolved B clonal lineage in tree format. graphs[[1]][[2]] contains the legend parameters of the first network of the evolved B clonal lineage. graphs[[1]][[3]] is the number of randomly considered nodes for the first network of the evolved B clonal lineage.

DAG a logical variable, when TRUE a directed acyclic graph is produced.

clonal_frequency a logical variable, TRUE if labeling of vertices is based on clonal frequency and FALSE otherwise.

scaleByClonalFreq logical variable with TRUE if vertex size is scaled by the number of unique sequences per clone and FALSE otherwise.

weight logical variable. When its value is FALSE, then the weights of outgoing edges from Germline node are set to 1. When its value is TRUE, the weights are set to the difference between the number of mutations among sequences in germline and connected nodes (value in the corresponding distance matrix) and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. In both cases, weights of remaining edges are extracted from the distance matrix. Outgoing edges from Germline represent the number of mutations of sequences having as common ancestor the Germline.
tie_flag  a string, with options 'rand', 'full', 'close_to_germ', 'far_from_germ', 'close_path_to_germ', 'far_path_from_germ', 'most_expanded' and 'least_expanded' for removing edges when equal distance (tie) in distance matrix. 'rand' means random pruning in one of nodes, 'full' means keeping all nodes, close_to_germ means pruning of node(s) farthest from germline (based on number of intermediate nodes), far_from_germ means pruning of node(s) closest to germline (based on number of intermediate nodes), close_path_to_germ means pruning of node(s) farthest from germline (based on edge path length), far_path_from_germ means pruning of node(s) closest to germline (based on edge path length), most_expanded means pruning of node(s) with the lowest B cell count (clonal frequency) and least_expanded, which means pruning of node(s) with the highest B cell count (clonal frequency). In cases of subsequent ties, a random node is selected.

opt  a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.

Value
metrics. A list of lists. Each list contains various metrics for the quantification of networks. E.g metrics[[1]][[1]] is the weighted Longest path from germline for the first network. metrics[[1]][[2]] is the length of weighted longest shortest path from germline for the first network. metrics[[1]][[3]] is the unweighted Longest path from germline for the first network. metrics[[1]][[4]] is the length of unweighted longest shortest path from germline for the first network. metrics[[1]][[5]] is the weighted shortest path network for the first network. metrics[[1]][[6]] is the unweighted shortest path network for the first network. metrics[[1]][[7]] is the average number of daughter cells for the first network. metrics[[1]][[8]] is the std number of daughter cells for the first network. metrics[[1]][[9]] is the min number of daughter cells for the first network. metrics[[1]][[10]] is the max number of daughter cells for the first network. metrics[[1]][[11]] is a ggplot object that contains the plot of Degree Distribution of daughter cells for the first network. metrics[[1]][[12]] is the weighted vertex degree for the first network. metrics[[1]][[13]] is a ggplot object that contains the plot of unweighted Degree Distribution of daughter cells for the first network. metrics[[1]][[14]] is the average number of isotypes for the first network. metrics[[1]][[15]] is a ggplot object that contains the plot of Distribution of isotypes for the first network. metrics[[1]][[16]] is the Isotypes/Clusters info data.frame with columns Parent, Child and Parent-Child, which contains the type of isotypes/clusters for each pair of nodes (Parent-Child relationship in tree) found in the first network. metrics[[1]][[17]] is a ggplot object that contains the plot of Isotype/Cluster Directionality for the first network. In particular, the frequency of all types of isotypes/clusters for each pair of nodes in the tree is depicted. metrics[[1]][[18]] is the vertex betweenness centrality for the first network. metrics[[1]][[19]] is the edge betweenness centrality for the first network. It is defined by the number of geodesics (shortest paths) going through a vertex according to igraph documentation. metrics[[1]][[20]] is the closeness centrality of vertices for the first network. It is defined by the inverse of the average length of the shortest paths to/from all the other vertices in the graph according to igraph documentation. metrics[[1]][[21]] is a ggplot object that contains the plot of Path length from Germline vs Node Degree for the first network. metrics[[1]][[22]] is a ggplot object that contains the plot of Number of edges from Germline vs Node Degree for the first network. metrics[[1]][[23]] is an igraph object that contains the Isotype/Cluster transition network for the first network. metrics[[1]][[24]] is the
global clustering coefficient for the first network. metrics[[1]][[25]] is the average clustering coefficient for the first network. metrics[[1]][[26]] is the mean clonal expansion for the first network, calculated as the mean of clonal frequencies of all vertices in the network. If the labeling or scaling of nodes in graph is based on clonal frequency (arguments: clonal_frequency==TRUE or scaleByClonalFreq==TRUE), then metrics[[1]][[27]] is the ratio: Number of edges from germline to each node with clonal frequency for the first network. metrics[[1]][[28]] is the mean ratio: Number of edges from germline to each node with clonal frequency for the first network. metrics[[1]][[29]] is a ggplot object that contains the ratio of Number of edges from germline to each node with clonal frequency for the first network. metrics[[1]][[30]] is the mean number of edges from germline for the first network. metrics[[1]][[31]] is the ratio: Total path length from germline to each node with clonal frequency for the first network. metrics[[1]][[32]] is the mean ratio: Total path length from germline to each node with clonal frequency for the first network. metrics[[1]][[33]] is a ggplot object that contains the ratio of Total path length from germline to each node with clonal frequency for the first network. metrics[[1]][[34]] is the mean Total path length from germline for the first network. metrics[[2]][[1]] is the weighted Longest path from germline for the second network.

See Also
AntibodyForest

Examples

```r
## Not run:
ForestMetrics(graphs,DAG=TRUE,clonal_frequency=TRUE, scaleByClonalFreq=TRUE,
weight=TRUE,tie_flag='close_to_germ',opt="cluster")
## End(Not run)
```

Description

PlotGraphs takes as input the output of AntibodyForest or ForestMetrics functions and plots all corresponding networks in the single cell immune repertoire or the corresponding ggplot object, the user specifies, from all clone lineages. The function gives the option in the user to store each tree or ggplot object within the repertoire in pdf format.

Usage

```r
AbForests_PlotGraphs(graphs, no_arg, topdf, filename)
```

Arguments

- **graphs**: a list of networks (Output of AntibodyForest function) or metrics (Output of ForestMetrics function).
- **no_arg**: element of list the user desires to plot: integer value, if the user desires to plot a metric and NULL, if the user desires to plot the networks.
AbForests_PlyloToMatrix

Conversion of phylogenetic tree to distance matrix

Description

PlyloToMatrix converts a previously existing phylogenetic tree to a corresponding distance matrix using the cophenetic distance. Then, there is the option to utilize this custom distance matrix as an input distance matrix to AntibodyForest function. The user is responsible for specifying a correct and valid distance matrix. In particular, the size of distance matrix must match the number of sequences for each network in each repertoire.

Usage

AbForests_PlyloToMatrix(tree_name)

Arguments

tree_name a phylogenetic tree (phylo object).

Value

dist_mat The corresponding distance matrix uses the cophenetic distance between two observations that have been clustered. This distance is defined to be the intergroup dissimilarity at which the two observations are first combined into a single cluster.

See Also

AntibodyForest
AbForests_RemoveNets

Examples

```r
## Not run:
PlyloToMatrix(tree_name)
## End(Not run)
```

AbForests_RemoveNets  
*Filter sub-repertoires with less than N unique sequences or with less than C unique cells*

Description

RemoveNets takes the output of SubRepertoires and performs the filtering of the 5 sub-repertoires. In particular, from these 5 sub-repertoires, networks with number of nodes or number of unique sequences below a specified threshold are eliminated.

Usage

```r
AbForests_RemoveNets(
  list,
  opt,
  distance_mat,
  tie_flag,
  weight,
  N,
  C,
  random.seed,
  alg_opt
)
```

Arguments

- **list**: a list of 5 sub-lists of data.frames. Each sub-list corresponds to the set of networks, in which a majority isotype is specified. `list[[1]]` or `list$list_IGHG` contains the networks, in data.frame format, with more IGG isotypes, `list[[2]]` or `list$list_IGHA` contains the networks, in data.frame format, with more IGA isotypes, `list[[3]]` or `list$list_IGHM` contains the networks, in data.frame format, with more IGM isotypes, `list[[4]]` or `list$list_IGAG` contains the networks, in data.frame format, with a tie in IGA and IGG isotypes and `list[[5]]` or `list$list_other` contains the networks, in data.frame format, with other isotypes apart from the aforementioned combinations. In each sub-list, each data.frame represents a clone lineage and contains 2 columns, one that describes the antibody sequences and the other which type of information (isotype) is considered in the analysis. This list of data.frames has been generated by SubRepertoires function based on the initial data input and user’s preferences.
AbForests_RemoveNets

**opt**
a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.

**distance_mat**
a custom integer distance matrix, or NULL for using the default distance matrix (calculated based on the levenshtein distance, which counts the number of mutations between sequences). Given the phylogenetic tree, a custom-made distance matrix can be produced by PlyloToMatrix function.

**tie_flag**
a string, with options 'rand', 'full', 'close_to_germ', 'far_from_germ', 'close_path_to_germ', 'far_path_from_germ', 'most_expanded' and 'least_expanded' for removing edges when equal distance (tie) in distance matrix. 'rand' means random pruning in one of nodes, 'full' means keeping all nodes, close_to_germ means pruning of node(s) farthest from germline (based on number of intermediate nodes), far_from_germ means pruning of node(s) closest to germline (based on number of intermediate nodes), close_path_to_germ means pruning of node(s) farthest from germline (based on edge path length), far_path_from_germ means pruning of node(s) closest to germline (based on edge path length), most_expanded means pruning of node(s) with the lowest B cell count (clonal frequency) and least_expanded, which means pruning of node(s) with the highest B cell count (clonal frequency). In cases of subsequent ties, a random node is selected.

**weight**
logical variable. When its value is FALSE, then the weights of outgoing edges from Germline node are set to 1. When its value is TRUE, the weights are set to the difference between the number of mutations among sequences in germline and connected nodes (value in the corresponding distance matrix) and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. In both cases, weights of remaining edges are extracted from the distance matrix.

**N**
the threshold of unique sequences below which networks are removed.

**C**
the threshold of unique cells below which networks are removed.

**random.seed**
a random seed, specified by the user, when random sampling of sequences happens in each of the cases described in tie_flag argument.

**alg_opt**
a string denoting the version of the edge selection algorithm used in the construction of networks. "0" means the naive version and "1" the advanced one.

**Value**
list a nested list of 5 sub-lists of data.frames. Each sub-list corresponds to the reduced set of networks according to threshold N, in which a majority isotype is specified. list[[1]] or list$list_IGHG contains the networks, in data.frame format, with more IGG isotypes, list[[2]] or list$list_IHGA contains the networks, in data.frame format, with more IGA isotypes, list[[3]] or list$list_IGHM contains the networks, in data.frame format, with more IGM isotypes, list[[4]] or list$list_IGAG contains the networks, in data.frame format, with a tie in IGA and IGG isotypes and list[[5]] or list$list_other contains the networks, in data.frame format, with other isotypes apart from the aforementioned combinations.

**See Also**
SubRepertoires, SubRepertoiresByUniqueSeq, PlyloToMatrix, AntibodyForest
Examples

## Not run:
RemoveNets(list, opt="cluster", distance_mat=NULL,
tie_flag='close_to_germ', weight=TRUE, N=4, C=NULL, random.seed=165)

## End(Not run)

AbForests_SubRepertoiresByCells

*Split single cell immune repertoire into sub-repertoires by isotype based on number of B cells*

Description

SubRepertoiresByCells separates the single cell immune repertoire into 5 sub-repertoires taking into account the number of cells. The goal is to determine the majority isotype per each network in the immune repertoire. Therefore, each sub-repertoire is dominated by isotype IGG, IGA, IGM, other and if there is an equal number of IGA and IGG isotypes in a network, IGA-IGG category exists respectively. In particular, in case there exists a tie in the number of IGA and IGM, the network is considered to contain IGA as majority isotype, while the same number of IGG and IGM in the network categorize this network as containing IGG as majority isotype. The function receives the output of CsvToDf or original data and can be given as input to ConvertStructure function.

Usage

AbForests_SubRepertoiresByCells(list)

Arguments

list a list of data.frames. Each data.frame represents a clone lineage and separates initial input data into subsets of networks.

Value

list a nested list of 5 sub-lists of data.frames. Each sub-list corresponds to the set of networks, in which a majority isotype is specified. list[[1]] or list$list_IGHG contains the networks, in data.frame format, with more IGG isotypes, list[[2]] or list$list_IGHA contains the networks, in data.frame format, with more IGA isotypes, list[[3]] or list$list_IGHM contains the networks, in data.frame format, with more IGM isotypes, list[[4]] or list$list_IGAG contains the networks, in data.frame format, with a tie in IGA and IGG isotypes and list[[5]] or list$list_other contains the networks, in data.frame format, with other isotypes apart from the aforementioned combinations.

See Also

ConvertStructure, CsvToDf
Examples

```r
## Not run:
SubReperatoiresByCells(list)

## End(Not run)
```

AbForests_SubReperatoiresByUniqueSeq

> Split single cell immune repertoire into sub-repertoires by isotype based on number of unique sequences

Description

SubReperatoiresByUniqueSeq separates the single cell immune repertoire into 5 sub-repertoires taking into account only unique sequences. The goal is to determine the majority isotype per each network in the immune repertoire. Therefore, each sub-repertoire is dominated by isotype IGG, IGA, IGM, other and if there is an equal number of IGA and IGG isotypes in a network, IGA-IGG category exists respectively. In particular, in case there exists a tie in the number of IGA and IGM, the network is considered to contain IGA as majority isotype, while the same number of IGG and IGM in the network categorize this network as containing IGG as majority isotype.

Usage

```r
AbForests_SubReperatoiresByUniqueSeq(
  list,
  opt,
  distance_mat,
  tie_flag,
  weight,
  random.seed,
  alg_opt,
  cdr3
)
```

Arguments

- **list**: a list of data.frames. Each data.frame represents a clone lineage and contains 2 columns, one that describes the antibody sequences and the other which type of information (isotype) is considered in the analysis. This list of data.frames has been generated by ConvertStructure function based on the initial data input or the output of CsvToDf and user's preferences.

- **opt**: a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.
**distance_mat**  
a custom integer distance matrix, or NULL for using the default distance matrix (calculated based on the levenshtein distance, which counts the number of mutations between sequences). Given the phylogenetic tree, a custom-made distance matrix can be produced by PlyloToMatrix function.

**tie_flag**  
a string, with options 'rand', 'full', 'close_to_germ', 'far_from_germ', 'close_path_to_germ', 'far_path_from_germ', 'most_expanded' and 'least_expanded' for removing edges when equal distance (tie) in distance matrix. 'rand' means random pruning in one of nodes, 'full' means keeping all nodes, close_to_germ means pruning of node(s) farthest from germline (based on number of intermediate nodes), far_from_germ means pruning of node(s) closest to germline (based on number of intermediate nodes), close_path_to_germ means pruning of node(s) farthest from germline (based on edge path length), far_path_from_germ means pruning of node(s) closest to germline (based on edge path length), most_expanded means pruning of node(s) with the lowest B cell count (clonal frequency) and least_expanded, which means pruning of node(s) with the highest B cell count (clonal frequency). In cases of subsequent ties, a random node is selected.

**weight**  
logical variable. When its value is FALSE, then the weights of outgoing edges from Germline node are set to 1. When its value is TRUE, the weights are set to the difference between the number of mutations among sequences in germline and connected nodes (value in the corresponding distance matrix) and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. In both cases, weights of remaining edges are extracted from the distance matrix.

**random.seed**  
a random seed, specified by the user, when random sampling of sequences happens in each of the cases described in tie_flag argument.

**alg_opt**  
a string denoting the version of the edge selection algorithm used in the construction of networks. "0" means the naive version and "1" the advanced one.

**cdr3**  
variable with values 0 if the user desires to select full length sequences (only when the input is a list of csv files), 1 for sequences in the CDR3 only (only when the input is a list of csv files) and NULL otherwise.

**Value**

list a nested list of 5 sub-lists of data.frames. Each sub-list corresponds to the set of networks, in which a majority isotype is specified. list[[1]] or list$list_IGHG contains the networks, in data.frame format, with more IGG isotypes, list[[2]] or list$list_IGHA contains the networks, in data.frame format, with more IGA isotypes, list[[3]] or list$list_IGHM contains the networks, in data.frame format, with more IGM isotypes, list[[4]] or list$list_IGAG contains the networks, in data.frame format, with a tie in IGA and IGG isotypes and list[[5]] or list$list_other contains the networks, in data.frame format, with other isotypes apart from the aforementioned combinations.

**See Also**

AntibodyForest, ConvertStructure, CsvToDf, PlyloToMatrix

**Examples**

## Not run:
**AbForests_UniqueAntibodyVariants**

Count the number of unique antibody variants per clonal lineage

**Description**

UniqueAntibodyVariants calculates the number of unique antibody sequences, as dictated by the different grouping sequences strategy, for each network in the immune repertoire.

**Usage**

```r
AbForests_UniqueAntibodyVariants(
  list,
  opt = "isotype",
  distance_mat = NULL,
  tie_flag = "close_to_germ",
  weight = TRUE,
  random.seed = 165,
  alg_opt = "naive",
  cdr3 = NULL
)
```

**Arguments**

- **list**
  
  a list of data.frames. Each data.frame represents a clone lineage and contains information on the antibody sequences and on the isotype/transcriptional cluster is considered in the analysis based the user's preferences.

- **opt**
  
  a string with options "isotype" and "cluster". The option "isotype" is utilized when the user desires to do an isotype analysis, while the selection of "cluster" denotes that an analysis based on transcriptome is requested.

- **distance_mat**
  
  a custom integer distance matrix, or NULL for using the default distance matrix (calculated based on the levenshtein distance, which counts the number of mutations between sequences). Given the phylogenetic tree, a custom-made distance matrix can be produced by PlyloToMatrix function.

- **tie_flag**
  
  a string, with options 'rand', 'full', 'close_to_germ', 'far_from_germ', 'close_path_to_germ', 'far_path_from_germ', 'most_expanded' and 'least_expanded' for removing edges when equal distance (tie) in distance matrix. 'rand' means random pruning in one of nodes, 'full' means keeping all nodes, close_to_germ means pruning of node(s) farthest from germline (based on number of intermediate nodes), far_from_germ means pruning of node(s) closest to germline (based on number of intermediate nodes), close_path_to_germ means pruning of node(s) farthest from germline (based on edge path length), far_path_from_germ means
pruning of node(s) closest to germline (based on edge path length), 'most_expanded' means pruning of node(s) with the lowest B cell count (clonal frequency) and least_expanded, which means pruning of node(s) with the highest B cell count (clonal frequency). In cases of subsequent ties, a random node is selected.

weight logical variable. When its value is FALSE, then the weights of outgoing edges from Germline node are set to 1. When its value is TRUE, the weights are set to the difference between the number of mutations among sequences in germline and connected nodes (value in the corresponding distance matrix) and the absolute value of the difference between the sequence lengths of germline and corresponding connected nodes. In both cases, weights of remaining edges are extracted from the distance matrix.

random.seed a random seed, specified by the user, when random sampling of sequences happens in each of the cases described in tie_flag argument.

alg_opt a string denoting the version of the edge selection algorithm used in the construction of networks. "0" means the naive version and "1" the advanced one.

cdr3 variable with values 0 if the user desires to select full length sequences (only when the input is a list of csv files), 1 for sequences in the CDR3 only (only when the input is a list of csv files) and NULL otherwise.

Value

uni_seq a vector, same size as list, which contains the number of unique antibody variants for each clonal lineage.

Examples

```r
## Not run:
UniqueAntibodyVariants(list, opt="cluster",
distance_mat=NULL, tie_flag=close_to_germ, weight=TRUE, random.seed=165, alg_opt="naive", cdr3=NULL)
## End(Not run)
```

automate_GEX

GEX processing wrapper in Platypus V2

Description

Automates the transcriptional analysis of the gene expression libraries from cellranger. This function will integrate multiple samples

Usage

```r
automate_GEX(
    GEX.outs.directory.list,
    GEX.list,
    integration.method,
    VDJ.gene.filter,
)```
Arguments

GEX.outs.directory.list
The path to the output of cellranger vdj runs. Multiple repertoires to be integrated together should be supplied as a character vector in the first element of a list. For example, if two separate VDJ repertoires should be integrated together (e.g. on the same tSNE plot), GEX.outs.directory.list[[1]] <- c("my.VDJ1.path/outs/","my.VDJ2.path/outs/") should be stored as input. If these repertoires should be analyzed separately, >GEX.outs.directory.list[[1]] <- "my.VDJ1.path/outs/" >GEX.outs.directory.list[[2]] <- "my.VDJ2.path/outs/") should be supplied. This can be left blank if supplying the clonotypes and all_contig files directly as input. Multiple analyses can be stored

GEX.list
List containing the output from Seurat Read10x. This must be supplied if GEX.out.directory is not provided.

integration.method
String specifying which data normalization and integration pipeline should be used. Default is "scale.data", which corresponds to the ScaleData function internal to harmony package. "sct" specifies SCTransform from the Seurat package. "harmony" should be specified to perform harmony integration. This method requires the harmony package from bioconductor.

VDJ.gene.filter
Logical indicating if variable genes from the b cell receptor and t cell receptor should be removed from the analysis. True is highly recommended to avoid clonal families clustering together.

mito.filter
Numeric specifying which percent of genes are allowed to be composed of mitochondrial genes. This value may require visual inspection and can be specific to each sequencing experiment. Users can visualize the percentage of genes corresponding to mitochondrial genes using the function "investigate_mitochondrial_genes".

norm.scale.factor
Scaling factor for the standard Seurat pipeline. Default is set to 10000 as reported in Seurat documentation.

n.feature.rna
Numeric that specifies which cells should be filtered out due to low number of detected genes. Default is set to 0. Seurat standard pipeline uses 2000.

n.count.rna.min
Numeric that specifies which cells should be filtered out due to low RNA count. Default is set to 0. Seurat standard pipeline without VDJ information uses 200.
n.count.rna.max
Numeric that specifies which cells should be filtered out due to high RNA count. Default is set to infinity. Seurat standard pipeline without VDJ information uses 2500.

n.variable.features
Numeric specifying the number of variable features. Default set to 2000 as specified in Seurat standard pipeline.

cluster.resolution
Numeric specifying the resolution that will be supplied to Seurat’s FindClusters function. Default is set to 0.5. Increasing this number will increase the number of distinct Seurat clusters. Suggested to examine multiple parameters to ensure gene signatures differentiating clusters remains constant.

neighbor.dim
Numeric vector specifying which dimensions should be supplied in the FindNeighbors function from Seurat. Default input is ‘1:10’.

mds.dim
Numeric vector specifying which dimensions should be supplied into dimensional reduction techniques in Seurat and Harmony. Default input is ‘1:10’.

groups
Integer specifying the groups of the different samples. This is needed if there are multiple biological replicates for a given condition sequenced and aligned through cellranger separately.

Value
Returns a processed Seurat object containing transcriptional information from all samples which can be supplied to the VDJ_GEX_integrate function.

Examples
```r
## Not run:
automate_GEX(out_directory=fullRepertoire.output,
rep.size=3*length(unlist(fullRepertoire.output[[1]])),
distribution="identical",
with.germline="FALSE")
## End(Not run)
```

---

Bcell_sequences_example_tree

*Example csv file 1*

Description
Example csv file 1

Usage
Bcell_sequences_example_tree
**Bcell_tree_2**

**Format**

An object of class `data.frame` with 170 rows and 1 columns.

**References**

R package Platypus: https://doi.org/10.1093/nargab/lqab023

---

**Example csv file 2**

**Description**

Example csv file 2

**Usage**

`Bcell_tree_2`

---

**call_MIXCR**

**Calls MiXCR VDJ object of Platypus V2**

**Description**

Extracts information on the VDJRegion level using MiXCR. This function assumes the user can run an executable instance of MiXCR and is eligible to use MiXCR as determined by license agreements. The VDJRegion corresponds to the recombined heavy and light chain loci starting from framework region 1 (FR1) and extending to framework region 4 (FR4). This can be useful for extracting full-length sequences ready to clone and further calculating somatic hypermutation occurrences.

**Usage**

`call_MIXCR(VDJ.per.clone, mixcr.directory, species)`
Arguments

VDJ.per.clone The output from the VDJ_per_clone function. This object should have information regarding the contigs and clonotype_ids for each cell.
mixcr.directory The directory containing an executable version of MiXCR. This must be downloaded separately and is under a separate license.
species Either "mmu" for mouse or "hsa" for human. These use the default germline genes for both species contained in MIXCR.

Value

Returns a nested list containing VDJRegion information as determined by MIXCR. The outer list corresponds to the individual repertoires in the same structure as the input VDJ.per.clone. The inner list corresponds to each clonal family, as determined by either the VDJ_clonotype function or the default nucleotide clonotyping produced by cellranger. Each element in the inner list corresponds to a dataframe containing repertoire information such as isotype, CDR sequences, mean number of UMIs. This output can be supplied to further package functions such as VDJ_extract_sequences and VDJ_GEX_integrate.

See Also

VDJ_extract_sequences

Examples

```r
## Not run:
call_MIXCR(VDJ.per.clone = VDJ.per.clone.output
, mixcr.directory = "~/Downloads/mixcr-3.0.12/mixcr", species = "mmu")
## End(Not run)
```

class.switch.prob.hum The probability matrix of class switching for human b cells. The row names of the matrix are the isotypes the cell is switching from, the column names are the isotypes the cell is switching to. All B cells start from IGHM, and switch to one of the other isotypes or remain the same.

Description

class.switch.prob.hum The probability matrix of class switching for human b cells. The row names of the matrix are the isotypes the cell is switching from, the column names are the isotypes the cell is switching to. All B cells start from IGHM, and switch to one of the other isotypes or remain the same.

Usage

data("class.switch.prob.hum")
class_switch_prob_mus

**Format**

A 8*8 matrix. The row and column names are "IGHM", "IGHD", "IGHG1", "IGHG2", "IGHG3", "IGHG4", "IGHE", "IGHA". The probability for a cell to switch from "IGHM" to "IGHD" is the value at class_switch_prob_hum[1,2].

class_switch_prob_mus

The probability matrix of class switching for mouse B cells. The row names of the matrix are the isotypes the cell is switching from, the column names are the isotypes the cell is switching to. All B cells start from IGHM, and switch to one of the other isotypes or remain the same.

**Description**

class_switch_prob_mus The probability matrix of class switching for mouse B cells. The row names of the matrix are the isotypes the cell is switching from, the column names are the isotypes the cell is switching to. All B cells start from IGHM, and switch to one of the other isotypes or remain the same.

**Usage**

data("class_switch_prob_mus")

**Format**

A 9*9 matrix. The row and column names are "IGHM", "IGHD", "IGHG1", "IGHG2A", "IGHG2B", "IGHG2C", "IGHG3", "IGHE", "IGHA". The probability for a cell to switch from "IGHM" to "IGHD" is the value at class_switch_prob_mus[1,2].

clonofreq

Plot clonal frequency barplot of the output simulated data

**Description**

Plot the top abundant clonal frequencies in a barplot with ggplot2

**Usage**

clonofreq(clonotypes, top.n, y.limit)

**Arguments**

- **clonotypes**: The clonotypes dataframe, which is the second element in the simulation output list.
- **top.n**: The top n abundant clones to be shown in the plot. If missing, all clones will be shown.
- **y.limit**: The upper limit for y axis in the plot.
clonofreq.isotype.data

*Get information about the clonotype counts grouped by isotype.*

**Description**

Return

**Usage**

clonofreq.isotype.data(all.contig.annotations, top.n)

**Arguments**

all.contig.annotations

The output dataframe all_contig_annotation from function simulate.repertoire.

top.n

The top n abundant clones to be shown in the plot. If missing, all clones will be shown.

**Value**

dataframes containing the top n abundant clonotypes and their frequency and isotype information for further processing.

clonofreq.isotype.plot

*Get information about the clonotype counts grouped by isotype.*

**Description**

Plot a stacked barplot for clonotype counts grouped by isotype.

**Usage**

clonofreq.isotype.plot(all.contig.annotations, top.n, y.limit, colors)
Arguments

all.contig.annotations
The output dataframe all_contig_annotation from function simulate.repertoire.

top.n
The top n abundant clones to be shown in the plot. If missing, all clones will be shown.

y.limit
The upper limit for y axis in the plot.

colors
A named character vector of colors, the names are the isotypes. If missing, the default has 11 colors corresponding to the default isotype names.

Value

a stacked barplot for clonotype counts grouped by isotype

---

clonofreq.trans.data  Get information about the clonotype counts grouped by transcriptome state(cell type).

Description

Dataframe with clonotype counts grouped by transcriptome state(cell type).

Usage

clonofreq.trans.data(all.contig.annotations, history, trans.names, top.n)

Arguments

all.contig.annotations
The output dataframe all_contig_annotation from function simulate.repertoire.

history
The dataframe history from simulate output.

trans.names
The names of cell types which are used in transcriptome.switch.prob argument in the simulation.

top.n
The top n abundant clones to be shown in the plot. If missing, all clones will be shown.

Value

a dataframe with clonotype counts grouped by transcriptome state(cell type).
### clonofreq.trans.plot

*Get information about the clonotype counts grouped by transcriptome state (cell type).*

#### Description

Plot a stacked barplot for clonotype counts grouped by transcriptome state (cell type).

#### Usage

```r
clonofreq.trans.plot(
  all.contig.annotations,
  history,
  trans.names,
  top.n,
  y.limit,
  colors
)
```

#### Arguments

- `all.contig.annotations`: The output dataframe `all_contig_annotation` from function `simulate.repertoire`.
- `history`: The dataframe `history` from `simulate` output.
- `trans.names`: The names of cell types which are used in `transcriptome.switch.prob` argument in the simulation.
- `top.n`: The top n abundant clones to be shown in the plot. If missing, all clones will be shown.
- `y.limit`: The upper limit for y axis in the plot.
- `colors`: A named character vector of colors, the names are the isotypes. If missing, the default has 11 colors corresponding to the default isotype names.

#### Value

A stacked barplot for clonotype counts grouped by transcriptome state (cell type).

### cluster.id.igraph

*Get clone network igraphs colored by seurat cluster id.*

#### Description

Get clone network igraphs colored by seurat cluster id.
colors

Usage

cluster.id.igraph(meta.data, history, igraph.index, empty.node)

Arguments

- **meta.data**: the meta.data dataframe from the Seurat object of the simulation. The object should be pre-processed and has cluster ids in the meta.data.
- **history**: The dataframe 'history' from the simulation output.
- **igraph.index**: The list 'igraph.index' from the simulation output.
- **empty.node**: If TRUE, there will be empty node in igraph. if FALSE, the empty node will be deleted.

Value

a list of clone network igraphs colored by seurat cluster id

colors

colors A vector of characters specifying colors used in igraph phylogenetic tree. Default colors: "#66C2A5", "#FC8D62", "#8DA0CB", "#E78AC3", "#A6D854"

Description

colors A vector of characters specifying colors used in igraph phylogenetic tree. Default colors: "#66C2A5", "#FC8D62", "#8DA0CB", "#E78AC3", "#A6D854"

Usage

data("colors")

Format

a character vector
Echidna_simulate_repertoire

Simulate immune repertoire and transcriptome data

Description

Simulate repertoire and transcriptome matrix, with igraph tree plot for each clone showing the evolution process. The node in the tree plot are colored with transcriptome state and isotype.

Usage

Echidna_simulate_repertoire(
  initial.size.of.repertoire,
  species,
  cell.type,
  cd4.proportion,
  duration.of.evolution,
  complete.duration,
  vdj.productive,
  vdj.model,
  vdj.insertion.mean,
  vdj.insertion.stdv,
  vdj.branch.prob,
  clonal.selection,
  cell.division.prob,
  sequence.selection.prob,
  special.v.gene,
  class.switch.prob,
  class.switch.selection.dependent,
  class.switch.independent,
  SHM.method,
  SHM.nuc.prob,
  SHM.isotype.dependent,
  SHM.phenotype.dependent,
  max.cell.number,
  max.clonotype.number,
  death.rate,
  igraph.on,
  transcriptome.on,
  transcriptome.switch.independent,
  transcriptome.switch.prob,
  transcriptome.switch.isotype.dependent,
  transcriptome.switch.SHM.dependent,
  transcriptome.switch.selection.dependent,
  transcriptome.states,
  transcriptome.noise,
  seq.name)
Arguments

**initial.size.of.repertoire**
The initial number of existing cells when the evolution starts. Default is 10.

**species**
The species of the simulated repertoire, can be "mus" for mouse or "hum" for human. Default is "mus".

**cell.type**
The cell type for the simulation. "B" or "T"

**cd4.proportion**
A number between 0 and 1 specifying the proportion of Cd4+ T cells, when cell.type is "T" and transcriptome states data is default. Default is 1, all the cells are Cd4. When user specify transcriptome data for T cells, mixture of CD4+ and CD8+ T cells are not applicable.

**duration.of.evolution**
The maxim time steps for simulation.

**complete.duration**
TRUE or FALSE. Default is TRUE. If TRUE, after cell number or clone number reaches the upper limit, the evolution (class switch, mutation, transcriptional state switch) will continue until the duration.of.evolution is complete. If FALSE, the evolution will stop when either cell number or clone number reaches the limit.

**vdj.productive**
"random": the sequence will be generated from random VDJ recombination, there might be a proportion of unproductive sequences. These VDJ genes were taken from IMGT. When more than one allele was present for a given gene, the first was used. "naive": the VDJ sequence be sampled from a pool of productive sequences obtained by filtering randomly simulated sequences with MIXCR. "vae": the VDJ sequence be sampled from a pool of productive sequences obtained by filtering sequences generated from vae models with MIXCR.

**vdj.model**
Specifies the model used to simulate V-D-J recombination. Can be either "naive" or "data". "naive" is chain independent and does not differentiate between different species. To rely on the default "experimental" options, this should be "data" and the parameter vdj.insertion.mean should be "default". This will allow for different mean additions for either the VD and JD junctions and will differ depending on species.

**vdj.insertion.mean**
Integer value describing the mean number of nucleotides to be inserted during simulated V-D-J recombination events. If "default" is entered, the mean will be normally distributed.

**vdj.insertion.stdv**
Integer value describing the standard deviation corresponding to insertions of V-D-J recombination. No "default" parameter currently supported but will be updated with future experimental data. This should be a number if using a custom distribution for V-D-J recombination events, but can be "default" if using the "naive" vdj.model or the "data", with vdj.insertion.mean set to "default".

**vdj.branch.prob**
Probability of new VDJ recombination event in each time step. When new VDJ recombination happen, a new cell with a new sequence will be generated. Default is 0.2.
clonal.selection
TRUE or FALSE. If TRUE, cells in clones with higher frequency have their division probability proportional to the clonal frequency. If FALSE, clones with higher frequency will have lower probability to expand.

cell.division.prob
Probability of cells to be duplicated in each time step. Default is 0.1. If uneven probability for different clones is needed, the input should be a vector of 2 numeric items, with the first item being the lower bound, the second item being the upper bound of the division rate. The most abundant clone will get the highest division rate, and division rate of other clones will follow arithmetic progression and keep decreasing until the last abundant clone with the lower limit of division rate. If input 3 values, the third value will be the division rate for cells with selected sequences. If a fourth number is given, the division probability of selected sequence will be sampled between the third number and the fourth number.

sequence.selection.prob
Probability of each unique sequence to be selected as expanding sequence. Expanding sequences can have their division rate specified in the third element of cell.division.prob.

special.v.gene
If TRUE, simulation will apply special sequence.selection.prob for heavy and light chain v gene combination specified in dataframe "special_v".

class.switch.prob
Probability matrix of class switching for b cells. The row names of the matrix are the isotypes the cell is switching from, the column names are the isotypes the cell is switching to. All B cells start from IGHM, and switch to one of the other isotypes or remain the same. Default values are in the attaching matrix "class_switch_prob_hum" and "class_switch_prob_mus". The order of isotype in rows and columns should be the same.

class.switch.selection.dependent
If TRUE, class switching will happen when the cell is selected, if the cell has IgM or IgD isotype.

class.switch.independent
If TRUE, class switching will happen randomly at each time step for all cells. If FALSE, random class switching will be switched off.

SHM.method
The mode of SHM speciation events. Options are either: "poisson", "data", "motif", "wrc", and "all". Specifying either "poisson" or "naive" will result in mutations that can occur anywhere in the heavy chain region, with each nucleotide having an equal probability for a mutation event. Specifying "data" focuses mutation events during SHM in the CDR regions (based on IMGT), and there will be an increased probability for transitions (and decreased probability for transversions). Specifying "motif" will cause neighbor dependent mutations based on a mutational matrix from high throughput sequencing data sets (Yaari et al., Frontiers in Immunology, 2013). "wrc" allows for only the WRC mutational hotspots to be included (where W equals A or T and R equals A or G). Specifying "all" will use all four types of mutations during SHM branching events, where the weights for each can be specified in the "SHM.nuc.prob" parameter.

SHM.nuc.prob
Specifies the rate at which nucleotides change during speciation (SHM) events. This parameter depends on the type of mutation specified by SHM.method. For
both "poisson" and "data", the input value determines the probability for each site to mutate (the whole sequence for "poisson" and the CDRs for "data"). For either "motif" or "wrc", the number of mutations per speciation event should be specified. Note that these are not probabilities, but the number of mutations that can occur (if the mutation is present in the sequence). If "all" is specified, the input should be a vector where the first element controls the poisson style mutations, second controls the "data", third controls the "motif" and fourth controls the "wrc".

**SHM.isotype.dependent**

If TRUE, somatic hypermutation of certain isotype will happen based on probability specified in dataframe "iso_SHM_prob".

**SHM.phenotype.dependent**

If TRUE, somatic hypermutation of certain phenotype will happen based on probability specified in dataframe "pheno_SHM_prob".

**max.cell.number**

Integer value describing maximum number of cells allowed. Default is 1500.

**max.clonotype.number**

Integer value describing maximum number of clones allowed. Cell derived from the same mother cell belong to same clone.

**death.rate**

Probability of cell death happen to each cell in each time step.

**igraph.on**

If TRUE, mutational network for every B cell clone will be in the output. If False, the igraphs will not be included.

**transcriptome.on**

If TRUE, the simulation will include transcriptome data. If FALSE, only vdj sequence will be simulated.

**transcriptome.switch.independent**

TRUE or FALSE value describing whether transcriptome state is allowed to switch independently, not dependent on class switching or somatic hypermutation. If TRUE, transcriptome.switch.prob should be specified to control the probability of transcriptome state switching.

**transcriptome.switch.prob**

Probability of transcriptome state switching independently. Default values are in the attaching matrix "trans_switch_prob_b" and "trans_switch_prob_t". The order of cell type in rows and columns should be the same, and the order of the cell type in the matrix should match cell type names in transcriptome.states.

**transcriptome.switch.isotype.dependent**

TRUE or FALSE value describing whether transcriptome state of a cell is allowed to switch depending on isotype switching. If TRUE, transcriptome state will switch once class switching happens.

**transcriptome.switch.SHM.dependent**

TRUE or FALSE value describing whether transcriptome state of a cell is allowed to switch depending on somatic hypermutation. If TRUE, transcriptome state will switch once somatic hypermutation happens.

**transcriptome.switch.selection.dependent**

If TRUE, selected cells will undergo transcriptome state switching if their transcriptome state is 1.
Echidna_vae_generate

Simulate B or T cell receptor sequences by variational autoencodes (VAEs) trained with experimental data.

**Description**

Simulate B or T cell receptor sequences by variational autoencodes (VAEs) trained with experimental data.

**Usage**

```r
echidna_vae_generate(
  sequence,
  n.train,
  n.sample,
  batch.size,
  latent.dim,
  intermediate.dim,
  epochs,
  epsilon.std,
  null.threshold
)
```
get.avr.mut.data

Arguments

- **sequence**: a vector of sequence the model to be trained on
- **n.train**: number of sequence to be used in training set, the rest will be in testing set
- **n.sample**: number of new sequence to generate from VAE model
- **batch.size**: set to larger to save time, set to smaller to same computing power
- **latent.dim**: parameter used in VAE model
- **intermediate.dim**: parameter used in VAE model
- **epochs**: parameter used in VAE model
- **epsilon.std**: parameter used in VAE model
- **null.threshold**: threshold of predicted value to be considered as an existing base, default is 0.05. When generated sequence is too short, lower this threshold.

Value

A simulated VDJ repertoire on the basis of the input experimental repertoire

get.avr.mut.data

Get information about somatic hypermutation in the simulation. This function return a barplot showing the average mutation.

Description

Get information about somatic hypermutation in the simulation. This function return a barplot showing the average mutation.

Usage

get.avr.mut.data(igraph.index.attr, history, clonotype.select, level)

Arguments

- **igraph.index.attr**: A list "igraph.index.attr" from the simulation output.
- **history**: A dataframe "history" from the simulation output.
- **clonotype.select**: The selected clonotype index, can be the output of the function "select.top.clone.clone".
- **level**: Can be "clone" or "cell". If "clone", the function will return average mutation on unique variant level. Otherwise it will return on cell level.

Value

A bar plot showing the average mutation on clone or cell level.
**get.avr.mut.plot**  
*Get information about somatic hypermutation in the simulation. This function return a barplot showing the average mutation.*

**Description**

Get information about somatic hypermutation in the simulation. This function return a barplot showing the average mutation.

**Usage**

```r
get.avr.mut.plot(igraph.index.attr, history, clonotype.select, level, y.limit)
```

**Arguments**

- `igraph.index.attr`: A list "igraph.index.attr" from the simulation output.
- `history`: A dataframe "history" from the simulation output.
- `clonotype.select`: The selected clonotype index, can be the output of the function "select.top.clone".
- `level`: Can be "node" or "cell". If "node", the function will return average mutation on unique variant level. Otherwise it will return on cell level.
- `y.limit`: The upper limit for y axis in the plot.

**Value**

a barplot showing the average mutation per node (same heavy and light chain set) or per cell.

---

**get.barplot.errorbar**  
*Return a barplot of mean and standard error bar of certain value of each clone.*

**Description**

Return a barplot of mean and standard error bar of certain value of each clone.

**Usage**

```r
get.barplot.errorbar(data, y.lab, y.limit)
```

**Arguments**

- `data`: A dataframe. Columns are different simulations, rows are the top clones. The first row is the top abundant clone.
- `y.lab`: A string specifies the y lable name of the barplot.
- `y.limit`: The upper limit for y axis in the plot.
get.elbow

Value

a barplot of mean and standard error bar of certain value of each clone.

get.elbow  Get the seurat object from simulated transcriptome output.

Description

Get the seurat object from simulated transcriptome output.

Usage

get.elbow(data)

Arguments

data  The output "transcriptome" dataframe from simulation output.

Value

the seurat object from simulated transcriptome output.

get.n.node.data  Get the number of unique variants in each clone in a vector. The output is the vector representing the numbers of unique variants.

Description

Get the number of unique variants in each clone in a vector. The output is the vector representing the numbers of unique variants.

Usage

get.n.node.data(data, clonotype.select)

Arguments

data  The output "igraph.index.attr" list from simulation output.
clonotype.select  The index of the clones to be shown. If missing, all clones will be included.

Value

the number of unique variants in each clone in a vector. The output is the vector representing the numbers of unique variants.
get.n.node.plot

Get the number of unique variants in each clone in a vector and the barplot. The first item in the output is the vector representing the numbers of unique variants, the second item is the barplot.

Description

Get the number of unique variants in each clone in a vector and the barplot. The first item in the output is the vector representing the numbers of unique variants, the second item is the barplot.

Usage

get.n.node.plot(igraph.index.attr, clonotype.select, y.limit)

Arguments

igraph.index.attr
The output "igraph.index.attr" list from simulation output.
clonotype.select
The index of the clones to be shown. If missing, all clones will be included.
y.limit
The upper limit for y axis in the plot.

Value

the number of unique variants in each clone in a vector and the barplot. The first item in the output is the vector representing the numbers of unique variants, the second item is the barplot.

get.seq.distance

Computing sequence distance according to the number of unmatched bases.

Description

Computing sequence distance according to the number of unmatched bases.

Usage

get.seq.distance(germline, sequence)

Arguments

germline
A string representing the germline sequence.
sequence
A string of the sequence to be compared, which has the same length as germline.

Value

the number of unmatched bases in 2 sequences.
get.uimap

Further process the seurat object from simulated transcriptome output and make UMAP ready for plotting.

Description
Further process the seurat object from simulated transcriptome output and make UMAP ready for plotting.

Usage
get.uimap(gex, d, reso)

Arguments
- gex: output from get.elbow function.
- d: dims argument of in Seurat::FindNeighbors() and Seurat::RunUMAP
- reso: resolution argument in Seurat::FindClusters()

Value
Further processed seurat object from simulated transcriptome output with UMAP ready for plotting.

get.vgu.matrix

Get paired v gene heavy chain and light chain matrix on clonotype level. A v gene usage heatmap can be obtained by pheatmap::pheatmap(vgu_matrix, show_colnames=T, main = "V Gene Usage"), where the vgu_matrix is the output of this function.

Description
Get paired v gene heavy chain and light chain matrix on clonotype level. A v gene usage heatmap can be obtained by pheatmap::pheatmap(vgu_matrix, show_colnames=T, main = "V Gene Usage"), where the vgu_matrix is the output of this function.

Usage
get.vgu.matrix(all.contig.annotations, level)

Arguments
- all.contig.annotations: The dataframe "all_contig_1 Annotation" from simulation output.
- level: Can be "clone" or "cell". If "clone", the function will return paired v gene usage matrix on clonotype level. Otherwise it will return on cell level.
Value

A paired `v` gene heavy chain and light chain matrix on clonotype level.

---

**GEX_clonotype**

*Platypus V2 GEX and VDJ integration for clonotypes*

Description

Platypus V2: Integrates VDJ and gene expression libraries by providing cluster membership `seq_per_vdj` object and the index of the cell in the Seurat RNA-seq object.

Usage

`GEX_clonotype(GEX.object, VDJ.per.clone)`

Arguments

- **GEX.object**: A single seurat object from `automate_GEX` function. This will likely be supplied as `automate_GEX.output[[1]]`.
- **VDJ.per.clone**: Output from the `VDJ_per_clone` function. Each element in the list should be found in the output from the `automate_GEX` function.

Value

Returns a dataframe containing repertoire information, such as isotype, CDR sequences, mean number of UMIs. This output can be supplied to further packages `VDJ_extract_sequences` and `VDJ_GEX_integrate`.

Examples

```r
## Not run:
GEX_clonotype(GEX.object=automate.GEX.output[[1]], VDJ.per.clone=vdj.per.clone.output)

## End(Not run)```
GEX_cluster_genes Differentially expressed genes between clusters or data subsets

Description

For more flexibility consider GEX_DEgenes(). Extracts the differentially expressed genes between two samples. This function uses the FindMarkers function from the Seurat package. Further parameter control can be accomplished by calling the function directly on the output of automate_GEX or VDJ_GEX_matrix.

Usage

GEX_cluster_genes(GEX, min.pct, filter, base, platypus.version)

Arguments

- **GEX**: Output Seurat object of either automate_GEX for platypus.version v2 or of VDJ_GEX_matrix for platypus.version v3 (usually VDJ_GEX_matrix.output[[2]])
- **min.pct**: The minimum percentage of cells expressing a gene in either of the two groups to be compared. Default is 0.25
- **filter**: Character vector of initials of the genes to be filtered. Default is c("MT-", "RPL", "RPS"), which filters mitochondrial and ribosomal genes.
- **base**: The base with respect to which logarithms are computed. Default: 2
- **platypus.version**: is set automatically

Value

Returns a dataframe containing the output from the FindMarkers function, which contains information regarding the genes that are differentially regulated, statistics (p value and log fold change), and the percent of cells expressing the particular gene. Each element in the list corresponds to the clusters in numerical order. For example, the first element in the list output[[1]] corresponds to the genes differentially expressed in cluster 0 in GEX

Examples

# Platypus version v2
# GEX_cluster_genes(GEX = automate_GEX_output[[i]], min.pct = .25, filter = c("MT-", "RPL", "RPS"))

# Platypus version v3
GEX_cluster_genes(GEX = subset(Platypus::small_vgm[[2]], seurat_clusters %in% c(0,1)), min.pct = .25, filter = c("MT-", "RPL", "RPS"))
**GEX_cluster_genes_heatmap**

*Heatmap of cluster defining genes*

**Description**

Produces a heatmap displaying the expression of the top genes that define each cluster in the Seurat object. The output heatmap is derived from DoHeatmap from Seurat and thereby can be edited using typical ggplot interactions. The number of genes per cluster and the number of cells to display can be specified by the user. Either the log fold change or the p value can be used to select the top n genes.

**Usage**

```r
GEX_cluster_genes_heatmap(  
  GEX,  
  GEX_cluster_genes.output,  
  n.genes.per.cluster,  
  metric,  
  max.cell,  
  group.colors,  
  slot,  
  platypus.version  
)
```

**Arguments**

- **GEX**
  Output Seurat object of either automate_GEX for platypus.version v2 or of VDJ_GEX_matrix for platypus.version v3 (usually VDJ_GEX_matrix.output[[2]])

- **GEX_cluster_genes.output**
  The output from the GEX_cluster_genes function - this should be a list with each list element corresponding to the genes, p values, logFC, pct expression for the genes differentially regulated for each cluster.

- **n.genes.per.cluster**
  An integer value determining how many genes per cluster to display in the output heatmap. This number should be adjusted based on the number of clusters. Too many genes per cluster and clusters may cause a problem with the heatmap function in Seurat.

- **metric**
  The metric that dictates which are the top n genes returned. Possible options are "p.value" (default), "avg_logFC", "top_logFC", "bottom_logFC". "top_logFC" returns the top expressed genes for each cluster, whereas "bottom_logFC" returns the least expressed genes per cluster both by log fold change.

- **max.cell**
  The max number of cells to display in the heatmap for each cluster, which corresponds to the number of columns. Default is set to 100 cells per cluster.

- **group.colors**
  Optional character vector. Array of colors with the same length as GEX_cluster_genes.output to color bars above the heatmap. Defaults to rainbow palette.
GEX_cluster_membership

slot
Seurat object slot from which to plot gene expression data.

platypus.version
is set automatically

Value
Returns a heatmap from the function DoHeatmap from the package Seurat, which is a ggplot object that can be modified or plotted. The number of genes is determined by the n.genes parameter and the number of cells per cluster is determined by the max.cell argument. This function gives a visual description of the top genes differentially expressed in each cluster.

Examples

```r
## Not run:
# For Platypus version 2
cluster_defining_gene_heatmap <- GEX_cluster_genes_heatmap(GEX = automate_GEX_output[[i]],
                                 GEX_cluster_genes.output=GEX_cluster_genes_output,
                                 n.genes.per.cluster=5,metric="p.value",max.cell=5)

# For Platypus version 3
cluster_defining_gene_heatmap <- GEX_cluster_genes_heatmap(GEX = VDJ_GEX_matrix.output[[2]],
                                 GEX_cluster_genes.output=GEX_cluster_genes_output,
                                 n.genes.per.cluster=5,metric="p.value",max.cell=5)

## End(Not run)
```

---

GEX_cluster_membership

_Cluster membership plots by sample_

Description
Plots the cluster membership for each of the distinct samples in the Seurat object from the automate_GEX function. The distinct samples are determined by "sample_id" field in the Seurat object.

Usage

GEX_cluster_membership(GEX, by.group, platypus.version)

Arguments

GEX
Output Seurat object containing gene expression data from automate_GEX (platypus.version = "v2") or VDJ_GEX_matrix (platypus.version = "v3", usually VDJ_GEX_matrix.output[[2]]) that contained at least two distinct biological samples. The different biological samples correspond to integer values (v2) or factor values (v3) in the order of the working directories initially supplied to the automate_GEX function.
by.group

Logical indicating whether to look at the cluster distribution per group (using the group_id column). Default is set to FALSE.

platypus.version

Version of platypus to use. Defaults to "v2". If an output of the GEX_automate function is supplied, set to "v2". If an output of the VDJ_GEX_matrix function is supplied set to "v3"

Value

Returns a ggplot object in which the values on the x axis correspond to each cluster found in the Seurat object. The y axis corresponds to the percentage of cells found in each cluster. The bar and color corresponds to the distinct sample_id.

Examples

#Platypus v2
#GEX_cluster_membership(GEX=automate_GEX_out[[2]], platypus.version = "v2")
#Platypus v3
GEX_cluster_membership(GEX= Platypus::small_vgm[[2]], platypus.version = "v3")

GEX_coexpression_coefficient

Description

Returns either a plot or numeric data of coexpression levels of selected genes. Coexpression % is calculated as the quotient of double positive cells (counts > 0) and the sum of total cells positive for either genes.

Usage

GEX_coexpression_coefficient(GEX, genes, subsample.n, plot.dotmap)

Arguments

GEX

GEX seurat object generated with VDJ_GEX_matrix (VDJ_GEX_matrix.output[[2]])

genes

Character vector. At least 2 genes present in rownames(GEX). Use "all" to include all genes. The number of comparisons to make is the length(genes)! (factorial). More than 100 genes are not recommended.

subsample.n

Integer. Number of cells to subsample. If set to 100, 100 cells will be randomly sampled for the calculation

plot.dotmap

Boolean. Whether to return a plot

Value

Returns a dataframe if plot.dotmap == F or a ggplot if plot.dotmap == T detailing the coexpression levels of selected genes within the given cell population
Examples

To return a dataframe with coefficients
```
#GEX_coexpression_coefficient(GEX = VDJ_GEX_matrix.output[[2]]
#, genes = c("CD19", "EBF1","SDC1"), subsample.n = "none", plot.dotmap = FALSE)
```

To return a dotplot detailing coexpression and overall expression
```
GEX_coexpression_coefficient(GEX = Platypus::small_vgm[[2]]
, genes = c("CD19", "CD83"), subsample.n = "none", plot.dotmap = FALSE)
```

GEX_DEgenes

**Wrapper for differential gene expression analysis and plotting**

Description

Extracts the differentially expressed genes between two groups of cells. These groups are defined as cells having either of two entries (group1, group2) in the grouping.column of the input Seurat object metadata. This function uses the FindMarkers function from the Seurat package.

Usage

```
GEX_DEgenes(
  GEX,
  FindMarkers.out,
  grouping.column,
  group1,
  group2,
  min.pct,
  filter,
  return.plot,
  logFC,
  color.p.threshold,
  color.log.threshold,
  color.by.threshold,
  up.genes,
  down.genes,
  base,
  label.n.top.genes,
  genes.to.label,
  platypus.version,
  size.top.colorbar
)
```

Arguments

- **GEX**: Output Seurat object from automate_GEX or VDJ_GEX_matrix_function (VDJ_GEX_matrix.output[[2]]) function that contained at least two distinct biological groups.
FindMarkers.out

OPTIONAL: the output of the FindMarkers function. This skips the DEG calculation step and outputs desired plots. All plotting parameters function as normal. Grouping parameters and min.pct are ignored.

grouping.column

Character. A column name of GEX@meta.data. In this column, group1 and group2 should be found. Defaults to "sample_id". Could also be set to "seurat_clusters" to generate DEGs between cells of 2 chosen clusters.

group1
either character or integer specifying the first group of cells that should be compared. (e.g. "s1" if sample_id is used as grouping.column)

group2
either character or integer specifying the first group of cells that should be compared. (e.g. "s2" if sample_id is used as grouping.column)

min.pct

The minimum percentage of cells expressing a gene in either of the two groups to be compared.

filter

Character vector of initials of the genes to be filtered. Default is c("MT-", "RPL", "RPS"), which filters mitochondrial and ribosomal genes.

return.plot

Character specifying if a "heatmap", "heatmap" or a "volcano" or "none" is to be returned. If not "none" then @return is a list where the first element is a dataframe and the second a plot (see @return). Defaults to none

logFC

Logical specifying whether the genes will be displayed based on logFC (TRUE) or pvalue (FALSE).

color.p.threshold

numeric specifying the adjusted p-value threshold for geom_points to be colored. Default is set to 0.01.

color.log.threshold

numeric specifying the absolute logFC threshold for geom_points to be colored. Default is set to 0.25.

color.by.threshold

Boolean. Set to TRUE to color by color.p.threshold and color.log.threshold. Set to FALSE for a continuous color scale by fold change.

up.genes

FOR HEATMAP Integer specifying the number of upregulated genes to be shown.

down.genes

FOR HEATMAP Integer specifying the number of downregulated genes to be shown.

base

The base with respect to which logarithms are computed. Default: 2

label.n.top.genes

FOR VOLCANO Integer. How many top genes to label either by Fold change (if logFC == TRUE) or by p.value (if logFC == FALSE). More than 50 are not recommended. Also works in conjunction with genes.to.label

genes.to.label

FOR VOLCANO Character vector of genes to label irregardless of their p value.

platypus.version

Function works with V2 and V3, no need to set this parameter

size.top.colorbar

Integer. Size of the top colorbar for heatmap plot.
GEX_DEgenes_persample  Platypus V2 Differentially expressed genes

Value

Returns a dataframe containing the output from the FindMarkers function, which contains information regarding the genes that are differentially regulated, statistics (p value and log fold change), and the percent of cells expressing the particular gene for both groups.

Examples

#Basic run between two samples
GEX_DEgenes(GEX = Platypus::small_vgm[[2]], min.pct = .25,
group1 = "s1", group2 = "s2", return.plot = "volcano")

#Getting DEGs between two seurat clusters
GEX_DEgenes(GEX = Platypus::small_vgm[[2]], min.pct = .25,
#grouping.column = "seurat_clusters", group1 = "0", group2 = "1")

#Plotting a heatmap by foldchange of sample markers
GEX_DEgenes(GEX = VDJ_GEX_matrix.output[[2]],
#, min.pct = .25, group1 = "s1", group2 = "s2", return.plot = "heatmap"
#, up.genes = 10, down.genes = 10, logFC = TRUE)

#Plotting volcano by p value of sample markers. Label additional genes of interest
GEX_DEgenes(GEX = VDJ_GEX_matrix.output[[2]], min.pct = .25
#, group1 = "s1", group2 = "s2", return.plot = "volcano", logFC = FALSE
#, label.n.top.genes = 40, genes.to.label = c("CD28", "ICOS")

#Generate a heatmap from an already existing FindMarkers output
GEX_DEgenes(GEX = VDJ_GEX_matrix.output[[2]],
#, FindMarkers.out = FindMarkers.output.dataframe, return.plot = "heatmap"
#, up.genes = 10, down.genes = 10, logFC = TRUE, platypus.version = "v3")

Description

!Only for Platypus version v2. For more flexibility and platypus v3 please refer to GEX_Degenes. Extracts the differentially expressed genes between two samples. This function uses the FindMarkers function from the Seurat package. Further parameter control can be accomplished by calling the function directly on the output of automate_GEX and further extracting sample information from the "sample_id" component of the Seurat object.

Usage

GEX_DEgenes_persample(
    automate.GEX,
    min.pct,
    sample1,
    sample2,
by.group,
filter,
return.plot,
logFC,
up.genes,
down.genes,
base
)

Arguments

automate.GEX          Output Seurat object from automate_GEX function that contained at least two
                      distinct biological samples. The differential biological samples correspond to
                      integer values in the order of the working directories initially supplied to the
                      automate_GEX function.

min.pct               The minimum percentage of cells expressing a gene in either of the two groups
                      to be compared.

sample1               either character or integer specifying the first sample that should be compared.

sample2               either character or integer specifying the first sample that should be compared.

by.group             Logical specifying if groups should be used instead of samples. If TRUE, then
                      the argument in sample1 and sample2 will correspond to cells found in the
                      groups from sample1 or sample2.

filter                Character vector of initials of the genes to be filtered. Default is c("MT-", "RPL",
                      "RPS"), which filters mitochondrial and ribosomal genes.

return.plot           Logical specifying if a heatmap of the DEX genes is to be returned. If TRUE
                      then @return is a list where the first element is a dataframe and the second a
                      heatmap (see @return)

logFC                 Logical specifying whether the genes will be displayed based on logFC (TRUE)
                      or pvalue (FALSE).

up.genes              Integer specifying the number of upregulated genes to be shown.

down.genes            Integer specifying the number of downregulated genes to be shown.

base                  The base with respect to which logarithms are computed. Default: 2

Value

Returns a dataframe containing the output from the FindMarkers function, which contains informa-

Examples

## Not run:
GEX_DEgenes_persample(automate.GEX=automate.GEX.output[[1]],
, min.pct = .25, sample1 = "1", sample2 = "2")

## End(Not run)
GEX_dottile_plot

Description

Outputs a dotplot for gene expression, where the color of each dot is scaled by the gene expression level and the size is scaled by the % of cells positive for the gene.

Usage

GEX_dottile_plot(GEX, genes, group.by, threshold.to.plot, platypus.version)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GEX</td>
<td>GEX seurat object generated with VDJ_GEX_matrix</td>
</tr>
<tr>
<td>genes</td>
<td>Character vector. Genes of those in rownames(GEX) to plot. Can be any num-</td>
</tr>
<tr>
<td></td>
<td>ber, but more then 30 is discouraged because of cluttering</td>
</tr>
<tr>
<td>group.by</td>
<td>Character. Name of a column in <a href="mailto:GEX@meta.data">GEX@meta.data</a> to split the plot by. If set to</td>
</tr>
<tr>
<td></td>
<td>&quot;none&quot;, a plot with a single column will be produced.</td>
</tr>
<tr>
<td>threshold.to.plot</td>
<td>Integer 1-100. % of cells which must be expressing the feature to plot a</td>
</tr>
<tr>
<td></td>
<td>point. If below, the field will be left empty</td>
</tr>
<tr>
<td>platypus.version</td>
<td>This is coded for &quot;v3&quot; only, but in practice any Seurat Object can be fed in</td>
</tr>
</tbody>
</table>

Value

Returns a ggplot object were the dot size indicates the percentage of expressing cells and the dot color indicates the expression level.

Examples

#To return a plot detailing the expression of common genes by seurat cluster
GEX_dottile_plot(GEX = Platypus::small_vgm[[2]], genes = c("CD19","CD83"),
group.by = "seurat_clusters", threshold.to.plot = 5)
GEX_GOterm

GEX GO-Term analysis and plotting

Description

Runs a GO term analysis on a submitted list of genes. Works with the output of GEX_topN_DE_genes_per_cluster or a custom list of genes to obtain GOterms.

Usage

```r
GEX_GOterm(
  GEX.cluster.genes.output,
  topNgenes,
  ontology,
  species,
  up.or.down,
  MT.Rb.filter,
  kegg,
  go.plots,
  top.N.go.terms.plots,
  kegg.plots,
  top.N.kegg.terms.plots
)
```

Arguments

- `GEX.cluster.genes.output`: Either output of Platypus::GEX_cluster_genes or custom character vector containing gene symbols. A custom gene list will not be further filtered or ordered.
- `topNgenes`: How many of the most significant up or down regulated genes should be considered for GO term analysis. All genes will be used if left empty.
- `ontology`: Ontology used for the GO terms. "MF", "BP" or "CC" possible. Default: "BP"
- `species`: The species the genes belong to. Default: "Mm" (requires the package "org.Mm.eg.db"). Set to "Hs" for Human (requires the package "org.Hs.eg.db")
- `up.or.down`: Whether up or downregulated genes should be used for GO term analysis if GEX_cluster_genes output is used. Default: "up"
- `MT.Rb.filter`: logical, if mitochondrial and ribosomal genes should be filtered out.
- `kegg`: logical, if KEGG pathway analysis should be conducted. Requires internet connection. Default: False.
- `go.plots`: logical, if top GO-terms should be visualized. Default: False. If True, for each cluster the top N (top.N.GO.terms.plots) Go-terms for each cluster will be plotted to the working directory and saved as a list element. Plots are made both based on padj and ratio.
GEX_GSEA

Description

Conducts a Gene Set Enrichment Analysis (GSEA) on a set of genes submitted in a data frame with a metric each. Works with the output of GEX_genes_cluster or a custom data frame containing the gene symbols either in a column "symbols" or as rownames and a metric for each gene. The name of the column containing the metric has to be declared via the input metric.colname.

Usage

GEX_GSEA(
  GEX.cluster.genes.output,
  MT.Rb.filter,
  filter,
  top.N.go.terms.plots,
  The number of most significant GO-terms to be included in the go.plots. Default: 10.
  kegg.plots,
  logical, if top KEGG-terms should be visualized. Default: False. If True, for each cluster the top N (top.N.kegg.terms.plots) KEGG-terms for each cluster will be plotted to the working directory and saved as a list element. Plots are made both based on padj and ratio.
  top.N.kegg.terms.plots,
  The number of most significant KEGG-terms to be included in the kegg.plots. Default: 10.
)

Value

Returns a list of data frames and plots containing the TopGO and the TopKEGG output containing the significant GO/KEGG terms and their visualizations.

Examples

## Not run:
GEX_GOterm(DE_genes_cluster, MT.Rb.filter = TRUE, ontology = "MF")
GEX_GOterm(rownames(DE_genes_cluster[[1]]), MT.Rb.filter = TRUE, species = "Mm", ontology = "BP", go.plots = TRUE)

# Install the needed database with
# if (!requireNamespace("BiocManager", quietly = TRUE))
# install.packages("BiocManager")
# BiocManager::install("org.Mm.eg.db")
# BiocManager::install("org.Hs.eg.db")

## End(Not run)
path.to.pathways,
metric.colname,
pval.adj.cutoff,
Enrichment.Plots,
my.own.geneset,
eps,
platypus.version,
verbose
)

Arguments

GEX.cluster.genes.output
Data frame containing the list of gene symbols and a metric. Function works
directly with GEX_cluster_genes output.

MT.Rb.filter Logical, should Mitotic and Ribosomal genes be filtered out of the geneset. True
by default.

filter Character vector containing the identifying symbol sequence for the genes which
should be filtered out, if MT.Rb.filter == T. By default set to c("MT-", "RPL",
"RPS").

path.to.pathways
Either a path to gmt file containing the gene sets (can be downloaded from
MSigDB) or vector where first element specifies species and second element
specifies the MSigDB collection abbreviation. E.g.: c("Homo sapiens", "H").
Mouse C7 (immunologic signature) gene set will be used by default.

metric.colname Name of column which contains the metric used for the ranking of the submitted
genelist. "avg_logFC" is used by default.

pval.adj.cutoff
Only genes with a more significant adjusted pvalue are considered. Default: 0.001

Enrichment.Plots
List of Gene-set names which should be plotted as Enrichment plots in addition
to the top 10 Up and Downregulated Genesets.

my.own.geneset A list, where each element contains a gene list and is named with the corresponding
pathway name. Default is set to FALSE, so that gene sets from MSigDB are
used. Should not contain ".gmt" in name.

eps Numeric, specifying boundary for calculating the p value in the GSEA.

platypus.version Function works with V2 and V3, no need to set this parameter.

verbose Print run parameters and status to console

Value

Returns a list containing a tibble with the gene sets and their enrichment scores and Enrichment
plots. List element [1]: Dataframe with Genesets and statistics. [2]: Enrichment plots of top10
Up regulated genesets. [3]: Enrichment plots of top10 Down regulated genesets. [4]: Enrich-
ment plots of submited gene-sets in parameter Enrichment Plot.
Examples

```r
## Not run:
df <- GEX_cluster_genes(gex_combined[[1]])

# Using gmt file to perform gsea
output <- GEX_GSEA(GEX.cluster.genes.output = df[[1]], MT.Rb.filter = TRUE,
                   path.to.pathways = "/c5.go.bp.v7.2.symbols.gmt")
cowplot::plot_grid(plotlist=output[[2]], ncol=2)
View(gex_gsea[[1]])

# Directly downloading gene set collection from MSigDB to perform gsea
output <- GEX_GSEA(GEX.cluster.genes.output = df[[1]], MT.Rb.filter = TRUE,
                   path.to.pathways = c("Mus musculus", "C7")

# Using your own gene list to perform gsea
output <- GEX_GSEA(GEX.cluster.genes.output = df[[1]], MT.Rb.filter = TRUE,
                   my.own.geneset = my_geneset)

## End(Not run)
```

---

### GEX_heatmap

**Flexible GEX heatmap wrapper**

#### Description

Produces a heatmap containing gene expression information at the clonotype level. The rows correspond to different genes that can either be determined by pre-made sets of B or T cell markers, or can be customized by the user. The columns correspond to individual cells and the colors correspond to the different clonotype families.

#### Usage

```r
GEX_heatmap(
  GEX,
  b.or.t,
  sample.index,
  clone.rank.threshold,
  custom.array,
  slot
)
```

#### Arguments

- **GEX**
  
  A single seurat object from clonotype_GEX function corresponding to all of the samples in a single VDJ_analyze object. This will likely be supplied as clonotype_GEX.output[[i]] if there were multiple, distinct transcriptomes.

- **b.or.t**
  
  Logical indicating if B or T cell gene panel should be used.
GEX_pairwise_DEGs

Description

Produces and saves a list of volcano plots with each showing differentially expressed genes between pairs groups. If e.g. seurat_clusters used as group.by, a plot will be generated for every pairwise comparison of clusters. For large numbers of this may take longer to run. Only available for platypus v3

Usage

GEX_pairwise_DEGs(  
  GEX,  
  group.by,  
  min.pct,  

sample.index  Corresponds to which repertoire should be used in the case that the length of clonotype.list has a length greater than 1. The transcriptional profiles from only one repertoire can be plotted at a time.

custom.array  Corresponds to which repertoire should be used in the case that the length of clonotype.list has a length greater than 1. The transcriptional profiles from only one repertoire can be plotted at a time.

Value

Returns a heatmap via Seurat::DoHeatmap of gene expression per clonotype

See Also

VDJ_extract_sequences

Examples

# prep the small_vgm sample dataset  
small_vgm <- Platypus::small_vgm  
small_vgm[[2]]$clone_rank <- c(1:nrow(small_vgm[[2]]@meta.data))  
GEX_heatmap(GEX = small_vgm[[2]], b.or.t = "custom",  
  clone.rank.threshold = 1, sample.index = "s1",  
  custom.array = c("CD24A", "CD83"), slot = "data")

GEX_pairwise_DEGs  
Wrapper for calculating pairwise differentially expressed genes

Description

Produces and saves a list of volcano plots with each showing differentially expressed genes between pairs groups. If e.g. seurat_clusters used as group.by, a plot will be generated for every pairwise comparison of clusters. For large numbers of this may take longer to run. Only available for platypus v3

Usage

GEX_pairwise_DEGs(  
  GEX,  
  group.by,  
  min.pct,  

clone.rank.threshold  
A numeric that specifies the threshold clonal rank that specifies which clonotypes to extract transcriptome information from. For example, if 10 is supplied then the gene expression for the top ten clones included on the heatmap, separated by clonotype.

custom.array  
Corresponds to which repertoire should be used in the case that the length of clonotype.list has a length greater than 1. The transcriptional profiles from only one repertoire can be plotted at a time.

slot  
Seurat data slot from which to plot values. Can be "raw.data", "data" or "scale.data"
GEX_phenotype

RP_MT.filter,
label.n.top.genes,
genes.to.label,
save.plot
)

Arguments

GEX Output Seurat object of the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[2]])
group.by Character. Defaults to "seurat_clusters" Column name of GEX@meta.data to use for pairwise comparisons. More than 20 groups are discouraged.
min.pct Numeric. Defaults to 0.25 passed to Seurat::FindMarkers
RP_MT.filter Boolean. Defaults to True. If True, mitochondrial and ribosomal genes are filtered out from the output of Seurat::FindMarkers
label.n.top.genes Integer. Defaults to 50. Defines how many genes are labelled via geom_text_repel. Genes are ordered by adjusted p value and the first label.n.genes are labelled
genes.to.label Character vector. Defaults to "none". Vector of gene names to plot independently of their p value. Can be used in combination with label.n.genes.
save.plot Boolean. Defaults to True. Whether to save plots as appropriately named .png files

Value

A nested list with out[i][1] being ggplot volcano plots and out[i][2] being source DEG dataframes.

Examples

GEX_pairwise_DEGs(GEX = Platypus::small_vgm[[2]], group.by = "sample_id", min.pct = 0.25, RP_MT.filter = TRUE, label.n.top.genes = 5, genes.to.label = c("CD24A"), save.plot = FALSE)

GEX_phenotype Assignment of cells to phenotypes based on selected markers

Description

Adds a column to a VGM[[2]] Seurat object containing cell phenotype assignments. Defaults for T and B cells are available. Marker sets are customizable as below

Usage

GEX_phenotype(seurat.object, cell.state.names, cell.state.markers, default)
ARGUMENTS

seurat.object  A single seurat object / VDJ_GEX_matrix.output[[2]] object

cell.state.names
Character vector containing the cell state labels defined by the markers in cell.state.markers parameter. Example is c("NaiveCd4","MemoryCd4").

cell.state.markers
Character vector containing the gene names for each state. ; is used to use multiple markers within a single gene state. Different vector elements correspond to different states. Order must match cell.state.names containing the c("CD4+;CD44-","CD4+;IL7R+;CD44+").

default  Default is TRUE - will use predefined gene sets and cell states.

Value

Returns the input Seurat object with an additional column

Examples

vgm.phenotyped <- GEX_phenotype(seurat.object = Platypus::small_vgm[[2]], default = TRUE)

GEX_phenotype_per_clone

Plotting of GEX phenotype by VDJ clone

Description

Integrates VDJ and gene expression libraries by providing cluster membership seq_per_vdj object and the index of the cell in the Seurat RNA-seq object. ! For platypus.version == "v3" and VDJ_GEX_matrix output the function will iterate over entries in the sample_id column of the GEX by default.

Usage

GEX_phenotype_per_clone(
  GEX,
  clonotype.ids,
  global.clonotypes,
  GEX.group.by,
  GEX.clonotypes,
  platypus.version
)
Arguments

GEX For platypus.version == "v3" the GEX object from the output of the VDJ_GEX_matrix function (VDJ_GEX_matrix.output \[\[2\]\]). For platypus.version == "v2" a single seurat object from automate_GEX function after labeling cell phenotypes using the GEX_phenotype function.

clonotype.ids For platypus.version == "v2" Output from either VDJ_analyze or VDJ_clonotype functions. This list should correspond to a single GEX.list object, in which each list element in clonotype.list is found in the GEX.object. Furthermore, these repertoires should be found in the automate_GEX library.

global.clonotypes Boolean. Defaults to FALSE. Set to True if clonotyping has been done across samples

GEX.group.by For platypus.version == "v3". Character. Column name of the GEX@meta.data to group barplot by. Defaults to seurat_clusters

GEX.clonotypes For platypus.version == "v3". Numeric vector with ids of clonotypes to plot e.g. c(1,2,3,4). Can also be set to "topclones"

platypus.version Set to either "v2" or "v3" depending on wether suppyling GEX_automate or VDJ_GEX_matrix\[\[2\]\] objects. Defaults to "v3"

Value

Returns a stacked barplot that visualizes the seurat cluster membership for different cell phenotypes.

Examples

#For testing: only a single clonotype in two samples
small_vgm_cl <- Platypus::small_vgm
small_vgm_cl[[2]]$clonotype_id_10x <- "clonotype1"
GEX_phenotype_per_clone(GEX = small_vgm_cl[[2]]
, GEX.clonotypes = c(1), GEX.group.by = "seurat_clusters", platypus.version = "v3")

GEX_proportions_barplot

Plots proportions of a group of cells within a secondary group of cells. E.g. The proportions of samples in seurat clusters, or the proportions of samples in defined cell subtypes

Description

Plots proportions of a group of cells within a secondary group of cells. E.g. The proportions of samples in seurat clusters, or the proportions of samples in defined cell subtypes

Usage

GEX_proportions_barplot(GEX, source.group, target.group, stacked.plot, verbose)
**GEX_scatter_coexpression**

Scatter plot for coexpression of two selected genes

**Description**

Plots a composite figure showing single marker expression as histograms and coexpression as a scatterplot.

**Usage**

```r
GEX_scatter_coexpression(GEX, gene.1, gene.2, color.theme)
```
**Arguments**

- **GEX**
  - GEX seurat object generated with VDJ_GEX_matrix
- **gene.1**
  - Character. Name of a gene in rownames(VDJ.matrix)
- **gene.2**
  - Character. Name of a gene in rownames(VDJ.matrix)
- **color.theme**
  - Character. A color to use for the composite plot

**Value**

Returns a gridplot showing coexpression scatterplot as well as histograms of gene.1 and gene.2

**Examples**

```r

gene1 <- "CD24A"
gene2 <- "CD83"
GEX_scatter_coexpression(GEX = Platypus::small_vgm[[2]], gene1, gene2)
```

---

**Description**

Organizes the top N genes that define each Seurat cluster and converts them into a single dataframe. This can be useful for obtaining insight into cluster-specific phenotypes.

**Usage**

```r
GEX_topN_DE_genes_per_cluster(GEX_cluster_genes.output, n.genes, by_FC, filter)
```

**Arguments**

- **GEX_cluster_genes.output**
  - The output from the GEX_cluster_genes function - this should be a list with each list element corresponding to the genes, p values, logFC, pct expression for the genes differentially regulated for each cluster.
- **n.genes**
  - The number of genes to be selected from each cluster. If n.genes is higher than the number of cells in a cluster then it is silently adjusted to be
- **by_FC**
  - Logical indicating if the top n genes are selected based on the logFC value instead of p value. Default is FALSE.
- **filter**
  - Character vector of initials of the genes to be filtered. Default is c("MT-", "RPL", "RPS"), which filters mitochondrial and ribosomal genes.

**Value**

Returns a dataframe in which the top N genes defining each cluster based on differential expression are selected.
## Not run:
GEX_topDE_genes_per_cluster(GEX_cluster_genes.output=list_of_genes_per_cluster, n.genes=20, by.FC=FALSE, filter=c("MT-", "RPS", "RPL"))

## End(Not run)

GEX_visualize_clones

Platypus V2 GEX and VDJ integration for visualizing clone clustering

Description

!Only for platypus version v2. For platypus v3 refer to: VDJ_GEX_overlay_clones() Visualize selected clonotypes on the tSNE or UMAP projection.

Usage

GEX_visualize_clones(  
  GEX.list,  
  VDJ.GEX.integrate.list,  
  highlight.type,  
  highlight.number,  
  reduction  
)

Arguments

- **GEX.list**: list of Seurat objects, output of the automate_GEX function.
- **VDJ.GEX.integrate.list**: Output of the VDJ_GEX_integrate function.
- **highlight.type**: (Optional) either "None" if representation highlighted by cluster, "clonotype" if want to highlight most expanded clonotypes, or "sample" if several samples are within the same Seurat object. Default is None.
- **highlight.number**: (Optional) an integer or list of integers representing the number of most expanded clonotypes or samples one wants to select eg 4 to highlight the 4th most expanded clonotype or 2:5 to highlight the top 2 to top 5 most expanded clonotype. Only compatible with highlight.type "clonotype" or "sample", will be ignored if type is "None". Default is 1.
- **reduction**: (Optional) Reduction to plot, either "tsne", "umap", or "harmony". Default is "tsne".

Value

concatenated ggplot2 plot with selected clonotypes highlighted (if None, the coloring is according to the clustering).
GEX_volcano

Examples

```r
## Not run:
GEX_visualize_clones(GEX.list=automate_GEX.output,
  VDJ.per.clone=VDJ_per_clone.output,
  highlight.type="clonotype",
  highlight.number=1:4,
  reduction="umap")

## End(Not run)
```

---

GEX_volcano

Flexible wrapper for GEX volcano plots

Description

Plots a volcano plot from the output of the FindMarkers function from the Seurat package or the GEX_cluster_genes function alternatively.

Usage

```r
GEX_volcano(
  DEGs.input,
  input.type,
  condition.1,
  condition.2,
  explicit.title,
  RP.MT.filter,
  color.p.threshold,
  color.log.threshold,
  label.p.threshold,
  label.logfc.threshold,
  n.label.up,
  n.label.down,
  by.logFC,
  maximum.overlaps,
  plot.adj.pvalue
)
```

Arguments

- **DEGs.input**: Either output data frame from the FindMarkers function from the Seurat package or GEX_cluster_genes list output.
- **input.type**: Character specifying the input type as either "findmarkers" or "cluster.genes". Defaults to "cluster.genes"
- **condition.1**: either character or integer specifying ident.1 that was used in the FindMarkers function from the Seurat package. Should be left empty when using the GEX_cluster_genes output.
condition.2 either character or integer specifying ident.2 that was used in the FindMarkers function from the Seurat package. Should be left empty when using the GEX_cluster_genes output.

explicit.title logical specifying whether the title should include logFC information for each condition.

RP.MT.filter Boolean. Defaults to TRUE. Whether to exclude ribosomal and mitochondrial genes.

color.p.threshold numeric specifying the adjusted p-value threshold for geom_points to be colored. Default is set to 0.01.

color.log.threshold numeric specifying the absolute logFC threshold for geom_points to be colored. Default is set to 0.25.

label.p.threshold numeric specifying the adjusted p-value threshold for genes to be labeled via geom_text_repel. Default is set to 0.001.

label.logfc.threshold numeric specifying the absolute logFC threshold for genes to be labeled via geom_text_repel. Default is set to 0.75.

n.label.up numeric specifying the number of top upregulated genes to be labeled via geom_text_repel. Genes will be ordered by adjusted p-value. Overrides the "label.p.threshold" and "label.logfc.threshold" parameters.

n.label.down numeric specifying the number of top downregulated genes to be labeled via geom_text_repel. Genes will be ordered by adjusted p-value. Overrides the "label.p.threshold" and "label.logfc.threshold" parameters.

by.logFC logical. If set to TRUE n.label.up and n.label.down will label genes ordered by logFC instead of adjusted p-value.

maximum.overlaps integer specifying removal of labels with too many overlaps. Default is set to Inf.

plot.adj.pvalue logical specifying whether adjusted p-value should by plotted on the y-axis.

Value

Returns a volcano plot from the output of the FindMarkers function from the Seurat package, which is a ggplot object that can be modified or plotted. Infinite p-values are set defined value of the highest -log(p) + 100.

Examples

```r
## Not run:
#using the findmarkers.output
GEX_volcano(findmarkers.output = FindMarkers.Output
, condition.1 = "cluster1", condition.2 = "cluster2"
, maximum.overlaps = 20)
```
GEX_volcano(findmarkers.output = FindMarkers.Output, condition.1 = "cluster1", condition.2 = "cluster2", n.label.up = 50, n.label.down = 20)

#using the GEX_cluster_genes output
GEX_volcano(findmarkers.output = GEX_cluster_genes.Output, cluster.genes.output = TRUE)

## End(Not run)

---

**hotspot_df**

*hotspot_df* Hotspot mutations taken from Yaari et al., *Frontiers in Immunology*, 2013. This contains transition probabilities for all 5mer combinations based on high throughput sequencing data. The transition probabilities are for the middle nucleotide in each 5mer set. This can be customized by changing the genes and sequences. Custom mutation hotspots can be supplied by modifying this dataframe. Repeating particular hotspot entries allows for the hotspot to mutate more than one time per SHM event.

**Description**

@format A data frame with 1024 rows and 6 variables:

- **pattern** Character array where each entry corresponds to a 5 base motif. The mutation probabilities correspond to the middle nucleotide in each 5mer.
- **toA** The probability for the middle nucleotide in "pattern" to mutate to an adenine
- **toC** The probability for the middle nucleotide in "pattern" to mutate to an cytosine
- **toG** The probability for the middle nucleotide in "pattern" to mutate to an guanine
- **toT** The probability for the middle nucleotide in "pattern" to mutate to an thymine

**Source** The origin of how this motif was discovered. Either Inferred or Experimental

**Usage**

data("hotspot_df")

**Format**

An object of class data.frame with 1024 rows and 6 columns.

**Source**

Yaari et al., *Frontiers in Immunology*, 2013
Description

human germline IgH (heavy chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customizing by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

Usage

data("hum_b_h")

Format

A list including 3 elements (data frames): v gene, d gene, j gene, respectively.

[[1]]
gene The v gene name
seq The corresponding sequence [[2]]
gene The d gene name
seq The corresponding sequence [[3]]
gene The j gene name
seq The corresponding sequence

Source

IMGT

Description

human germline IgH (light chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customizing by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

Usage

data("hum_b_l")
**hum_t_h**

**Format**

A list including 2 elements (data frames): v gene, d gene, j gene, respectively.

- **gene** The v gene name
- **seq** The corresponding sequence [[2]]
- **gene** The j gene name
- **seq** The corresponding sequence

**Source**

IMGT

---

**hum_t_h**

---

**Description**

human germline TRB (heavy chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customized by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

**Usage**

```r
data("hum_t_h")
```

**Format**

A list including 3 elements (data frames): v gene, d gene, j gene, respectively.

- **gene** The v gene name
- **seq** The corresponding sequence [[2]]
- **gene** The d gene name
- **seq** The corresponding sequence [[3]]
- **gene** The j gene name
- **seq** The corresponding sequence

**Source**

IMGT
Description

human germline TRA (light chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customized by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

Usage

data("hum_t_l")

Format

A list including 2 elements (data frames): v gene, d gene, j gene,respectively.

[1]

gene  The v gene name
seq   The corresponding sequence

[2]

gene  The j gene name
seq   The corresponding sequence

Source

IMGT

iso_SHM_prob

A probability dataframe specifying SHM.nuc.prob for cells of different isotypes. The first column is the names of isotypes, while the second column is the SHM.nuc.prob of cell of that isotype. user can define different SHM.nuc.prob for isotypes.

Description

iso_SHM_prob A probability dataframe specifying SHM.nuc.prob for cells of different isotypes. The first column is the names of isotypes, while the second column is the SHM.nuc.prob of cell of that isotype. user can define different SHM.nuc.prob for isotypes.

Usage

data("iso_SHM_prob")

Format

a dataframe with 2 columns
Description
C57BL/6 germline IgH (heavy chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customized by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

Usage
data("mus_b_h")

Format
A list including 3 elements (data frames): v gene, d gene, j gene, respectively.

[[1]]
gene The v gene name
seq The corresponding sequence [[2]]
gene The d gene name
seq The corresponding sequence [[3]]
gene The j gene name
seq The corresponding sequence

Source
IMGT

Description
C57BL/6 germline IgH (light chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customized by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

Usage
data("mus_b_l")
Format

A list including 2 elements (data frames): `v gene`, `d gene`, `j gene`, respectively.

[[1]]

`gene` The `v gene` name

`seq` The corresponding sequence [[2]]

`gene` The `j gene` name

`seq` The corresponding sequence

Source

IMGT

---

**mus_b_trans**

`mus_b_trans` A data frame contains mouse B cell average gene expression for multiple cell types, with the rows representing the gene names, column names representing the cell type names. The original single cell sequencing data is retrieved from 10xgenomics and combined with experimental data from. The expression level for different cell types are obtained by calculating the average expression after sorting the original data by markers as shown below.

- **NaiveBcell** `Cd19+;Cd27-;Cd38-
- **GerminalcenterBcell** `Fas+;Cd19+
- **Plasmacell** `Sdc1+
- **MemoryBcell** `Cd38+;Fas-

Description

`mus_b_trans` A data frame contains mouse B cell average gene expression for multiple cell types, with the rows representing the gene names, column names representing the cell type names. The original single cell sequencing data is retrieved from 10xgenomics and combined with experimental data from. The expression level for different cell types are obtained by calculating the average expression after sorting the original data by markers as shown below.

- **NaiveBcell** `Cd19+;Cd27-;Cd38-
- **GerminalcenterBcell** `Fas+;Cd19+
- **Plasmacell** `Sdc1+
- **MemoryBcell** `Cd38+;Fas-

Usage

`data("mus_b_trans")`
**Format**

A data frame with 26538 rows and 4 variables, with the rows representing the gene names, column names representing the cell type names.

**Source**

https://support.10xgenomics.com/single-cell-vdj/datasets/3.0.0/vdj_v1_mm_c57bl6_pbmc_5gex
https://support.10xgenomics.com/single-cell-vdj/datasets/3.0.0/vdj_v1_mm_balbc_pbmc_5gex

**Description**

C57BL/6 germline TRB (heavy chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customized by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

**Usage**

```r
data("mus_t_h")
```

**Format**

A list including 3 elements (data frames): v gene, d gene, j gene,respectively.

- `gene` The v gene name
- `seq` The corresponding sequence
- `gene` The d gene name
- `seq` The corresponding sequence
- `gene` The j gene name
- `seq` The corresponding sequence

**Source**

IMGT
Description

C57BL/6 germline TRA (light chain v,d,j gene segments). When multiple alleles were present, the first one was included. These names and sequences can be changed by customized by changing this dataframe. Additionally, repeating elements can give certain germline gene elements a larger probability of being used during repertoire evolution.

Usage

data("mus_t_l")

Format

A list including 2 elements (data frames): v gene, d gene, j gene,respectively.

[[1]]
  gene  The v gene name
  seq  The corresponding sequence [[2]]
  gene  The j gene name
  seq  The corresponding sequence

Source

IMGT

no.empty.node

Get clone network igraphs without empty mode. Empty node represents the 'extincted' sequences, that are not in any living cell but once existed.

Description

Get clone network igraphs without empty mode. Empty node represents the 'extincted' sequences, that are not in any living cell but once existed.

Usage

no.empty.node(history, igraph.index)
**Arguments**

- **history**
  The dataframe 'history' from the simulation output.

- **igraph.index**
  The list 'igraph.index' from the simulation output.

- **empty.node**
  If TRUE, there will be empty node in igraph. If FALSE, the empty node will be deleted.

**Value**

a list of clone network igraphs without empty mode.

**Description**

WRC hotspot mutations taken from Yaari et al., Frontiers in Immunology, 2013. These include only the mutations following the WRC pattern, where W equals A or T and R equals A or G. Custom mutation hotspots can be supplied by modifying this dataframe. Repeating particular hotspot entries allows for the hotspot to mutate more than one time per SHM event.

**Usage**

data("one_spot_df")

**Format**

A data frame with 32 rows and 6 variables:

- **pattern** Character array where each entry corresponds to a 5 base motif. The mutation probabilities correspond to the middle nucleotide in each 5mer.
- **toA** The probability for the middle nucleotide in "pattern" to mutate to an adenine
- **toC** The probability for the middle nucleotide in "pattern" to mutate to an cytosine
- **toG** The probability for the middle nucleotide in "pattern" to mutate to an guanine
- **toT** The probability for the middle nucleotide in "pattern" to mutate to an thymine

**Source**

The origin of how this motif was discovered. Either Inferred or Experimental

Yaari et al., Frontiers in Immunology, 2013
**Pheno_SHM_prob**

A probability dataframe specifying SHM.nuc.prob for cells of different phenotypes. The first column is the names of phenotypes, while the second column is the SHM.nuc.prob of cell of that phenotype. User can define different SHM.nuc.prob for phenotypes.

**Description**

Pheno_SHM_prob A probability dataframe specifying SHM.nuc.prob for cells of different phenotypes. The first column is the names of phenotypes, while the second column is the SHM.nuc.prob of cell of that phenotype. User can define different SHM.nuc.prob for phenotypes.

**Usage**

```r
data("pheno_SHM_prob")
```

**Format**

A dataframe with 2 columns

-------------------

**PlatypusDB_AIRR_to_VGM**

**AIRR to Platypus V3 VGM compatibility function**

**Description**

PlatypusDB_AIRR_to_VGM_AIRR to Platypus V3 VGM compatibility function Loads in and converts input AIRR-compatible tsv file(s) into the Platypus VGM object format. All compulsory AIRR data columns are needed. Additionally, the following columns are required: v_call, cell_id, clone_id. If trim.and.align is set to TRUE additionally the following columns are needed: v_sequence_start, j_sequence_end. Note on TRUST4 input: TRUST4 (https://doi.org/10.1038/s41592-021-01142-n2) is a newly alignment tool for VDJ data by the Shirley lab. It is able to also extract VDJ sequences from 10x GEX data. We are actively testing TRUST4 as an alternative to Cellranger and can not give recommendations as of now. This function does support the conversion of TRUST4 airr output data into the Platypus VGM format. In that case, an extra column will be added describing whether the full length VDJ sequence was extracted for any given cell and chain.

**Usage**

```r
PlatypusDB_AIRR_to_VGM(
    AIRR.input,
    get.VDJ.stats,
    VDJ.combine,
    trim.and.align,
    filter.overlapping.barcodes.VDJ,
    group.id,
    verbose
)
```
PlatypusDB_fetch

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AIRR.input</td>
<td>Source of the AIRR table(s) as a list. There are 2 available input options: 1. 1. List with local paths to .tsv files / 3. List of AIRR tables loaded in as R objects within the current R environment.</td>
</tr>
<tr>
<td>get.VDJ.stats</td>
<td>Boolean. Defaults to TRUE. Whether to generate summary statistics on repertoires and output those as output_VGM[[3]]</td>
</tr>
<tr>
<td>VDJ.combine</td>
<td>Boolean. Defaults to TRUE. Whether to integrate repertoires. A sample identifier will be appended to each barcode both. Highly recommended for all later functions</td>
</tr>
<tr>
<td>trim.and.align</td>
<td>Boolean. defaults to FALSE. Whether to trim VJ/VDJ seqs and add information from alignment in AIRR dataframe columns. ! No alignment is done here, instead, columns containing alignment information in the AIRR dataframes are reformatted.</td>
</tr>
<tr>
<td>filter.overlapping.barcodes.VDJ</td>
<td>Boolean. defaults to TRUE. Whether to remove barcodes which are shared among samples in the GEX analysis. Shared barcodes normally appear at a very low rate.</td>
</tr>
<tr>
<td>group.id</td>
<td>vector with integers specifying the group membership. c(1,1,2,2) would specify the first two elements of the input AIRR list are in group 1 and the third/fourth input elements will be in group 2.</td>
</tr>
<tr>
<td>verbose</td>
<td>Writes runtime status to console. Defaults to FALSE</td>
</tr>
</tbody>
</table>

**Value**

A VDJ_GEX_Matrix object used in Platypus V3 as an input to most analysis and plotting functions

**Examples**

```r
VGM <- PlatypusDB_AIRR_to_VGM(AIRR.input = list("/path/to/s1/airr_rearrangement.tsv", "/path/to/s2/airr_rearrangement.tsv"),
VDJ.combine = TRUE, group.id = c(1,2), filter.overlapping.barcodes.VDJ = TRUE)
```

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

PlatypusDB_fetch(
  PlatypusDB.links,
  save.to.disk,
  load.to.environment,
  load.to.list,
  path.to.save,
  combine.objects
)

Arguments

PlatypusDB.links
Character vector. One or more links to files in the PlatypusDB. Links are constructed as follows: 
"%Project id%/%sample_id%/%filetype%". Any of the three can be "ALL", to download all files fitting the other link elements. If %filetype% is gexVGM, vdjVGM or metadata, %sample_id% needs to be "ALL", as these are elements which are not divided by sample. See examples for clarification. See last example on how to download AIRR compliant data. Feature Barcode (FB) data will be downloaded both for GEX and VDJ if present and does not need to be specified in the path. For sample_id entries the the metadata table for a given project via the function PlatypusDB_list_projects()

save.to.disk
Boolean. Defaults to FALSE. Whether to save downloaded files individually to the directory specified in path.to.save

load.to.environment
Boolean. Defaults to TRUE. Whether to load objects directly into the current .GlobalEnv. An array of the names of the loaded objects will be returned. !Be aware of RAM limitations of your machine when downloading multiple large files.

load.to.list
Boolean. Defaults to FALSE. Whether to return loaded objects as a list. !Be aware of RAM limitations of your machine when downloading multiple large files.

path.to.save
System path to save files to.

combine.objects
Boolean. Defaults to TRUE. Whether to combine objects if appropriate. e.g. VDJ and GEX RData objects for a sample are saved as two independent objects and downloaded as such, to allow for flexibility. If combine.objects is set to TRUE, the function will coerce RData objects of each loaded sample or of each loaded VDJ_GEX_matrix appropriately. Combined input of VDJ and GEX Rdata objects can be directly supplied to the VDJ_GEX_matrix function.

Value

A list of loaded project files as R objects if load.to.list = T or a name of these object loaded to the environment if load.to.environment = T.
Examples

## Not run:

```r
# Get a list of available projects by name
names(PlatypusDB_list_projects())

# Load the VDJ_GEX_matrix of a project as an object and
# also save it to disk for later.
# This will download the VDJ and GEX part of the VDJ_GEX_matrix and combine
PlatypusDB_fetch(PlatypusDB.links = c("Kuhn2021a//ALL"),
    save.to.disk = FALSE, load.to.environment = TRUE, load.to.list = FALSE,
    combine.object = TRUE, path.to.save = "/Downloads")

# Load VDJ dataframe of the VDJ GEX matrix for all samples of one project
loaded_list <- PlatypusDB_fetch(PlatypusDB.links = c("Kuhn2021a//VDJmatrix"),
    save.to.disk = FALSE, load.to.environment = FALSE, load.to.list = TRUE)

# Load the VDJ and GEX RData of 2 samples from
# 2 different projects which can be directly passed
# on to the VDJ_GEX_matrix function to integrate
# downloaded_objects <- PlatypusDB_fetch(
#    PlatypusDB.links = c("Project1/s1/ALL", "Project1/s2/ALL"),
#    save.to.disk = FALSE, load.to.environment = FALSE, load.to.list = TRUE)
#    combine.objects = TRUE)

# Integrate_samples <- VDJ_GEX_matrix_DB(data.in = downloaded_objects)

# Download metadata objects for projects
list_of_metadata_tables <- PlatypusDB_fetch(
    PlatypusDB.links = c("Kuhn2021a//metadata"),
    save.to.disk = FALSE, load.to.environment = FALSE, load.to.list = TRUE)

# Download of airr_rearrangement.tsv
# Load VDJ.RData into a list
# downloaded_objects <- PlatypusDB_fetch(
#    PlatypusDB.links = c("Project1/ALL/VDJ.RData"), save.to.disk = FALSE
#    load.to.environment = FALSE, load.to.list = TRUE)

# Extract airr_rearrangement table for sample 1
# airr_rearrangement <- downloaded_objects[[1]][[1]][[6]]
# Index hierarchy: Sample, VDJ or GEX, VDJ element

# Save for import to AIRR compatible pipeline
# write.table(airr_rearrangement, file = "airr_rearrangement_s1.tsv", sep = "\t",
#         row.names = FALSE, quote=FALSE)
```

## End(Not run)
PlatypusDB_find_CDR3s  

**CDR3 query function for PlatypusDB**

### Description
Queries for the occurrence of CDR3 sequences in public datasets on PlatypusDB.

### Usage

```r
PlatypusDB_find_CDR3s(VDJ.cdr3s.aa, VJ.cdr3s.aa, projects.to.search)
```

### Arguments
- **VDJ.cdr3s.aa**: Character A VDJ CDR3s amino acid sequence to search for
- **VJ.cdr3s.aa**: Character A VJ CDR3s amino acid sequence to search for
- **projects.to.search**: Optional character vector. Defaults to "ALL". Names of projects to search within.

### Value
A list of subsets of VDJ matrices from projects containing the query VDJ CDR3 (out[[1]]), the VJ CDR3 (out[[2]]) and cells containing both the query VDJ and VJ CDR3s (out[[3]])

### Examples

```r
## Not run:
public_clones <- PlatypusDB_find_CDR3s(VDJ.cdr3s.aa = "CMRYGNYWYFDVW", VJ.cdr3s.aa = "CLQHGESPFTF", projects.to.search = "ALL")
## End(Not run)
```

---

PlatypusDB_list_projects

**Metadata download by project for PlatypusDB**

### Description
Lists metadata tables of available projects on PlatypusDB.

### Usage

```r
PlatypusDB_list_projects(keyword)
```
Arguments

keyword Character. Keyword by which to search project ids (First Author, Year) in the
database. Defaults to an empty string (""), which will list all projects currently
available.

Value

A list of metadata tables by project. List element names correspond to project ids to use in the
PlatypusDB_fetch function.

Examples

## Not run:

# Get list of all available projects and metadata.
PlatypusDB_projects <- PlatypusDB_list_projects()

# Names of list are project ids to use in PlatypusDB_fetch function
names(PlatypusDB_projects)
# Common format: first author, date, letter a-z (all lowercase)

# View metadata of a specific project
print(PlatypusDB_projects[["Kuhn2021a"]])

## End(Not run)

---

**PlatypusDB_load_from_disk**

*PlatypusDB utility for import of local datasets*

**Description**

Utility function for loading in local dataset as VDJ_GEX_matrix and PlatypusDB compatible R
objects. Especially useful when wanting to integrate local and public datasets. This function only
imports and does not make changes to format, row and column names. Exception: filtered_contig.fasta
are appended to the filtered_contig_annotations.csv as a column for easy access.

**Usage**

PlatypusDB_load_from_disk(
    VDJ.out.directory.list,
    GEX.out.directory.list,
    FB.out.directory.list,
    batches
)
Arguments

VDJ.out.directory.list
List containing paths to VDJ output directories from cell ranger. This pipeline assumes that the output file names have not been changed from the default 10x settings in the /outs/ folder. This is compatible with B and T cell repertoires (both separately and simultaneously).

GEX.out.directory.list
List containing paths the outs/ directory of each sample or directly the raw or filtered_feature_bc_matrix folder. Order of list items must be the same as for VDJ. This outs directory may also contain Feature Barcode (FB) information. Do not specify FB.out.directory in this case.

FB.out.directory.list
List of paths pointing at the outs/ directory of output of the Cellranger counts function which contain Feature barcode counts. Any input will overwrite potential FB data loaded from the GEX input directories. Length must match VDJ and GEX directory inputs. (in case of a single FB output directory for multiple samples, please specify this directory as many times as needed)

batches
Integer vector. Defaults to all 1, yielding all samples with batch number "b1". Give a batch number to each sample (each entry in the VDJ/GEX input lists). This will be saved as element 5 in the sample list output.

Value

Large nested list object containing all needed Cellranger outputs to run the VDJ_GEX_matrix function. Level 1 of the list are samples, level 2 are VDJ GEX and metadata information. (e.g. out[[1]][[1]] corresponds to VDJ data objects of sample 1)

Examples

```r
## Not run:
VDJ.in <- list()
VDJ.in[[1]] <- c("~/VDJ/S1/")
VDJ.in[[2]] <- c("~/VDJ/S2/")
GEX.in <- list()
GEX.in[[1]] <- c("~/GEX/S1/")
GEX.in[[2]] <- c("~/GEX/S2/")
PlatypusDB_load_from_disk(VDJ.out.directory.list = VDJ.in, GEX.out.directory.list = GEX.in)
```

## End(Not run)

PlatypusDB_VGM_to_AIRR

### Platypus V3 VGM to AIRR compatibility function

Description

Exports AIRR compatible tables supplemented with VDJ and GEX information from the Platypus VGM object and the cellranger output airr_rearrangements.tsv
Usage

PlatypusDB_VGM_to_AIRR(
    VGM,
    VDJ.features.to.append,
    GEX.features.to.append,
    airr.rearrangements,
    airr.integrate
)

Arguments

VGM Output object of the VDJ.GEX_matrix function generated with VDJ.combine = T, GEX.combine = T (to merge all samples) and integrate.VDJ.to.GEX = T (to integrate VDJ and GEX data)

VDJ.features.to.append Character vector. Defaults to "none". Can be either "all" or column names of the VGM VDJ matrix (VGM[[1]]) to append to the AIRR compatible table.

GEX.features.to.append Character vector. Defaults to "none". Can be either "all" or GEX metadata column names or Gene names of the VGM GEX object (VGM[[2]])(passed to Seurat::FetchData()) to append to the AIRR compatible table. For a list of available features run: names(VGM[[2]]@meta.data) and rownames(VGM[[2]])

airr.rearrangements Source of the airr_rearrangements.tsv file as generated by Cellranger. There are 3 available input options: 1. R list object from Platypus_DB_load_from_disk or Platypus_DB_fetch / 2. List with local paths to airr_rearrangements.tsv / 3. List of airr_rearrangements.tsv loaded in as R objects within the current R environment. ! Order of input list must be identical to that of sample_ids in the VGM ! If not provided or set to "none" CIGAR strings in output will be empty.

airr.integrate Boolean. Defaults to TRUE, whether to integrate output AIRR tables

Value

A list of length of samples in VGM containing a AIRR-compatible dataframe for each sample if airr.integrate = F or a single dataframe if airr.integrate = T ! Cave the format: VGM object => 1 cell = 1 row; AIRR table 1 cell = as many rows as VDJ and VJ chains available for that cell. GEX cell-level information is attached to all rows containing a chain of that cell.

Examples

## Not run:  
#complete workflow below

#usage with airr rearrangement tables from PlatypusDB_load_from_disk  
#or PlatypusDB_fetch list object
airr.list.out <- PlatypusDB_VGM_to_AIRR(VGM = VGM
  , VDJ.features.to.append = c("VDJ_cdr3s_aa")
  , GEX.features.to.append = c("CTLA4", "TOX")
  , airr.rearrangements = Data.in)
#usage with airr rearrangement tables from disk
airr.list.out <- PlatypusDB_VGM_to_AIRR(VGM = VGM
, VDJ.features.to.append = c("VDJ_cdr3s_aa")
, GEX.features.to.append = c("CTLA4", "TOX"),
airr.rearrangements =list("~/path_to/s1/airr.rearrangement.tsv"
,"~/path_to/s2/airr.rearrangement.tsv")
)

#usage with airr rearrangement tables from objects in R environment
airr.list.out <- PlatypusDB_VGM_to_AIRR(VGM = VGM
, VDJ.features.to.append = c("VDJ_cdr3s_aa")
, GEX.features.to.append = c("CTLA4", "TOX"),
airr.rearrangements = list(airr_rearrangements.s1, airr_rearrangements_2)
)

#Complete workflow
#set paths of cellranger directories containing
#also the airr_rearrangements.tsv file
VDJ.out.directory.list <- list()
VDJ.out.directory.list[[1]] <- c("~/cellrangerVDJ/s1")
VDJ.out.directory.list[[2]] <- c("~/cellrangerVDJ/s2")
GEX.out.directory.list <- list()
GEX.out.directory.list[[1]] <- c("~/cellrangerGEX/s1")
GEX.out.directory.list[[2]] <- c("~/cellrangerGEX/s2")
#Run VGM with GEX and VDJ integration
VGM <- VDJ_GEX_matrix(VDJ.out.directory.list = VDJ.out.directory.list,
GEX.out.directory.list = GEX.out.directory.list,
GEX.integrate = TRUE, VDJ.combine = TRUE, integrate.GEX.to.VDJ = TRUE
, integrate.VDJ.to.GEX = TRUE,
get.VDJ.stats = FALSE, trim.and.align = FALSE)
#Generate AIRR compatible table supplemented by GEX information
airr.list.out <- PlatypusDB_VGM_to_AIRR(VGM = VGM,
VDJ.features.to.append = c("VDJ_sequence_nt_trimmed","VJ_sequence_nt_trimmed"),
GEX.features.to.append = c("UMAP_1","UMAP_2","CTLA4", "TOX"),
airr.rearrangements = c("~/cellrangerVDJ/s1/airr_rearrangement.tsv"
,"~/cellrangerVDJ/s2/airr_rearrangement.tsv")
)

#To save a dataframe as .tsv
write.table(airr_dataframe, file = "supplemented_airr_rearrangements.tsv"
, sep= '\t', row.names = FALSE, quote=FALSE)

## End(Not run)

---

**select.top.clone**

Get the index of top ranking clones.

**Description**

Get the index of top ranking clones.
Usage
   select.top.clone(clonotypes, top.n)

Arguments
   clonotypes   The output "clonotypes" dataframe from simulation output.
   top.n        The top n abundant clones to be selected.

Value
   a vector of indexes of top ranking clones

---

**small_vgm**

Small VDJ GEX matrix (VGM) for function testing purposes

**Description**

Small VDJ GEX matrix (VGM) for function testing purposes

**Usage**

```r
small_vgm
```

**Format**

An object of class `list` of length 5.

**References**

R package Platypus : https://doi.org/10.1093/nargab/lqab023

---

**special_v**

special_v a dataframe, of heavy and light chain v gene combination and their probability to be selected for expansion.

**Description**

special_v a dataframe, of heavy and light chain v gene combination and their probability to be selected for expansion.

**Usage**

```r
data("special_v")
```

**Format**

An object of class `data.frame` with 5 rows and 3 columns.
trans_switch_prob_b

The probability for B cell transcriptome states switching. The row names of the matrix are the cell states the cell is switching from, the column names are the cell states the cell is switching to.

**Description**

trans_switch_prob_b The probability for B cell transcriptome states switching. The row names of the matrix are the cell states the cell is switching from, the column names are the cell states the cell is switching to.

**Usage**

```r
data("trans_switch_prob_b")
```

**Format**

A 4x4 matrix. The row and column names are: "GerminalcenterBcell","NaiveBcell","Plasmacell","MemoryBcell". The probability for a cell to switch from "GerminalcenterBcell" to "Plasmacell" is the value at `trans_switch_prob_b[1,3]`.

trans_switch_prob_t

The probability for T cell transcriptome states switching. The row names of the matrix are the cell states the cell is switching from, the column names are the cell states the cell is switching to.

**Description**

trans_switch_prob_t The probability for T cell transcriptome states switching. The row names of the matrix are the cell states the cell is switching from, the column names are the cell states the cell is switching to.

**Usage**

```r
data("trans_switch_prob_t")
```

**Format**

A 7x7 matrix. The row and column names are: "NaiveCd4","ActivatedCd4","MemoryCd4","NaiveCd8","EffectorCd8","MemoryCd8","ExhaustedCd8".
umap.top.highlight  Set idents for top abundant clones in Seurat object, get ready for highlight the top abundant clones in UMAP.

Description
Set idents for top abundant clones in Seurat object, get ready for highlight the top abundant clones in UMAP.

Usage
umap.top.highlight(gex, all.contig.annotations, top.n)

Arguments
- gex: output from get.umap function.
- all.contig.annotations: The output dataframe all_contig_annotations from simulation.
- top.n: The top n abundant clones to be shown in the plot. If missing, all clones will be shown.

Value
a Seurat object ready for highlight the top abundant clones in UMAP

VDJ_abundances  Calculate abundances/counts of specific features for a VDJ dataframe

Description
Calculate abundances/counts of specific features for a VDJ dataframe per an optional specific grouping factor (e.g., 'clonotype_id') and an optional sample factor(e.g., 'sample_id'). Outputs either a count dataframe of the specific feature or a ggplot2 barplot.

Usage
VDJ_abundances(
  VDJ,
  feature.columns,
  proportions,
  specific.features,
  grouping.column,
  max.groups,
  specific.groups,
Arguments

VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.

feature.columns
vector of strings, denoting the columns of the VDJ/VDJ.GEX.matrix[[1]] object from which to extract the unique feature values (for which we will calculate the counts or proportions).

proportions
string, 'absolute' will return the absolute counts, 'group.level.proportions' will return the counts divided by the total number or elements/values in the specific groups (group level proportions), 'sample.level.proportions' will return the counts divided by the total number of elements in the sample.

specific.features
vector of specific feature values (or NULL) for which to calculate counts/proportions, from the specified feature.columns parameter (only works if a single feature column is specified in feature.columns).

grouping.column
string or 'none', represents the column from the VDJ/VDJ.GEX.matrix[[1]] object by which to group counting process. This is usually the 'clonotype_id' column to calculate frequencies at the clonotype level. If 'none', no grouping will be done. For example, if feature.columns='VDJ_cgene' and grouping.column='clonotype_id', we will obtain a count dataframe of the frequencies of each isotype per unique clonotype (per sample if sample.column='sample_id').

max.groups
integer or NULL, the maximum number of groups for which to count features. If NULL, it will count for all groups.

specific.groups
vector of strings (or 'none'), if the counting should be done only for specific groups (e.g., count the frequency of isotype only for clonotypes 1 and 2 if feature.columns='VDJ_cgene', grouping.column='clonotype_id' and specific.groups=c('clonotype1', 'clonotype2'))

sample.column
string, represents the sample column if your VDJ/VDJ.GEX.matrix[[1]] object has multiple samples (usually 'sample_id')

VDJ.VJ.1chain
boolean, if T will remove aberrant cells (more than 1 VDJ of VJ chain), if F it will keep them.

treat.incomplete.groups
string, method of dealing with groups which are missing the features in the feature.columns parameter (e.g., a clonotype which does not have any transcriptomic clusters annotations if feature.columns='transcript_cluster'). 'exclude' -
excludes groups with no cells for the specific features, 'unknown' - sets them as unknown

`treat.incomplete.features`

string, method of dealing with missing feature values (e.g., a clonotype has several NA values for the 'VDJ_cgene' feature column - cells with NA values). 'unknown' - counted as unknown, 'exclude' - excludes completely, 'max.global' - replaces value by max value of that feature across the repertoire, 'max.group' - replaced by the max feature value inside that group, 'proportional' - iteratively assigns the missing values to the known groups, keeping the same proportions.

`combine.features`

boolean - if T and we have two columns in feature.columns, will combine the feature values for each cell in the VDJ object, counting them as a single feature when calculating proportions.

`treat.combined.features`

string, method of dealing with combined features with missing values. 'exclude' will be treated similarly to excluding incomplete feature values (excluding them completely if a single value is missing from the combination), or 'include' and will be treated as a new feature value.

`specific.feature.colors`

named list of specific colors to be used in the final barplots, for each unique feature value in the VDJ object's feature.columns values. For example, if we have a feature column of binders with unique values=c('yes', 'no'), specific.feature.colors=list('yes'='blue', 'no'='red') will color them accordingly.

`output.format`

string, either 'plots' to obtain barplots, 'abundance.df' to obtain the count dataframe, or 'abundance.df.list' to obtain a list of count dataframes, for each sample.

**Value**

Either a count dataframe with the following columns: group=unique group value, e.g., 'clonotype1' if grouping.column='clonotype_id'), sample, group_frequency, unique_feature_values, feature_value_counts, total_feature_names or a barplot of the counts/proportions per feature, per group.

**Examples**

```
VDJ_abundances(VDJ = small_vgm[[1]],
feature.columns='VDJ_cgene',proportions='absolute',
grouping.column='clonotype_id',specific.groups='none',
output.format='abundance.df')
```

---

**VDJ_alpha_beta_Vgene_circos**

*Circos plot for VDJ and VJ pairings*

**Description**

Produces a Circos plot from the VDJ_GEX_matrix[[1]] output. Connects the V-alpha with the corresponding V-beta gene for each clonotype.
Usage

VDJ_alpha_beta_Vgene_circos(
  VDJ,  
  V.or.J, 
  B.or.Tcells, 
  label.threshold, 
  c.threshold, 
  cell.level, 
  clonotype.per.gene.threshold, 
  c.count, 
  platypus.version, 
  filter1H1L
)

Arguments

VDJ
For Platypus platypus.version v3, VDJ_GEX_matrix.output[[1]] has to be supplied. For Platypus V2 the output of the VDJ_GEX_integrate function is necessary.

V.or.J
Determines whether to plot the alpha beta gene pairing of the V or J genes. "V", "J" or "both" as possible inputs. Default: "both".

B.or.Tcells
Specify whether B or T cells are being analyzed ("B" or "T"). If not specified, function attempts to decide based on gene names.

label.threshold
Minimal amount of clonotypes per gene necessary to add a gene label to the sector. Default: 0.

c.threshold
Only clonotypes are considered with a frequency higher than c.threshold. Allows to filter for only highly expanded clonotypes.

cell.level
Logical, defines whether weight of connection should be based on number of clonotypes or number of cells. Default: number of clonotypes.

clonotype.per.gene.threshold
How many clonotypes are required to plot a sector for a gene. Filters the rows and columns of the final adjacency matrix.

c.count
Show clonotype or cell count on Circos plot. Default = T.

platypus.version
Which platypus.version of platypus is being used. Default = "v3".

Value

Returns list of plots. The first n elements contain the circos plot of the n datasets from the VDJ.analyze function. The n+1 element contains a list of the n adjacency matrices for each dataset.
VDJ_analyze

Examples

```r
## Not run:
plots <- VDJ_alpha_beta_Vgene_circos(Platypus::small_vgm[[1]],
, platypus.version="v3")

## End(Not run)
```

---

**Description**

Platypus V2 Processes and organizes the repertoire sequencing data from cellranger vdj and returns a list of dataframes, where each dataframe corresponds to an individual repertoire. The function will return split CDR3 sequences, germline gene information, filter out those clones with either incomplete information or doublets (multiple CDR3 sequences for a given chain). This function should be called once for desired integrated repertoire and transcriptome. For example, if there are 3 VDJ libraries and 3 GEX libraries and the goal is to analyze all three GEX libraries together (e.g. one UMAP/tSNE reduction) this then function should be called one time and the three VDJ directories should be provided as input to the single function call.

**Usage**

```r
VDJ_analyze(
  VDJ.out.directory,
  filter.1HC.1LC,
  clonotype.list,
  contig.list,
  filtered.contigs
)
```

**Arguments**

- **VDJ.out.directory**  
  Character vector with each element containing the path to the output of cellranger vdj runs. Multiple repertoires to be integrated in a single transcriptome should be supplied as multiple elements of the character vector. This can be left blank if supplying the clonotypes and contig files directly as input. This pipeline assumes that the output file names have not been changed from the default 10x settings in the /outs/ folder. This is compatible with B and T cell repertoires (both separately and simultaneously).

- **filter.1HC.1LC**  
  Logical indicating whether only those clones containing 1 VH/TRB and VL/TRA should be maintained for further analysis. Default is set to TRUE, which restricts the analysis to only clones with exactly 1 heavy chain and 1 light chain (or 1 beta + 1 alpha in the case of T cells).
**clonotype.list**  List of dataframes containing clonotyping information for each repertoire. The column names should correspond to the clonotypes.csv file from cellranger vdj output.

**contig.list**  List of dataframes containing the contig information for each repertoire. The column names should correspond to the all_contigs.csv file from cellranger vdj output.

**filtered.contigs**  Logical indicating if the filtered contigs file should be used. TRUE will read VDJ information from only the filtered output of cellranger. FALSE will read the all contigs file from cellranger. Default set to TRUE (filtered output)

**Value**

Returns a list of dataframes where each dataframe corresponds to one input directory. If only one file is supplied, the output list will only contain one element. This output can be supplied as input to other functions including VDJ_per_clone, VDJ_network, VDJ_germline_genes, VDJ_expansion, visualize_clones_GEX, VDJ_phylo, VDJ_clonotype. Germline gene information is based on the majority of cells within each clonotype. For example, if the majority of cells in clonotype1 have the IGHG1 isotype then the entire clonal family will be determined as IGHG1. For a cell-specific investigation, the output of this function can be supplied to the function VDJ_per_clone, which will provide isotype, sequence, germline gene, etc information for each cell within the each clone.

**Examples**

```r
# Not run:
example.vdj.analyze <- VDJ_analyze(
  VDJ.out.directory = "~/path/to/cellranger/vdj/outs/", filter.1HC.1LC = T)

# End(Not run)
```

---

**VDJ_antigen_integrate**  *Integrates antigen-specific information into the VDJ/VDJ.GEX.matrix[[1]] object*

**Description**

Integrate antigen-specific information from a list of antigen dataframes or antigen csv file paths. The antigen data should contain either the clonotypes, cell barcodes, or sequences with the specific column names of the VDJ/VDJ.GEX.matrix[[1]] object. These columns will be used to rematch the binder information at the cell, sequence, or clonotype level into the main VDJ.GEX.matrix[[1]].

**Usage**

`VDJ_antigen_integrate(
  VDJ,
  antigen.data.list,
  VDJ.antigen_integrate`
VDJ_antigen_integrate

antigen.features,
binder.threshold,
VDJ.VJ.1chain,
match.by,
matching.type,
distance.threshold,
sample.id,
aberrant.chosen.sequences,
output.format
)

Arguments

VDJ
VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.

antigen.data.list
list of antigen csv file paths or antigen dataframes for the specific antigen datasets. To ease matching, the column names by which we will match should be the same as the column names in the original VDJ/VDJ.GEX.matrix[[1]] object.

antigen.features
vector of columns of antigen features to be integrated from the antigen csv files into the VDJ/VDJ.GEX.matrix[[1]] object. The vector can also use unique, short-hand names of the columns to add (e.g., ‘affinity’ for ‘octet.affinity.[nM]’).

binder.threshold
list or nested list of threshold values and specific features by which to define binders in the VDJ. For example, if binder.threshold=list(list(‘affinity’, 0.2), list(‘elisa’, 0.8)), we will have two new binder columns: binders_affinity if the values are greater than 0.2, binders_elisa if they are greater than 0.8.

VDJ.VJ.1chain
boolean, if T will remove aberrant cells (more than 1 VDJ of VJ chain), if F it will keep them in the VDJ when matching antigen data.

match.by
string, represents the method by which to match the antigen data and integrate it into the VDJ/VDJ.GEX.matrix[[1]] object. ‘clonotype’ will match by ‘clonotype_id’ (needs to be present in the antigen data), ‘clonotype.v3’ will match by v3 cellranger clonotypes (you need a v3_clonotypes column in the VDJ/VDJ.GEX.matrix[[1]], ‘cdr3.aa’ by VDJ and VJ cdr3s amino acid sequences, ‘cdrh3.aa’ by VDJ cdr3s amino acid sequences, ‘VDJ.VJ.aa’ by full VDJ and VJ aa sequences, ‘VDJ.VJ.nt’ by trimmed nt VDJ and VJ sequences (must run VDJ_call_MIXCR first on the VDJ), ‘cdr3.nt’ by VDJ and VJ cdr3s as nucleotides, ‘cdrh3.nt.’ by VDJ cdr3s as nucleotides, ‘absolut’ will match the VDJ_cdr3s_aa with the CDR3 column in Absolut! datasets.

matching.type
string, either ‘exact’ for exact sequence matching if the match.by parameter is a sequence type, or ‘homology’ for homology matching (matches if the Levenshtein distance is less than the distance.threshold parameter).

distance.threshold
integer, maximum string distance value by which to match sequences in the antigen data and sequences in the VDJ object (to further integrate the antigen data).
sample.id: boolean, if T then will also match by the 'sample_id' column in the antigen dataframes.

aberrant.chosen.sequences: boolean, if T will add a column of the chosen aberrant sequences (which matched a sequence in the antigen data) if matching by sequence (and VDJ.VJ.1chain=F).

output.format: string, 'vgm' - returns the full VDJ object, 'dataframe.per.sample' - list of VDJ dataframes for each sample.

Value

Either the original VDJ dataframe with additional columns of the antigen features integrated, a list of VDJ dataframes per sample.

Examples

```r
## Not run:
VDJ_antigen_integrate_v2(VDJ, antigen.directory.list=antigen.directory.list, antigen.feature=c('elisa', 'affinity'), VDJ.VJ.1chain=T, match.by='clonotype', sample.id=T, output.format='vgm')

## End(Not run)
```

VDJ_assemble_for_PnP

Ab sequence assembly for recombinant PnP expression

Description

Assembles sequences from MIXCR output into inserts for expression in PnP cells. For details check https://doi.org/10.1038/ncomms12535 ! ALWAYS VALIDATE INDIVIDUAL SEQUENCE IN GENEIOUS OR OTHER SOFTWARE BEFORE ORDERING SEQUENCES FOR EXPRESSION ! Check notes on column content below ! Only cells with 1 VDJ and 1 VJ sequence are considered. Warnings are issued if sequences do not pass necessary checks.

Usage

```r
VDJ_assemble_for_PnP(
  VDJ.mixcr.matrix,
  id.column,
  species,
  manual_IgKC,
  manual_2A,
  manual_VDJLeader,
  write.to.disk,
  filename,
  verbose
)
```
**Arguments**

- **VDJ.mixcr.matrix**
  Output dataframe from the VDJ_call_MIXCR function or a dataframe generated using the VDJ_GEX_matrix function and supplemented with MIXCR information (Needed columns: All Framework and CDR sequences)

- **id.column**
  Character. Column name of VDJ.mixcr.matrix to use as ID for the assembled sequences. Defaults to "barcode"

- **species**
  Character. Which IgKC sequence to use. Can be "human" or "mouse". Defaults to "mouse"

- **manual_IgKC**
  Character. Manual overwrite for sequence used as IgKC.

- **manual_2A**
  Character. Manual overwrite for sequence used as Furine 2A site.

- **manual_VDJLeader**
  Character. Manual overwrite for sequence used as VDJ Leader and signal peptide.

- **write.to.disk**
  Boolean. Defaults to TRUE. Whether to save assembled sequences to working directory

- **filename**
  Character. Output file name for .fasta and .csv files if write.to.disk == T. Defaults to PnP_assembled_seqs.fasta/.csv

- **verbose**
  Print runtime message to console. Defaults to FALSE

**Value**

Returns the input VGM matrix with one additional column containing the assembles sequences. If write.to.disk == T writes a CSV containing key columns of the VGM as well as a .FASTA file to the current working director (getwd())! Important notes on column content: 1. The column "seq_length_check" contains either "passed" or "FAILED". If FAILED, this means that at least one of the sequences (e.g. FRL1) was shorter than 9NTs and therefore considered invalid. Please check for missing sequences if you find any warnings 2. The column "seq_codon_check" is deemed "passed" if all CDR and FR input sequences of a cell contain only full codons (i.e. are divisible by 3) 3. The column "PnP_assembled_seqs" contains the assembled sequences / inserts for PnP expression. These should be validated manually in Geneious or other software and can then be ordered to be synthesized. 4. The column "PnP_assembled_annotations" contains a string of annotations for the respective assembled sequence. The structure is | [Sequence element] -> [index (starting from 1) of last nucleotide of the sequence element] ... 5. The column "PnP_assembled_translations" contains the amino acid translation of the full contig that will result from the assembled insert in the backbone PnP vector. Please note: the sequences in the PnP_assembled_translation resulted from pasting the VJ leader sequence (contained in the PnP vector backbone), the PnP_assembled_seqs (The insert itself) and a surrogate stop codon ATAA. If correct, the translation should only contain one * (stop codon) at the very end. For reference: VJLeader sequence: ATGGATTTTCAAGGT-GCAGATTTTTCAGCTTCCGCTAAATCGGCTTCCAGTTATAATGTCGCCG 6. The column "seq_VJCDR3_check" is deemed "passed" if the translated sequence of the input VJ CDR3 is found in the translated assembled sequence. If this test fails, there is likely an issue with the VJ segment 7. The column "seq_Fur2A_check" is deemed "passed" if correct AA sequence of the 2A site is found in the translated assembled sequence. If this test fails, and the seq_VJCDR3_check was passed, there is likely an issue at the border between VJ and IgKC/2A sequences 8. The column "seq_VDJCDR3_check" is deemed "passed" if the translated sequence of the input VDJ CDR3
is found in the translated assembled sequence. 9. The column "seq_splicessite_check" is deemed passed if the last 6 nucleotides of the assembled sequence are one of the following: "TCCTCA", "TCTTCA", "TCGTCA", "TCATCA".

Examples

```r
## Not run:
VGM_with_PnP_seq <- VDJ_assemble_for_PnP(VDJ.mixcr.matrix = VDJ_call_MIXCR.output,
  id.column = "barcode", species = "mouse", manual_IgKC = "none", manual_2A = "none",
  manual_VDJLeader = "none", write.to.disk = TRUE, filename = "PnP_seq_example")
## End(Not run)
```

### VDJ_bulk_to_vgm

**Utility function for bulk data to standard Platypus format conversion**

**Description**

The **VDJ_bulk_to_vgm** function converts bulk output files from MIXCR or MAF into a vgm-format compatible with most downstream Platypus functions used for VDJ repertoire analysis.

**Usage**

```r
VDJ_bulk_to_vgm(
  VDJ.bulk.out.directory.list,
  input.type, 
  integrate.MIXCR.output,
  vgm.expanded,
  clone.strategy,
  group.id,
  cell.type,
  batches,
  best.match.only
)
```

**Arguments**

```
VDJ.bulk.out.directory.list
  List containing paths to bulk VDJ output files from MIXCR or MAF.

input.type
  Character vector. Defaults to "MIXCR". "MIXCR" and "MAF" are supported.

integrate.MIXCR.output
  Boolean. Defaults to TRUE. Whether to include in the VGM output additional MiXCR (49-78) columns.

vgm.expanded
  Boolean. Defaults to TRUE. Whether to include vgm[[9]] in the output list, where vgm[[9]] is the expanded version of vgm[[1]] having 1 line per read. For some Platypus functions, only vgm[[9]] (and not vgm[[1]]) may be compatible.
```
clone.strategy Character vector to specify the clonotyping strategy. Defaults to "cdr3.aa". Note that MIXCR input comes with clonotypes already assigned, and therefore clone.strategy should be specified only when the user wants to change the clonotyping strategy, and if no clone.strategy is provided, re-clonotyping will not be performed. Meanwhile, MAF inputs do not come with the clonotypes pre-assigned. Hence, if no clone.strategy is specified, "cdr3.aa" will be used as the default clonotyping strategy. The clonotyping strategies available in this function are: "cdr3.aa", "VDJJ.VJJ", "VDJJ.VJJ.cdr3length".

group.id Numeric vector. Defaults to NA. The user can specify to which group does each file belong to (e.g. a group could correspond to some specific treatment). The length of this numeric vector should match the number of samples in the VDJ.bulk.out.directory.list input.

cell.type Character vector. Defaults to NA. Cell type (e.g., "Bcell") of the MIXCR or MAF file that is provided as input.

batches Numeric vector. Defaults to NA. An additional grouping parameter that can be specified by the user. The length of this numeric vector should match the number of samples in the VDJ.bulk.out.directory.list input.

best.match.only Boolean. Whether only the highest scoring gene (V,J,D,C gene should) should be included in the output, or all matching genes in MIXCR should be included (MAF outputs: for the same read we can only have one possible V,J,D or C gene). Defaults to TRUE.

Value

A VGM object (vgm.bulk.list). vgm.bulk.list[[1]]: each line correspond to a clonotype. vgm.bulk.list[[9]] (if vgm.expanded==TRUE): each line correspond to a read. The other (2-8) entries of the list are left empty for compatibility with Platypus functions.

Examples

```r
## Not run:
Run from local directory using MIXCR/MAF bulk VDJ-repertoire files as inputs:
VDJ.bulk.out.directory.list <- list()
VDJ.bulk.out.directory.list[[1]] <- c("~/MIXCR_vdj_cdr3_clonotyping/C4.txt")
VDJ.bulk.out.directory.list[[2]] <- c("~/MIXCR_vdj_cdr3_clonotyping/C6.txt")
bulk.vgm.MIXCR <- VDJ_bulk_to_vgm(VDJ.bulk.out.directory.list = VDJ.bulk.out.directory.list,
input.type = "MIXCR",
integrate.MIXCR.output = TRUE,
group.id = c(1,2),
cell.type = "Bcells",
batches = c(1,1),
vgm.expanded = TRUE,
best.match.only = FALSE)

To re-clonotype MIXCR samples based on e.g., the CDR3 a.a. sequence:
bulk.vgm.MIXCR <- VDJ_bulk_to_vgm(VDJ.bulk.out.directory.list = VDJ.bulk.out.directory.list,
input.type = "MIXCR",
integrate.MIXCR.output = TRUE,
```
group.id = c(1,2),
cell.type = "Bcells",
batches = c(1,1),
vgm.expanded = TRUE,
best.match.only = FALSE,
clone.strategy = "cdr3.aa"

## End(Not run)

**VDJ_call_MIXCR**  
*MiXCR wrapper for Platypus V3 VDJ object*

### Description

Extracts information on the VDJRegion level using MiXCR on WINDOWS, MAC and UNIX systems for input from both Platypus v2 (VDJ_per_clone) or v3 (Output of VDJ_GEX_matrix) This function assumes the user can run an executable instance of MiXCR and is eligible to use MiXCR as determined by license agreements. ! FOR WINDOWS USERS THE EXECUTABLE MIXCR.JAR HAS TO PRESENT IN THE CURRENT WORKING DIRECTORY ! The VDJRegion corresponds to the recombined heavy and light chain loci starting from framework region 1 (FR1) and extending to frame work region 4 (FR4). This can be useful for extracting full-length sequences ready to clone and further calculating somatic hypermutation occurrences.

### Usage

```
VDJ_call_MIXCR(
  VDJ,
  operating.system,
  mixcr.directory,
  species,
  simplify,
  platypus.version
)
```

### Arguments

**VDJ**  
For platypus.version = "v2" the output from the VDJ_per_clone function. This object should have information regarding the contigs and clonotype_ids for each cell. For platypus.version = "v3" the VDJ dataframe output of the VDJ_GEX_matrix function (VDJ.GEX.matri.output[[1]])

**operating.system**  
Can be either "Windows", "Darwin" (for MAC) or "Linux". If left empty this is detected automatically

**mixcr.directory**  
The directory containing an executable version of MiXCR. FOR WINDOWS USERS THIS IS SET TO THE CURRENT WORKING DIRECTORY (please paste the content of the MIXCR folder after unzipping into your working directory. Make sure, that mixcr.jar is not within any subfolders.)
**species**

Either "mmu" for mouse or "hsa" for human. These use the default germline genes for both species contained in MIXCR. Defaults to "hsa"

**simplify**

Only relevant when platypus.version = "v3". Boolean. Defaults to TRUE. If FALSE the full MIXCR output and computed SHM column is appended to the VDJ If TRUE only the framework and CDR3 region columns and computed SHM column is appended. To discriminate between VDJ and VJ chains, prefixes are added to all MIXCR output columns

**platypus.version**

Character. Defaults to "v3". Can be "v2" or "v3" dependent on the input format

### Value

For platypus.version = "v3" returns input VDJ dataframe supplemented with MIXCR output information. For platypus.version = "v2" returns a nested list containing VDJRegion information as determined by MIXCR. The outer list corresponds to the individual repertoires in the same structure as the input VDJ_per.clone. The inner list corresponds to each clonal family, as determined by either the VDJ_clonotype function or the default nucleotide clonotyping produced by cellranger. Each element in the inner list corresponds to a dataframe containing repertoire information such as isotype, CDR sequences, mean number of UMIs. This output can be supplied to further package functions such as VDJ_extract_sequences and VDJ_GEX_integrate.

### See Also

VDJ_extract_sequences

### Examples

```r
## Not run:
# For platypus version 2
VDJ_call_MIXCR(VDJ = VDJ_per.clone.output, 
mixcr.directory = "~/Downloads/mixcr-3.0.12/mixcr", species = "mmu")

# For platypus version 3 on a Windows system
VDJ_call_MIXCR(VDJ = VDJ_GEX_matrix.output[[1]], 
mixcr.directory = "WILL BE SET TO CURRENT WORKING DIRECTORY", 
species = "mmu", platypus.version = "v3", simplify = TRUE)
## End(Not run)
```

---

**VDJ_call_recon**

Calls the Kaplinsky/RECON tool

### Description

Calls the Kaplinsky/RECON tool on the VDJ/VDJ.GEX.matrix[[1]] object to infer the parent distribution of clonotypes and estimate their diversity. Outputs either a dataframe of the resulting means and weights of the RECON clonotype parent distribution estimation or a plot of the original clonotype distribution along resampled values from the reconstructed parent distribution.
Usage

VDJ_call_recon(
    VDJ,
    recon.directory,
    sample.id,
    clone.list,
    max.clones,
    size.threshold,
    resample,
    plot.results,
    max.clone.size,
    reticulate,
    output.format,
    operating.system
)

Arguments

VDJ | VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.
recon.directory | directory containing recon executable. Defaults to working directory/Recon
sample.id | boolean - if F, clonotypes will be considered at a global level, irrespective of samples.
clone.list | list - if empty, RECON will be used to estimate the diversity of all clonotypes, else it will only consider the specified clonotypes.
max.clones | integer or 'all' - maximum number of clones ot be considered for the RECON estimation. If 'all', will consider all clonotypes.
size.threshold | integer - the size threshold parameter for the RECON tool, as specified by the '-t' parameter.
resample | boolean - if T, will also perform and output a resample of the clonotype frequencies/sizes from the inferred parent distribution.
plot.results | boolean - if T, will save a pdf of the resampled clonotype frequencies/sizes from the RECON-inferred distribution along the original frequencies.
max.clone.size | integer - the maximum size of clones to be considered in the resulting plot (maximum number of elements on the x axis).
reticulate | boolean - if T, will create a new environment to install python and run the RECON tool, else, your environment must have a python version compatible with RECON installed.
output.format | string - 'vgm' will append the means and weights of the RECON-inferred distribution to the VDJ/VDJ.GEX.matrix[[1]] object, 'recon' will output a new dataframe of these weights, 'plots' will output the ggplot2 objects (if plot.results=T).
operating.system | string - operating system on which RECON will be run. 'Windows' for Windows, 'Linux' for Linux, 'Darwin' for MacOS.
Value

The resulting means and weights of the RECON-inferred distribution as a separate dataframe or appended to the VDJ, or a plot of resampled clonotype sizes from the inferred distribution vs original sizes/frequencies.

Examples

## Not run:
VDJ_call_recon(VDJ, recon.directory= './Recon',
max.clones='all', sample.id=T, resample=F,
plot.results=T, output.format='vgm',
reticulate=T, operating.system='Darwin')

## End(Not run)

VDJ_circos

Internal utility for circos functions

Description

Plots a Circos diagram from an adjacency matrix. Uses the Circlize chordDiagram function. Is called by VDJ_clonotype_clusters_circos(), VDJ_alpha_beta_Vgene_circos() and VDJ_VJ_usage_circos() functions or works on its own when supplied with an adjacency matrix.

Usage

VDJ_circos(Adj_matrix, group, grid.col, label.threshold, axis, c.count)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adj_matrix</td>
<td>Adjacency matrix to be plotted. Rownames and Colnames correspond to genes to be matched and entries determine the weight of the connection between the genes (e.g. number of clonotypes expressing these two genes).</td>
</tr>
<tr>
<td>group</td>
<td>Named list of genes, with list elements corresponding to group-names, and element names being the gene-names. Is generated by VDJ_VJ_usage and VDJ_alpha_beta_Vgene_circos.</td>
</tr>
<tr>
<td>grid.col</td>
<td>Named list of genes, with list elements corresponding to color and element names being gene-names. If not supplied it is generated randomly within the function. Is also generated by VDJ_VJ_usage and VDJ_alpha_beta_Vgene_circos.</td>
</tr>
<tr>
<td>label.threshold</td>
<td>Genes are only labeled if the count is larger then the label.threshold. By default all label.threshold = 0 (all genes are labeled).</td>
</tr>
<tr>
<td>axis</td>
<td>Option to choose the count axis for each gene. &quot;default&quot;, &quot;percent&quot; or &quot;max&quot; possible. Default: &quot;max&quot;.</td>
</tr>
<tr>
<td>c.count</td>
<td>Show clonotype or cell count on Circos plot.</td>
</tr>
</tbody>
</table>
Value

Returns the Circos plot from input of other functions. Do not run as standalone

Examples

```r
## Not run:
VDJ_circos() #Do not run as standalone. Called by other circos functions
## End(Not run)
```

**VDJ_clonal_donut**

Circular VDJ expansion plots

Description

Generate circular plots of clonal expansion per repertoire directly from the VDJ matrix of the VDJ_GEX_matrix function

Usage

```r
VDJ_clonal_donut(
  VDJ,            # VDJ dataframe generated using the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]]).
  counts.to.use,  # Plots will be made by sample and using the clonal frequencies specified by
  label.size,     # counts.to.use
  not.expanded.label.vjust,
  not.expanded.label.hjust,
  total.label.vjust,
  total.label.hjust,
  expanded.colors,
  non.expanded.color
)
```

Arguments

- **VDJ**: VDJ dataframe generated using the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]]).
  Plots will be made by sample and using the clonal frequencies specified by counts.to.use

- **counts.to.use**: How to count clonotypes and cells. A column name of the VDJ matrix containing clonotype IDs. This defaults to "clonotype_id_10x", which reflects clonotypes by Cellranger in an unaltered VGM. To use counts from the VDJ_clonotype_v3 function set this parameter to the relevant column e.g. "clonotype_id_cdr.aa" or "global_clonotype_id_cdr.aa" are two examples.

- **label.size**: Size of text labels. All parameters below are purely for graphical purposes and optional. If necessary changes should be made in small (0.1) increments. ! It is recommended to optimize these ONLY once a format for saving the plot is set.

- **not.expanded.label.vjust**: Numeric. Regulates the vertical position of the label for non expanded cells
not.expanded.label.hjust
  Numeric. Regulates the horizontal position of the label for non expanded cells

total.label.vjust
  Numeric. Regulates the vertical position of the center label

total.label.hjust
  Numeric. Regulates the horizontal position of the center label

expanded.colors
  Character vector. Colors to use for expanded clones. Should be more than 3 for better visibility. Defaults to a "darkorchid3"-based palette.

non.expanded.color
  Character. Color to use for non expanded clones. Defaults to "black"

Value

Returns a list of circular plots showing proportions of expanded clones and non-expanded clones. One plot is generated for each sample in the sample_id column

Examples

VDJ_clonal_donut(VDJ = Platypus::small_vgm[[1]])

VDJ_clonal_expansion

Flexible wrapper for clonal expansion barplots by isotype, GEX cluster etc.

Description

Clonal frequency plot displaying clonal expansion for either T and B cells with Platypus v3 input. Only available for Platypus "v3" available. For v2 plotting of B cell clonotype expansion and isotypes please refer to VDJ_isotypes_per_clone.

Usage

VDJ_clonal_expansion(
  VDJ,
  celltype,
  clones,
  subtypes,
  isotypes.to.plot,
  species,
  treat.incomplete.clones,
  treat.incomplete.cells,
  group.by,
  color.by,
  variant.plot
)
Arguments

**VDJ**  
VDJ dataframe generated using the VDJ\_GEX\_matrix function (VDJ\_GEX\_matrix\_output[[1]])

**celltype**  
Character. Either "Tcells" or "Bcells". If set to Tcells bars will not be colored by default and the parameters treat\_incomplete\_cells, treat\_incomplete\_clones, subtypes and species are ignored. The color.by and group.by arguments work identically for both celltypes. If none provided it will detect this param from the celltype column.

**clones**  
numeric value indicating the number of clones to be considered for the clonal expansion plot. Default value is 50. For a standard plot more than 50 is discouraged. When showing only one - possibly rare - isotype via isotypes\_to\_plot it may be useful to set this number higher (e.g. 100-200)

**subtypes**  
Logical indicating whether to display isotype subtypes or not.

**isotypes\_to\_plot**  
Character vector. Defaults to "all". This can be set to any number of specific Isotypes, that are to be shown exclusively. For example, to show only clones containing IgG, input "IGHG". If only wanting to check clones with IgA and IgD input c("IGHA","IGHD"). Works equally if subtypes are set to TRUE. Is ignored if color.by is not set to "isotype"

**species**  
Character indicating whether the samples are from "Mouse" or "Human". Default is "Human".

**treat\_incomplete\_clones**  
Character indicating how to proceed with clonotypes lacking a VDJC (in other words, no cell within the clonotype has a VDJC). "exclude" removes these clonotypes from the analysis. "include" keeps these clonotypes in the analysis. In the plot they will appear has having an unknown isotype.

**treat\_incomplete\_cells**  
Character indicating how to proceed with cells assigned to a clonotype but missing a VDJC. "proportional" to fill in the VDJ isotype according to the proportions present in of clonotype (in case present proportions are not replicable in the total number of cells e.g. 1/3 in 10 cells, values are rounded to the next full integer and if the new counts exceed the total number of cells, 1 is subtracted from the isotype of highest frequency. If the number is below the number of cell, 1 is added to the isotype with lowest frequency to preserve diversity), "exclude" to exclude them from analysis and rank clonotypes only by the number of cells with a heavy chain. This ranking may deviate from the frequency column in the clonotype table. CAVE: if treat\_incomplete\_cells is set to "exclude", clonotypes lacking a VDJC entirely will be removed from the analysis. This results in a similar but not identical output as when treat\_incomplete\_clones is set to true. The two parameters are thereby non-redundant.

**group\_by**  
Character. Defaults to "sample_id". Column name of VDJ to split VDJ by. For each unique entry in that column a plot will be generated. Therefore plots can be generated by sample_id, group_id or any other metadata item. To get plots for the whole repertoire set to "none"

**color\_by**  
Character. Defaults to "isotype". If set to "isotype" bars are colored by the respective IgH chain or in grey for T cells. This can alternatively be set to any column name of the VDJ. This allows coloring clones by their V\_gene usage or by GEX clusters
variant.plot Logical indicating whether to plot the output showing the variants or not.

Value

Returns a nested list. out[[1]] are plots out[[2]] are raw datatables containing also barcode and CDR3 information

Examples

# Standard B cell plot for platypus version v3
# Will generate one plot per sample (from sample_id column)
clonal_out <- VDJ_clonal_expansion(VDJ = Platypus::small_vgm[[1]],
  celltype = "Bcells", clones = 30, subtypes = FALSE, species = "Mouse"
  , treat.incomplete.clones = "exclude"
  , treat.incomplete.cells = "proportional"
)

# Regrouped and recolored plot in v3
# Will generate a plot for each sample.
# Bars are filled by the sample with the highest proportion of cells in a given clonotype
clonal_out <- VDJ_clonal_expansion(VDJ = Platypus::small_vgm[[1]],
  celltype = "Bcells", clones = 30, subtypes = FALSE, species = "Mouse"
  , treat.incomplete.clones = "exclude"
  , treat.incomplete.cells = "proportional"
  , color.by = "seurat_clusters") # change grouping with group.by = "column name"
clonal_out[[1]] # list of plots
clonal_out[[2]] # list of source dataframes

# T cell plot with recoloring by vgene
# VDJ_clonal_expansion(VDJ = Platypus::small_vgm[[1]]
# ,celltype = "Tcells", clones = 30, group.by = "sample_id"
# , color.by = "VDJ_vgene")

# Plotting only IgD clones. Increased the value for clones to scan more of the dataset
# VDJ_clonal_expansion(VDJ = Platypus::small_vgm[[1]]
# ,celltype = "Bcells", clones = 150, subtypes = FALSE
# , species = "Mouse", treat.incomplete.clones = "include"
# , treat.incomplete.cells = "proportional", isotypes.to.plot = "IGHD")

# Plotting only clones containing cells with the IGHD2c isotype (For murine data only!)
# VDJ_clonal_expansion(VDJ = Platypus::small_vgm[[1]]
# ,celltype = "Bcells", clones = 150, subtypes = TRUE, species = "Mouse"
# , treat.incomplete.clones = "exclude"
# , treat.incomplete.cells = "proportional", isotypes.to.plot = "IGHD2c")
**Description**

Wraps the VDJ_abundances function and output a barplot of clonotypes ranked by expansion (x axis) with counts of the specific feature values per clonotype (y axis). For a more in-depth configuration of the barplots (e.g., including clonotypes with missing features, different strategies for NA values, etc.), use the VDJ_abundances function with output.format='plots'.

**Usage**

```r
VDJ_clonal_expansion_abundances(
  VDJ,
  features,
  count.level,
  max.clonotypes,
  rank.clonotypes,
  specific.feat.colors
)
```

**Arguments**

- **VDJ**
  VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.
- **features**
  string or vector of strings, denoting the columns of the VDJ/VDJ.GEX.matrix[[1]] object from which to extract the unique feature values.
- **count.level**
  string, 'absolute' will return the absolute counts, 'group.level.proportions' will return the counts divided by the total number or elements/values in the specific groups (group level proportions), 'sample.level.proportions' will return the counts divided by the total number of elements in the sample.
- **max.clonotypes**
  integer or NULL, the maximum number of clonotypes for which to count features. If NULL, it will count for all clonotypes.
- **rank.clonotypes**
  boolean, if T - clonotypes will be ranked and order according to their expansion.
- **specific.feat.colors**
  named list (or NULL) of specific colors to be used in the final barplots.

**Value**

Either a count dataframe with the following columns: group(=unique group value, e.g., 'clonotype1' if grouping.column='clonotype_id'), sample, group_frequency, unique_feature_values, feature_value_counts, total_feature_names or a barplot of the counts/proportions per feature, per group.

**Examples**

```r
## Not run:
VDJ_clonal_expansion_abundances(VDJ = small_vgm[[1]],
                                 features='VDJ_cgene',count.level='absolute',
                                 max.clonotypes=30, rank.clonotypes=T, specific.feat.colors=NULL)

## End(Not run)
```
**Description**

Only Platypus V2 Organizes and extracts full-length sequences for clonal lineage inference. The output sequence can either contain the germline sequence as determined by cellranger or can just contain the sequences contained in each clonal family.

**Usage**

```r
VDJ_clonal_lineages(  
  VDJ,  
  VDJ_extract_germline.output,  
  as.nucleotide,  
  with.germline,  
  platypus.version  
)
```

**Arguments**

- **VDJ**
  For platypus v2 the output of the call_MIXCR function containing the full-length VDJRegion sequences. For v3 the VDJ matrix output of the VDJ_GEX_matrix function ran with trim.and.align = TRUE. (VDJ_GEX_matrix.output[[1]])

- **VDJ_extract_germline.output**
  The output from the VDJ_extract_germline function. This should have the germline information. This needs to be supplied if the with.germline argument is set to true.

- **as.nucleotide**
  Logical determining whether the full-length VDJRegion sequence should use nucleotide sequence. TRUE indicates nucleotide sequences and FALSE will extract amino acid sequences.

- **with.germline**
  Logical determining whether the germline sequence as determined by cellranger should be included in the output list of sequences. If so, the germline will be added to the last row of each dataframe object.

- **platypus.version**
  Default is "v3".

**Value**

returns a list containing the sequences for each clonal family as determined by the input clonotyping strategy to call_MIXCR and VDJ_extract_germline. The outer list corresponds to distinct repertoires supplied to the call_MIXCR function (e.g. VDJ.clonal.lineage.output[[i]][[j]] will contain a dataframe of the j'th clone in the i'th repertoire)
Examples

```r
## Not run:
clonal_lineages <- VDJ_clonal_lineages(VDJ=call_MIXCR_output,
VDJ_extract_germline.output=VDJ_extract_germline_output,as.nucleotide=F,with.germline=T)
## End(Not run)
```

---

**VDJ_clonotype**

* Deprecated Platypus V2 clonotyping wrapper *

**Description**

Deprecated function for Platypus V2 with options for Platypus V3. For revised hierarchical clonotyping please use VDJ_clonotype_v3() Returns a list of clonotype dataframes following additional clonotyping. This function works best following filtering to ensure that each clone only has one heavy chain and one light chain.

**Usage**

```r
VDJ_clonotype(
  VDJ,
  clone.strategy,
  homology.threshold,
  hierarchical,
  VDJ.VJ.1chain,
  global.clonotype,
  output.format,
  platypus.version
)
```

**Arguments**

- **VDJ**
  For platypus v2 output from VDJ_analyze function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire. For platypus v3 VDJ output from the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]])

- **clone.strategy**
  (Updated keywords, previous format is also functional) String describing the clonotyping strategy. Possible options include 'cdr3.nt', 'cdr3.aa', 'VDJJ.VJJ', 'VDJJ.VJJ.cdr3lengths', 'VDJJ.VJJ.cdr3length.VDJCDR3.homology', 'cdr3.homology', or 'VDJcdcr3.homology'.
  'cdr3.aa' will convert the default cell ranger clonotyping to amino acid based.
  'Hvj.Lvj' groups B cells with identical germline genes (V and J segments for both heavy chain and light chain. Those arguments including 'CDR3length' will group all sequences with identical CDRH3 and CDRL3 sequence lengths. Those arguments including 'CDR3.homology' will additionally impose a homology requirement for CDRH3 and CDRL3 sequences. 'CDR3.homology', or 'CDRH3.homology' will group sequences based on homology only (either of
the whole CDR3 sequence or of the CDRH3 sequence respectively). All homology calculations are performed on the amino acid level.

**homology.threshold**

Numeric value between 0 and 1 corresponding to the homology threshold for the clone.strategy arguments that require a homology threshold. Default value is set to 70 percent sequence homology. For 70 percent homology, 0.3 should be supplied as input.

**hierarchical**

Boolean. Defaults to FALSE. This is an extension specifically for cells with aberrant numbers of chains (i.e. 0VDJ 1VJ, 1VDJ 0VJ, 0VDJ 2VJ, 2VDJ 0VJ). Cells with 2VDJ 2VJ are filtered out as these are most likely doublets. Aberrant cells are clonotyped hierarchically in post, following this procedure: 1. define clonotypes classically with all cells containing exactly 1VDJ 1VJ chains. 2. For cells with only a single chain (either VDJ or VJ), check if any clone exists, which matches the clonotyping criteria for this chain. If true, add this cell to that clone. If false, create a new clone containing that cell. In case that more than 1 existing clone matches the aberrant cell, the cell is assigned to the most frequent existing clone. Two reasons are behind this decision: 2.1. The aberrant cells is numerically more likely to be a part of the more frequent existing clone. 2.2 In case of a wrong assignment, the effect of the error is lower, if an already expanded clone is increase by one count, rather than a existing non-expanded clone being assigned a second entry and thereby resulting as expanded. 3. For cells with 3 chains, verify the clonotyping criteria on both combinations of chains (i.e. VDJ1 - VJ1, VDJ2-VJ1 in case of a cell with 2VDJ 1VJ).

**VDJ.VJ.1chain**

Logical specifying whether cells with multiple VDJ and VJ chains should be removed from the clonotyping. Can be either T or F for those definitions not requiring germline genes or homology thresholds, as calculating the later is difficult when multiple chains are present.

**global.clonotype**

Logical specifying whether clonotyping should occur across samples or only within a single sample.

**output.format**

String specifies function output format. Options are "vgm" (default), "dataframe.per.sample", "clone.level.dataframes", or "phylo.dataframe". "vgm" will update the existing $clonotype_id column of the input vgm, which is the output from VDJ_GEX_matrix. "dataframe.per.sample" will return a list of VDJ dataframes, where each dataframe contains the cell-level information for a given sample. "clone.level.dataframes" will convert the per.cell matrix to a clonal dataframe, in which cells of the same clone will be merged into a single row. "dataframe.per.clone" will generate nested lists of dataframes, where each dataframe contains cell-level information of a given clone.

**platypus.version**

Default is "v3". To use the output of VDJ_GEX_matrix function, one should change this argument to "v3".

**Value**

Returns a list of clonotype dataframes where each list element matches the repertoire index in the input clonotype.list object. The dataframes will be updated with clonal frequencies based on the new clonotying definition.
Examples

reclonotyped_vgm <- VDJ_clonotype(VDJ=Platypus::small_vgm[[1]],
clone.strategy="VDJJ.VJJ",
homology.threshold=".3", platypus.version = "v3")

---

VDJ_clonotype_clusters_circos

Circos plot for clonotype - GEX cluster pairings

---

Description

Makes a Circos plot from the VDJ_GEX_integrate output. Connects the clonotypes with the corresponding clusters.

Usage

VDJ_clonotype_clusters_circos(
  VDJ,
  topX,
  label.threshold,
  axis,
  c.count,
  n_cluster,
  platypus.version
)

Arguments

VDJ       The output of the VDJ_GEX_integrate function (Platypus platypus.version v2). A list of data frames for each sample containing the clonotype information and cluster membership information. For Platypus platypus.version v3, the VDJ output of the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]]) has to be supplied.
topX      Plots only the top X most expanded clonotypes. By default all clonotypes are shown.
label.threshold Minimal amount of clonotypes per gene neccessary to add a gene label to the sector. Default: 0.
axis      Character. Defaults to "max". Passed to VDJ_circos
c.count   Show clonotype or cell count on Circos plot. Default = T.
n_cluster Integer. No default.
platypus.version Which platypus.version of platypus is being used. Default = "v3".
Value

Returns list of plots. The first n elements contain the circos plot of the n datasets from the VDJ.analyze function. The n+1 element contains a list of the n adjacency matrices for each dataset.

Examples

```r
# Platypus version 3
# prepare the small toy dataset
small_vgm <- Platypus::small_vgm
small_vgm[[1]]$clonotype_id_10x <- "clonotype1"
small_vgm[[1]]$clonotype_frequency <- nrow(small_vgm[[1]])
VDJ_clonotype_clusters_circos(small_vgm[[1]], topX=100, label.threshold=5
, platypus.version = "v3", n_cluster = 2)
```

Description

Updated clonotyping function based on implications for cells with different chain numbers than 1 VDJ 1 VJ chains.

This function offers two types of hierarchical clonotyping. The hierarchical option "single.chains" only merges cell with a single chain into clonotypes composed of cells with 1 VDJ 1 VJ chain. This is based on the assumption, that during mRNA capture and RT-PCR in GEMs, not all transcripts are captured and therefore cells may result missing a VDJ or VJ chain. The hierarchical option "double.and.single.chains" is based on the assumption, that cells with 1 VDJ and 2 VJ chains exist. For a review of the work concerning such cells as well as 2 VDJ 1 VJ cells please consult: https://doi.org/10.4049/jimmunol.1800904. The user may set a threshold of occurrence number above which cells with 1 VDJ 2 VJ chains are considered to be true and other cells with 1 VDJ 1 VJ, 1 VDJ 0 VJ and 0 VDJ 1 VDJ may be merged into the same clonotype by the strategy provided by the user. Cells with 2 VDJ chains are currently not considered in this process, as these are reported to be much rarer and, if appearing in the dataset are more likely to be doublets. We advice the user to carefully examine the output after hierarchical clonotyping before proceeding with further analysis. We thank Prof. Vijayanand as well as Vicente and Emmanuel from his lab for the discussions that have helped with improving the original Platypus clonotyping strategy.

Usage

```r
VDJ_clonotype_v3_v3(
    VDJ,
    clone.strategy, homology.threshold,
    hierarchical, triple.chain.count.threshold,
    global.clonotype, VDJ.VJ.1chain,
```
output.format, platypus.version

Arguments

VDJ  For platypus v2 output from VDJ_analyze function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire. For platypus v3 VDJ output from the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]])

clonotype.strategy  (Updated keywords, previous format is also functional) String describing the clonotyping strategy. Possible options are 10x.default, cdr3.nt, cdr3.aa, VDJ.JJ, VDJ.JJ.cdr3length, VDJ.JJ.cdr3length.cdr3.homology, VDJ.JJ.cdr3length.cdr3.homology, VDJ.JJ.cdr3length.VDJ.cdr3.homology, cdr3.homology, VDJ.cdr3.homology. cdr3.aa will convert the default cell ranger clonotyping to amino acid based. 'VDJJ.JJ' groups B cells with identical germline genes (V and J segments for both heavy chain and light chain. Those arguments including 'cdr3length' will group all sequences with identical VDJ and VJ CDR3 sequence lengths. Those arguments including 'cdr3.homology' will additionally impose a homology requirement for CDRH3 and CDR3 sequences.'CDR3.homology', or 'CDRH3.homology' will group sequences based on homology only (either of the whole CDR3 sequence or of the VDJ CDR3 sequence respectively). All homology calculations are performed on the amino acid level.

homology.threshold  Numeric value between 0 and 1 corresponding to the homology threshold for the clone.strategy arguments that require a homology threshold. Default value is set to 70 percent sequence homology. For 70 percent homology, 0.3 should be supplied as input.

hierarchical  Character. Defaults to "none". This is an extension specifically for cells with aberrant numbers of chains (i.e. 0VDJ 1VJ, 1VDJ 0VJ, 0VDJ 2VJ, 2VDJ 0VJ). Cells with 2VDJ 2VJ are filtered out as these are most likely doublets. If set to "none" aberrant cells are assigned to their own clonotypes. If set to "single.chains" the function will proceed in two steps: 0. Prefiltering: cells with 2 VDJ 2 VJ chains as well as cells with 2 VDJ and any number of VJ chains are filtered out. 1. Define clonotypes classically with all cells containing exactly 1VDJ 1VJ chains. 2. For cells with only a single chain (either VDJ or VJ), check if any clone exists, which matches the clonotyping criteria for this chain. If true, add this cell to that clone. If false, create a new clone containing that cell. In case that more than 1 existing clone matches the aberrant cell, the cell is assigned to the most frequent existing clone. Two reasons are behind this decision: 2.1. The aberrant cells is numerically more likely to be a part of the more frequent existing clone. 2.2 In case of a wrong assignment, the effect of the error is lower, if an already expanded clone is increase by one count, rather than a existing non-expanded clone being assigned a second entry and thereby resulting as expanded. Cells If set to "double.and.single.chains" the function will proceed as if set to "single.chains" but include two more steps 3. Check the frequency of each cell 1 VDJ 2 VJ chain exact clone (by exact nucleotide CDR3 matching). Only if this count exceeds the triple.chain.count.threshold, the clone
is used as a "hub clone". This protects from merging clonotypes on the basis of rare doublets. 4. Merge existing clonotypes into the 1 VDJ 2 VJ clonotypes as they match with the assumption that e.g. a cell with 1 VDJ 1 VJ is part of that same clonotype, but missing a VJ chain due to stochastical sampling.

**triple.chain.count.threshold**

Minimal occurrence frequency for any cell with more than 2 of either VDJ or VJ chain (e.g. 2 VDJ 1 VJ) for it to be considered as a trustworthy clone for hierarchical clonotyping ONLY when hierarchical is set to "double.and.single.chains". Defaults to 3, meaning that, an exact combination of three chains needs to appear in the dataset at least 3 times for it to be considered as a clone, into which other cells are merged. (For the counting of exact combination of chains CDR3 nucleotide string matching is used, even if clonotyping by homology)

**global.clonotype**

Logical specifying whether clonotyping should occur across samples or only within a single sample (grouping via sample_id column).

**VDJ.VJ.1chain**

Logical specifying whether cells other than once with 1 VDJ and 1 VJ chains should be considered.

**output.format**

Parameter output.format is deprecated. If non VGM-style output is required please refer to the function VDJ_clonotype. Output is VGM style VDJ by cell dataframe.

**platypus.version**

Only "v3" available

**Value**

Returns a VGM[[1]]-type dataframe. The columns clonotype_id and clonotype_frequency are updated with the new clonotyping strategy. They represent the "active strategy" that downstream functions will use. Furthermore extra columns are added with clonotyping information. New columns are named by clonotyping strategy so to allow for multiple clonotyping identifiers to be present in the same VDJ dataframe and make comparisons between these straightforward.

**Examples**

```r
reclonotyped_vgm <- VDJ_clonotype_v3(VDJ=Platypus::small_vgm[[1]],
clone.strategy="cdr3.nt",
 hierarchical = "none", global.clonotype = TRUE)

reclonotyped_vgm <- VDJ_clonotype_v3(VDJ=Platypus::small_vgm[[1]],
clone.strategy="cdr3.homology", homology.threshold = 0.5,
 hierarchical = "single.chains", global.clonotype = TRUE)
```

---

**VDJ_contigs_to_vgm**

Local VDJ dataframe to VDJ.GEX.matrix format utility
**Description**

Formats "VDJ_contigs_annotations.csv" files from cell ranger to match the VDJ_GEX_matrix output using only cells with 1VDJ and 1VJ chain.

**Usage**

VDJ_contigs_to_vgm(directory, sample.names, platypus.version)

**Arguments**

- directory: list containing paths to the "filtered_contig_annotations.csv" files from cell ranger.
- sample.names: vector specifying sample names.
- platypus.version: Function based on VGM object from V3, no need to set this parameter.

**Value**

data frame with column names that match the VDJ_GEX_matrix output. Can be appended to the VDJ_GEX_matrix output

**Examples**

```r
## Not run:
directory.list <- list()
directory.list[[1]] <- c("~/Dataset_1/filtered_contig_annotations.csv")
directory.list[[2]] <- c("~/Dataset_1/filtered_contig_annotations.csv")
filtered_contig_vgm <- VDJ_contigs_to_vgm(directory = directory.list, sample.names = c(s3,s4))
## End(Not run)
```

---

**VDJ_db_annotate**

Wrapper function of VDJ_antigen_integrate function

**Description**

Wraps the VDJ_antigen_integrate function and uses it to annotate a VDJ dataframe with antigen information. Needs to VDJ_db_load to be executed first, with preprocess=T and vgm.names=T to obtain the same column names as in the VDJ (to allow for sequence matching).

**Usage**

VDJ_db_annotate(VDJ, db.list, database.features, match, homology, lv.distance)
VDJ_db_load

Load and preprocess a list of antigen-specific databases

Description

Preprocessing function for several antigen databases for both TCRs (VDJdb, McPAS-TCR, TBAdb) and BCRs (TBAdb), saving them either at a specified path, or loading them as a database list for downstream integration/analyses.

Usage

VDJ_db_load(
  databases,
  file.paths,
  preprocess,
  species,
  filter.sequences,
  remove.na,
  vgm.names,
  keep.only.common,
  output.format,
  saving.path
)
### Arguments

- **databases**: list of databases to be processed and saved. Currently supported ones include: VDJdb(‘vdjdb’), McPAS-TCR(‘mcpas’), TBAdb(‘tbadb_tcr’ or ‘tbadb_bcr’).
- **file.paths**: list of file paths for the specified databases (in the database parameter). If NULL, will try to locally download the databases from the archived download links.
- **preprocess**: boolean - if T, will preprocess each database individually.
- **species**: string - either ‘Human’ or ‘Mouse’, the species for the processed database. Needs preprocess=T.
- **filter.sequences**: string - ‘VDJ’ to remove rows with NA VDJ sequences, ‘VJ’ to remove rows with NA VJ sequences, ‘VDJ.VJ’ to remove rows with both VDJ and VJ sequences missing. Needs preprocess=T.
- **remove.na**: string or NULL - ‘all’ will remove all rows with missing values from the database, ‘common’ will remove only rows with missing values for the shared columns among all databases (‘VJ_cdr3s_aa’, ‘VDJ_cdr3s_aa’, ‘Species’, ‘Epitope’, ‘Antigen species’), ‘vgm’ will remove missing values for columns shared with the VDJ object (specific to each database). Needs preprocess=T.
- **vgm.names**: boolean - if T, will change all column names of the shared columns (with VDJ) to match those from VDJ. Use this to integrate the antigen data into VDJ using VDJ_antigen_integrate or VDJ_db_annotate. Needs preprocess=T.
- **keep.only.common**: boolean - if T, will only keep the columns shared between all databases (‘VJ_cdr3s_aa’, ‘VDJ_cdr3s_aa’, ‘Species’, ‘Antigen species’) for each processed database. Needs preprocess=T.
- **output.format**: string - ‘df.list’ to save all databases as a list, ‘save’ to save them as csv files.
- **saving.path**: string - directory where the processed databases should be locally saved if output.format=’save’.

### Value

Processed antigen-specific databases for both TCRs and BCRs.

### Examples

```r
## Not run:
VDJ_db_load(databases=list(’vdjdb’), file.paths=NULL, preprocess=TRUE, species=’Mouse’, filter.sequences=’VDJ.VJ’, remove.na=’vgm’, vgm.names=TRUE, keep.only.common=TRUE, output.format=’df.list’)
```

## End(Not run)
**Description**

Calculates and plots common diversity and overlap measures for repertoires and alike. Require the `vegan` package.

**Usage**

```r
VDJ_diversity(
  VDJ,  # VDJ dataframe output from either the VDJ_analyse (platypus.version = "v2") or from the VDJ_GEX_matrix function (platypus.version = "v3") (VDJ_GEX_matrix.output[[1]])
  feature.columns,  # Character vector. One or more column names from the VDJ of which diversity or overlap metrics are calculated. if more than one column is provided (e.g. c("VDJ_cdr3s_aa","VJ_cdr3s_aa")) these columns will be pasted together before metric calculation. Defaults to "CDRH3_aa" if platypus.version == "v2" and "VDJ_cdr3s_aa" if platypus.version == "v3".
  grouping.column,  # Character. Column name of a column to group metrics by. This could be "sample_id" to calculate the metric for each sample. This column is required if metric = "simpson". If so, the simpson overlap index will be calculated pairwise for all combinations of elements in the grouping.column. Defaults to "none".
  metric,  # Character. Diversity or overlap metric to calculate. Can be c("richness", "bergerparker", "simpson", "ginisimpson", "shannon", "shannonevenness", "jaccard"). Defaults to "shannon". If jaccard is selected, a heatmap with the pairwise comparisons between all groups is returned. If any of the others is selected, a dotplot is returned
  subsample.to.same.n,  # Boolean defaults to TRUE. Whether to subsample larger groups down to the size of the smallest group
  pvalues.label.size,  # Numeric. Only used if overlap indices are calculated. Defaults to 4. Is passed on to ggplot theme

)  
```

**Arguments**

- **VDJ**: VDJ dataframe output from either the VDJ_analyse (platypus.version = "v2") or from the VDJ_GEX_matrix function (platypus.version = "v3") (VDJ_GEX_matrix.output[[1]])
- **feature.columns**: Character vector. One or more column names from the VDJ of which diversity or overlap metrics are calculated. if more than one column is provided (e.g. c("VDJ_cdr3s_aa","VJ_cdr3s_aa")) these columns will be pasted together before metric calculation. Defaults to "CDRH3_aa" if platypus.version == "v2" and "VDJ_cdr3s_aa" if platypus.version == "v3".
- **grouping.column**: Character. Column name of a column to group metrics by. This could be "sample_id" to calculate the metric for each sample. This column is required if metric = "simpson". If so, the simpson overlap index will be calculated pairwise for all combinations of elements in the grouping.column. Defaults to "none".
- **metric**: Character. Diversity or overlap metric to calculate. Can be c("richness", "bergerparker", "simpson", "ginisimpson", "shannon", "shannonevenness", "jaccard"). Defaults to "shannon". If jaccard is selected, a heatmap with the pairwise comparisons between all groups is returned. If any of the others is selected, a dotplot is returned
- **subsample.to.same.n**: Boolean defaults to TRUE. Whether to subsample larger groups down to the size of the smallest group
- **pvalues.label.size**: Numeric. Only used if overlap indices are calculated. Defaults to 4. Is passed on to ggplot theme
**axis.label.size**

Numeric. Only used if overlap indices are calculated. Defaults to 12. Is passed on to ggplot theme.

**platypus.version**

Version of platypus to use. Defaults to "v3". If an output of the VDJ_analyze function is supplied, set to "v2". If an output of the VDJ_GEX_matrix function is supplied set to "v3"

**Value**

Returns a ggplot with the calculated metric for each group (if provided). Data is accessible via ggplot.output$data

**Examples**

```r
#Calculate shannon index for VDJ CDR3s by sample
plot <- VDJ_diversity(VDJ = Platypus::small_vgm[[1]], platypus.version = "v3"
 ,feature.columns = c("VDJ_cdr3s_aa"), grouping.column = "sample_id"
 ,metric = "shannon")
#For raw values use
plot$data

#Calculate Gini-simpson and Simpson index for VDJ and VJ CDR3s by sample
VDJ_diversity(VDJ = Platypus::small_vgm[[1]], platypus.version = "v3"
 ,feature.columns = c("VDJ_cdr3s_aa","VJ_cdr3s_aa"), grouping.column = "sample_id"
 ,metric = c("ginisimpson"))

#Calculate Jaccard index of J gene usage between two samples
VDJ_diversity(VDJ = Platypus::small_vgm[[1]], platypus.version = "v3"
 ,feature.columns = c("VDJ_jgene"), grouping.column = "sample_id"
 ,metric = "jaccard")
```

---

**VDJ_dublets**

*Platypus V2 annotation utility*

**Description**

Only Platypus v2 produces a matrix indicating either the number of cells or clones which contain multiple heavy or light chains (or alpha/beta in the case of T cells).

**Usage**

VDJ_dublets(clonotype.list, clone.level)
**Arguments**

- `clonotype.list` Output from VDJ_analyze function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire.
- `clone.level` Logical indicating whether the matrix should display information on the clone level. TRUE will result in matrices containing information about the number of chains on the clonal level. FALSE will result in matrices depicting the number of cells.

**Value**

Returns a list of matrices containing the number of heavy/light chains per either cell or clone depending on the `clone.level` parameter. This can then be supplied to heatmap functions directly. Each list element corresponds to each of the input list elements of clonotypes.

**Examples**

```r
## Not run:
example.vdj.analyze <- VDJ_dublets(clonotype.list = "VDJ.analyze.output", clone.level=T)
## End(Not run)
```

---

**VDJ_dynamics**

Tracks a specific VDJ column across multiple samples/timepoints.

**Description**

Track a VDJ column across multiple samples or timepoints. Tracking consists of creating a per sample/timepoint dataframe of unique values for the VDJ column and their respective counts inside that timepoints/repertoire. Also creates alluvial plots to show the temporal dynamics of the tracked elements.

**Usage**

```r
VDJ_dynamics(
  VDJ, 
  columns.to.track, 
  starting.point.repertoire, 
  track.all.elements, 
  track.only.common, 
  max.elements.to.track, 
  specific.elements.to.track, 
  additional.grouping.column, 
  max.additional.groups, 
  specific.additional.groups, 
  timepoints.column, 
  proportions.level, 
)```
Arguments

**VDJ**

VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.

**columns.to.track**

string or list of strings - VDJ column with values to track (e.g., 'VDJ_cgene’ will track the changes in isotype counts/proportions across multiple timepoints, defined by the timepoints.column). If two columns are provided and tracked, then a new values will be created by combining the values from each column.

**starting.point.repertoire**

string or integer - the repertoire from which to start tracking (1 = will start at the first repertoire, 's3' will start at repertoire 's3').

**track.all.elements**

boolean - if T (and track.only.common=F), it will track all elements across all repertoires/timepoints.

**track.only.common**

boolean - if T (and track.all.elements=F), it will only track the common elements across all repertoires/timepoints.

**max.elements.to.track**

integer or NULL - the maximum number of elements to track (elements are first sorted by frequency/abundance). If NULL, it will track all elements.

**specific.elements.to.track**

vector of strings or NULL - specific elements we want tracked. If NULL, all elements will be tracked.

**additional.grouping.column**

string or 'none' - VDJ column for calculating the frequency/counts of elements on a per-group level. If output.format='plot’, each unique group will have its own bar plot of timepoints/repertoires (x axis) and feature counts (y axis). If NULL, no additional grouping will be done.

**max.additional.groups**

integer or NULL - the maximum number of additional groups to consider (groups are first ordered by their frequency = total number of cells in that group in the VDJ matrix). If NULL, all groups will be considered.

**specific.additional.groups**

vector of strings or NULL - specific grouping factors we want to consider. If NULL, all grouping factors will be considered.

**timepoints.column**

string - VDJ column with either timepoints or repertoires across which we want to track our elements (usually 'sample_id').

**proportions.level**

string - 'absolute.counts’ for absolute counts, ‘group’ for per group proportions, 'repertoire’ for per repertoire/timepoint proportions.
VDJExpandAberrants

output.format string - 'plot' for alluvial barplots, 'df' for count/proportions dataframes of the tracked elements.

ignore.legend boolean - if T, the legend will not be included in the resulting ggplot object.

Value

Either a count dataframe of the tracked elements across multiple timepoints/repertoires, or alluvial barplot.

Examples

VDJ_dynamics(VDJ = small_vgm[[1]], columns.to.track='clonotype.id', starting.point.repertoire=1, max.elements.to.track=10, timepoints.column='sample.id', output.format='plot')

VDJ_expand_aberrants

Expand the aberrant cells in a VDJ dataframe by converting them into additional rows

Description

Expand the aberrant cells in a VDJ dataframe by converting them into additional rows. Aberrant cells consist of cells with more than 1 VDJ or VJ chain.

Usage

VDJ_expand_aberrants(
  VDJ,
  chain.to.expand,
  add.barcode.prefix,
  additional.VDJ.features,
  additional.VJ.features,
  add.CDR3aa,
  add.expanded.number,
  recalculate.clonotype.frequency
)

Arguments

VDJ VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.

chain.to.expand string, 'VDJ' to expand VDJ aberrants, 'VJ' to expand VJ aberrants, 'VDJ.VJ' for both.

add.barcode.prefix boolean - if T, a new barcode will be added for each expanded aberrant.
additional.VDJ.features
vector of strings - VDJ_expand_aberrants will only expand across the sequence columns of VDJ. If you have additional columns with aberrant cell features (e.g., both 'yes' and 'no' binders for a single sequence), where the aberrants are VDJ-specific, include them here.

additional.VJ.features
vector of strings - VDJ_expand_aberrants will only expand across the sequence columns of VDJ. If you have additional columns with aberrant cell features (e.g., both 'yes' and 'no' binders for a single sequence), where the aberrants are VJ-specific, include them here.

add.CDR3aa
boolean - if T, will create a new column 'CDR3aa' with pasted VDJ_cdr3s_aa and VJ_cdr3s_aa.

add.expanded.number
boolean - if T, will add the number of new cells resulting from an aberrant one.

recalculate.clonotype.frequency
boolean - if T, will recalculate the clonotype frequencies for the resulting, expanded VDJ.

Value
Returns a VDJ format dataframe in which cells with more than one VDJ or VJ chain are split into multiple rows each containing only one VDJ VJ chain combination.

Examples
VDJ_expand_aberrants(VDJ = small_vgm[[1]],
chain.to.expand='VDJ.VJ',
add.barcode.prefix=TRUE, recalculate.clonotype.frequency=FALSE)

VDJ_extract_germline  Platypus V2 utility for full germline sequence via MiXCR

Description
Only Platypus v2. Extracts the full-length germline sequence as determined by cellranger. This function returns an object that now contains the reference germline for each of the clones. If multiple clones (as determined by cellranger) have been merged using the VDJ_clonotype function then these sequences may have distinct germline sequences despite being in the same clonal family (nested list). This is particularly possible when homology thresholds were used to determine the clonotypes.
**Usage**

```
VDJ_extract_germline(
    VDJ.per.clone,
    mixcr.directory,
    extract.VDJRegion,
    species
)
```

**Arguments**

- **VDJ.per.clone**  The output from the VDJ_per_clone function. This object should have information regarding the contigs and clonotype_ids for each cell.
- **mixcr.directory**  The directory containing an executable version of MiXCR. This must be downloaded separately and is under a separate license.
- **extract.VDJRegion**  Default is TRUE. Future iterations will allow for distinct gene regions to be extracted.
- **species**  Either "mus" or "hsa" for mouse and human respectively. Default is set to mouse.

**Value**

Returns a dataframe containing repertoire information, such as isotype, CDR sequences, mean number of UMI's. This output can be supplied to further packages VDJ_extract_sequences and VDJ_GEX_integrate

**Examples**

```r
## Not run:
VDJ_extract_germline(VDJ.per.clone=VDJ.per.clone.output,
mixcr.directory="~/Downloads/mixcr-3.0.12/mixcr",
extract.VDJRegion=T,species = "mmu")
## End(Not run)
```

---

**Desription**

Function to get shared/public elements across multiple repertoires, specified by the feature.columns parameter (a column of the VDJ matrix). If two columns are specified in feature.columns, the resulting shared features will combine the values from each column (at a per-cell level).
Usage

VDJ_get_public(
    VDJ,
    feature.columns,
    repertoire.column,
    specific.repertoires,
    find.public.all,
    find.public.percentage,
    treat.combined.features,
    output.format
)

Arguments

VDJ  VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.

feature.columns  Character or character vector columns of features to be assayed

repertoire.column  string - the repertoire-defining column (default to 'sample_id').

specific.repertoires  vector of strings or NULL - if only the shared elements from specific repertoires should be taken into account. If NULL, will output the shared/public elements across all repertoires.

find.public.all  boolean - if T, will look for the public elements across all repertoires

find.public.percentage  list - the first element denotes the percentage of repertoires to get shared elements for, the second element is the maximum number of repertoire combinations to consider (can be NULL to consider all).

treat.combined.features  string - 'exclude' will exclude combined features with one element missing, 'include' will include and considers them as a new feature value.

output.format  string - 'df' to get a shared element dataframe (with columns = Repertoire and Public), 'list' for a list of shared elements.

Value

Either a dataframe of public elements across multiple repertoires or a list.

Examples

VDJ_get_public(VDJ = small_vgm[[1]],
               feature.columns='VDJ_cdr3s_aa', find.public.all=TRUE,
               output.format='df')
**VDJ_GEX_clonal_lineage_clusters**

*Platypus V2 lineage - GEX integration utility*

**Description**

only Platypus v2 Integrates the transcriptional cluster information into the clonal lineages. This requires that automate_GEX, VDJ_clonal_lineages, and VDJ_GEX_integrate have already been ran. The transcriptional cluster will be added to the end of the Name for each sequence.

**Usage**

```r
VDJ_GEX_clonal_lineage_clusters(
  VDJ_GEX_integrate.output,
  VDJ_clonal_lineages.output
)
```

**Arguments**

`VDJ_GEX_integrate.output`

The output from the VDJ_GEX_integrate function that is performed on the VDJ_per_clone level. This involves a nested list where the outer list corresponds to the repertoire and inner lists correspond to specific clones based on the clonotyping strategy.

`VDJ_clonal_lineages.output`

Output from VDJ_clonal_lineages. This should be nested list, with the outer list element corresponding to the individual repertoire and the inner list corresponding to individual clonal lineages based on the initial clonotyping strategy in the form of a dataframe with either Seq or Name. The Name currently contains the barcode following the last "_".

**Value**

a nested list in the identical format to the VDJ_clonal_lineages.output but the name of each sequence will have been changed to include the transcriptional cluster corresponding to that barcode from the GEX library. This requires first running the

**Examples**

```r
## Not run:
clonal_lineages <- VDJ_clonal_lineages(call_MIXCR.output=call_MIXCR_output,
  VDJ_extract_germline.output=VDJ_extract_germline_output,
  as.nucleotide=FALSE,with.germline=TRUE)

## End(Not run)
```
**Description**

only Platypus v2 Integrates VDJ and gene expression libraries by providing cluster membership seq_per_vdj object. Output will plot which transcriptional cluster (GEX) that the cells of a given clonotype are found in.

**Usage**

```r
VDJ_GEX_expansion(
  GEX.list,
  VDJ.GEX.integrate.list,
  highlight.isotype,
  highlight.number
)
```

**Arguments**

- **GEX.list**
  The output of the automate_GEX function.

- **VDJ.GEX.integrate.list**
  Output from VDJ_GEX_integrate function. This object needs to have the GEX and VDJ information combined and integrated. This should be on the CLONAL level from the VDJ_GEX_integrate function.

- **highlight.isotype**
  (Optional) isotype to plot, choose between ["None","A","E","M","G","G1","G2A","G2B","G2C","G3"]. Default is None.

- **highlight.number**
  A vector corresponding to the rank of the clones that should be specified. Default is set to "20", which will present the cluster distribution for the top 20 clones.

**Value**

`ggplot2` plot that breaks down clonotype membership per cluster for the specified input clones.

**Examples**

```r
## Not run:
vdj.gex.expansion <- VDJ_GEX_expansion(GEX.list=GEX.list.output[[1]],
                                       VDJ.GEX.integrate.list=vdj.gex.integrate.output,
                                       highlight.isotype = "None",highlight.number=1:20)
## End(Not run)
```
**VDJ_GEX_integrate**

only Platypus v2 Integrates VDJ and gene expression libraries by providing cluster membership seq_per_vdj object and the index of the cell in the Seurat RNA-seq object.

**Description**

only Platypus v2 Integrates VDJ and gene expression libraries by providing cluster membership seq_per_vdj object and the index of the cell in the Seurat RNA-seq object.

**Usage**

`VDJ_GEX_integrate(GEX.object, clonotype.list, VDJ.per.clone, clonotype.level)`

**Arguments**

- **GEX.object**
  A single seurat object from automate_GEX function. This will likely be supplied as automate_GEX.output[[1]].

- **clonotype.list**
  Output from either VDJ_analyze or VDJ_clonotype functions. This list should correspond to a single GEX.list object, in which each list element in clonotype.list is found in the GEX.object. Furthermore, these repertoires should be found in the automate_GEX library.

- **VDJ.per.clone**
  Output from the VDJ_per_clone function. Each element in the list should be found in the output from the automate_GEX function.

- **clonotype.level**
  Logical specifying whether the integration should occur on the cellular level (VDJ_per_clone) or on the clonotype level (e.g. output from VDJ_analyze or VDJ_clonotype). TRUE specifies that the clonotype level will be selected - e.g. the clonotype.list object will now contain information from the GEX object regarding clonal membership.

**Value**

Returns a nested list containing information corresponding to either the clonal level or the sequence level, depending on the input argument "clonotype.level". This function essentially will update the output of the analyze_VDJ or the VDJ_per_clone functions.

**Examples**

```r
## Not run:
testing_integrate <- VDJ_GEX_integrate(GEX.object = automate.gex.output[[1]],
,clonotype.list = VDJ.analyze.output
,VDJ.per.clone = VDJ.per.clone.output,clonotype.level = TRUE)

## End(Not run)
```
Description

This function is designed as a common input to the Platypus pipeline. Integration of datasets as well as VDJ and GEX information is done here. Please check the Platypus V3 vignette for a detailed walkthrough of the output structure. In short: output[[1]] = VDJ table, output[[2]] = GEX Seurat object and output[[3]] = statistics [FB] Feature barcode (FB) technology is getting increasingly popular, which is why Platypus V3 fully supports their use as sample delimiters. As of V3, Platypus does not support Cite-seq data natively, also the VDJ_GEX_matrix function is technically capable of loading a Cite-seq matrix and integrating it with VDJ. For details on how to process sequencing data with FB data and how to supply this information to the VDJ_GEX_matrix function, please consult the dedicated vignette on FB data.

Usage

```
VDJ_GEX_matrix(
    VDJ.out.directory.list,
    GEX.out.directory.list,
    FB.out.directory.list,
    Data.in,
    Seurat.in,
    GEX.read.h5,
    VDJ.combine,
    GEX.integrate,
    integrate.GEX.to.VDJ,
    integrate.VDJ.to.GEX,
    exclude.GEX.not.in.VDJ,
    filter.overlapping.barcode.GEX,
    filter.overlapping.barcode.VDJ,
    exclude.on.cell.state.markers,
    get.VDJ.stats,
    numcores,
    trim.and.align,
    append.raw.reference,
    select.excess.chains.by.umi.count,
    excess.chain.confidence.count.threshold,
    gap.opening.cost,
    gap.extension.cost,
    parallel.processing,
    integration.method,
    VDJ.gene.filter,
    mito.filter,
    norm.scale.factor,
    n.feature.rna,
    n.count.rna.min,
)```
n.count.rna.max,
n.variable.features,
cluster.resolution,
neighbor.dim,
mds.dim,
subsample.barcodes,
FB.count.threshold,
FB.ratio.threshold,
FB.exclude.pattern,
group.id,
verbose
}

Arguments

**VDJ.out.directory.list**
List containing paths to VDJ output directories from cell ranger. This pipeline assumes that the output file names have not been changed from the default 10x settings in the /outs/ folder. This is compatible with B and T cell repertoires. Necessary files within this folder: filtered_contig_annotations.csv, clonotypes.csv, concat_ref.fasta, all_contig_annotations.csv (only if trim.and.align == T) and metrics_summary.csv (Optional, will be appended to stats table if get.VDJ.stats == T)

**GEX.out.directory.list**
List containing paths the outs/ directory of each sample or directly the raw or filtered_feature_bc_matrix folder. Order of list items must be the same as for VDJ.

**FB.out.directory.list**
[FB] List of paths pointing at the outs/ directory of output from the Cellranger counts function which contain Feature barcode counts. Single list elements can be a path or "PLACEHOLDER", if the corresponding input in the VDJ or GEX path does not have any adjunct FB data. This is only the case when integrating two datasets of which only one has FB data. See examples for details. Any input will overwrite potential FB data loaded from the GEX input directories. This may be important, if wanting to input unfiltered FB data that will cover also cells in VDJ not present in GEX.

**Data.in**
Input for R objects from either the PlatypusDB_load_from_disk or the PlatypusDB_fetch function. If provided, input directories should not be specified. If you wish to integrate local and downloaded data, please load them via load_from_disk and fetch and provide as a list (e.g. Data.in = list(load_from_disk.output, fetch.output))

**Seurat.in**
Alternative to GEX.out.directory.list. A seurat object. VDJ.integrate has to be set to TRUE. In metadata the column of the seurat object, sample_id and group_id must be present. sample_id must contain ids in the format "s1", "s2" ... "sn" and must be matching the order of VDJ.out.directory.list. No processing (i.e. data normalisation and integration) will be performed on these objects. They will be returned as part of the VGM and with additional VDJ data if integrate.VDJ.to.GEX = T. Filtering parameters such as overlapping barcodes, exclude.GEX.not.in.VDJ and exclude.on.cell.state.markers will be applied to the
Seurat.in GEX object(s).

GEX.read.h5
Boolean. defaults to FALSE. Whether to read GEX data from an H5 file. If set to true, please provide the each directory containing a cellranger H5 output file or a direct path to a filtered_feature_bc_matrix.h5 as one GEX.out.directory.list element.

VDJ.combine
Boolean. Defaults to TRUE. Whether to integrate repertoires. A sample identifier will be appended to each barcode both in GEX as well as in VDJ. Recommended for all later functions

GEX.integrate
Boolean. Defaults to TRUE. Whether to integrate GEX data. Default settings use the seurat scale.data option to integrate datasets. Sample identifiers will be appended to each barcode both in GEX and VDJ. This is helpful when analysing different samples from the same organ or tissue, while it may be problematic when analysing different tissues.

integrate.GEX.to.VDJ
Boolean. defaults to TRUE. Whether to integrate GEX metadata (not raw counts) into the VDJ output dataframe! Only possible, if GEX.integrate and VDJ.combine are either both FALSE or both TRUE

integrate.VDJ.to.GEX
Boolean. defaults to TRUE. Whether to integrate VDJ data into GEX seurat object as metadata. ! Only possible, if GEX.integrate and VDJ.combine are either both FALSE or both TRUE

exclude.GEX.not.in.VDJ
Boolean. defaults to FALSE. Whether to delete all GEX cell entries, for which no VDJ information is available. Dependent on data quality and sequencing depth this may reduce the GEX cell count by a significant number

filter.overlapping.barcodes.GEX
Boolean. defaults to TRUE. Whether to remove barcodes which are shared among samples in the GEX analysis. Shared barcodes normally appear at a very low rate.

filter.overlapping.barcodes.VDJ
Boolean. defaults to TRUE. Whether to remove barcodes which are shared among samples in the GEX analysis. Shared barcodes normally appear at a very low rate.

exclude.on.cell.state.markers
Character vector. If no input is provided or input is "none", no cells are excluded. Input format should follow: Character vector containing the gene names for each state. ; is used to use multiple markers within a single gene state. Different vector elements correspond to different states. Example: c("CD4+;CD44-","CD4+;IL7R+;CD44+"). All cells which match any of the given states (in the example case any of the 2) are excluded. This is useful in case different and non lymphocyte cells were co-sequenced. It should give the option to e.g. exclude B cells in the analysis of T cells in a dataset.

get.VDJ.stats
Boolean. defaults to TRUE. Whether to generate general statistics table for VDJ repertoires. This is appended as element [3] of the output list.

get.VDJ.stats
Number of cores used for parallel processing. Defaults to number of cores available. If you want to check how many cores are available use the library Parallel
and its command detectCores() (Not setting a limit here when running this function on a cluster may cause a crash)

trim.and.align  Boolean. Defaults to FALSE. Whether to trim VJ/VDJ seqs, align them to the 10x reference and trim the reference. This is useful to get full sequences for antibody expression or numbers of somatic hypermutations. Setting this to TRUE significantly increases computational time

append.raw.reference  Boolean. Defaults to TRUE. This appends the raw reference sequence for each contig even if trim.and.align is set to FALSE.

select.excess.chains.by.umi.count  Boolean. Defaults to FALSE. There are several methods of dealing with cells containing reads for more than 1VDJ and 1VJ chain. While many analyses just exclude such cells, the VGM is designed to keep these for downstream evaluation (e.g. in VDJ_clonotype). This option presents an evidenced-based way of selectively keeping or filtering only one of the present VDJ and VJ chains each. This works in conjunction with the parameter excess.chain.confidence.count.threshold (below) Idea source: Zhang W et al. Sci Adv. 2021 (10.1126/sciadv.abf5835)

excess.chain.confidence.count.threshold  Integer. Defaults to 1000. This sets a umi count threshold for keeping excessive chains in a cell (e.g. T cells with 2 VJ and 1 VDJ chain) and only has an effect if select.excess.chains.by.umi.count is set to TRUE. For a given cell with chains and their UMI counts: VDJ1 = 3, VDJ2 = 7, VJ1 = 6. If count.threshold is kept at default (1000), the VDJ chain with the most UMIs will be kept (VDJ2), while the other is filtered out (VDJ1), leaving the cell as VDJ2, VJ1. If the count.threshold is set to 3, both chains VDJ chains of this cell are kept as their UMI counts are equal or greater to the count.threshold and therefore deemed high confidence chains. In the case of UMI counts being equal for two chains AND below the count.threshold, the first contig entry is kept, while the second is filtered. To avoid filtering excess chains, set select.excess.chains.by.umi.count to FALSE. For further notes on the implication of these please refer to the documentation of the parameter hierarchical in the function VDJ_clonotype_v3.

gap.opening.cost  Argument passed to Biostrings::pairwiseAlignment during alignment to reference. Defaults to 10

gap.extension.cost  Argument passed to Biostrings::pairwiseAlignment during alignment to reference. Defaults to 4

parallel.processing  Character string. Can be "parlapply" for Windows system, "mclapply" for unix and Mac systems or "none" to use a simple for loop (slow!). Default is "none" for compatibility reasons. For the parlapply option the packages parallel, doParallel and the dependency foreach are required

integration.method  String specifying which data normalization and integration pipeline should be used. Default is "scale.data", which correspondings to the ScaleData function internal to harmony package. 'anchors' scales data individually and then finds and align cells in similar states as described here: https://satijalab.org/seurat/articles/integration_introduction.html
'sct' specifies SCTransform from the Seurat package. "harmony" should be specified to perform harmony integration. This method requires the harmony package from bioconductor.

**VDJ.gene.filter**
Logical indicating if variable genes from the b cell receprot and t cell receptor should be removed from the analysis. True is highly recommended to avoid clonal families clustering together.

**mito.filter**
Numeric specifying which percent of genes are allowed to be composed of mitochondrial genes. This value may require visual inspection and can be specific to each sequencing experiment. Users can visualize the percentage of genes corresponding to mitochondrial genes using the function "investigate_mitochondial_genes".

**norm.scale.factor**
Scaling factor for the standard Seurat pipeline. Default is set to 10000 as reported in Seurat documentation.

**n.feature.rna**
Numeric that specifies which cells should be filtered out due to low number of detected genes. Default is set to 0. Seurat standard pipeline uses 2000.

**n.count.rna.min**
Numeric that specifies which cells should be filtered out due to low RNA count. Default is set to 0. Seurat standard pipeline without VDJ information uses 200.

**n.count.rna.max**
Numeric that specifies which cells should be filtered out due to high RNA count. Default is set to infinity. Seurat standard pipeline without VDJ information uses 2500.

**n.variable.features**
Numeric specifying the number of variable features. Default set to 2000 as specified in Seurat standard pipeline.

**cluster.resolution**
Numeric specifying the resolution that will be supplied to Seurat’s FindClusters function. Default is set to 0.5. Increasing this number will increase the number of distinct Seurat clusters. Suggested to examine multiple parameters to ensure gene signatures differentiating clusters remains constant.

**neighbor.dim**
Numeric vector specifying which dimensions should be supplied in the FindNeighbours function from Seurat. Default input is ‘1:10’.

**mds.dim**
Numeric vector specifying which dimensions should be supplied into dimensional reduction techniques in Seurat and Harmony. Default input is ‘1:10’.

**subsample.barcodes**
For development purposes only. If set to TRUE the function will run on 100 cells only to increase speeds of debugging

**FB.count.threshold**
Numeric. Defaults to 10. For description of Feature Barcode assignment see parameter FB.ratio.threshold above

**FB.ratio.threshold**
Numeric. Defaults to 2 Threshold for assignment of feature barcodes by counts. A feature barcode is assigned to a cell if its counts are >FB.count.threshold and if its counts are FB.ratio.threshold-times higher than the counts of the feature barcode with second most counts.
FB.exclude.pattern

Character (regex compatible). If a feature barcode matches this pattern it will be excluded from the hashing sample assignments. This may be necessary if CITE-seq barcodes and hashing barcodes are sequenced in the same run.

group.id

vector with integers specifying the group membership. c(1,1,2,2) would specify the first two elements of the input VDJ/GEX lists are in group 1 and the third/fourth input elements will be in group 2.

verbose

if TRUE prints runtime info to console. Defaults to TRUE

Value

Single cell matrix including VDJ and GEX info. Format is a list with out[[1]] = a VDJ dataframe (or list of dataframes if VDJ.combine == F, not recommended) containing also selected GEX information of integrate.GEX.to.VDJ = T. out[[2]] = GEX Seurat object with the metadata also containing GEX information if integrate.VDJ.to.GEX = T. out[[3]] = Dataframe with statistics on GEX and VDJ. out[[4]] = runtime parameters. out[[5]] = session info

Examples

## Not run:

#FOR EXAMPLES see Platypus vignette at https://alexyermanos.github.io/Platypus/index.html

#Run from local directory input. For run from PlatypusDB input see

#PlatypusDB vignette

VDJ.out.directory.list <- list()
VDJ.out.directory.list[[1]] <- c("~/VDJ/S1/")
VDJ.out.directory.list[[2]] <- c("~/VDJ/S2/")
GEX.out.directory.list <- list()
GEX.out.directory.list[[1]] <- c("~/GEX/S1/")
GEX.out.directory.list[[2]] <- c("~/GEX/S2/")

VGM <- VDJ_GEX_matrix(
VDJ.out.directory.list = VDJ.out.directory.list,
GEX.out.directory.list = GEX.out.directory.list,
integrate.GEX.to.VDJ = T,
integrate.VDJ.to.GEX = T,
exclude.GEX.not.in.VDJ = F,
filter.overlapping.barcodes.GEX = F,
filter.overlapping.barcodes.VDJ = F,
get.VDJ.stats = T,
parallel.processing = "none",
subsample.barcodes = F,
trim.and.align = F,
group.id = c(1,2))

# With Feature Barcodes

## Option 1: Cellranger multi or Cellranger count with --libraries output

VDJ.out.directory.list <- list()
VDJ.out.directory.list[[1]] <- "~/VDJ/S1/" #point to outs or per_sample_outs directory content
VDJ.GEX_overlay_clones

VDJ.out.directory.list[[2]] <- "~/VDJ/S2/
GEX.out.directory.list <- list()
GEX.out.directory.list[[1]] <- "~/GEX/S1/
GEX.out.directory.list[[2]] <- "~/GEX/S2/
# These directories contain two matrices (GEX and FB)
VGM <- VDJ_GEX_matrix(
  VDJ.out.directory.list = VDJ.out.directory.list,
  GEX.out.directory.list = GEX.out.directory.list,
  FB.ratio.threshold = 2)

Option 2: Separate input of FB data from separate Cellranger count run
VDJ.out.directory.list <- list()
VDJ.out.directory.list[[1]] <- "~/VDJ/S1/
VDJ.out.directory.list[[2]] <- "~/VDJ/S2/
GEX.out.directory.list <- list()
GEX.out.directory.list[[1]] <- "~/GEX/S1/
GEX.out.directory.list[[2]] <- "~/GEX/S2/
FB.out.directory.list[[1]] <- "~/FB/S1/
FB.out.directory.list[[2]] <- "~/FB/S1/
VGM <- VDJ_GEX_matrix(
  VDJ.out.directory.list = VDJ.out.directory.list,
  GEX.out.directory.list = GEX.out.directory.list,
  FB.out.directory.list = FB.out.directory.list,
  FB.ratio.threshold = 2)

Option 3: FB input for two datasets of which only one contains FB data
VDJ.out.directory.list <- list()
VDJ.out.directory.list[[1]] <- "~/study1/VDJ/S1/
VDJ.out.directory.list[[2]] <- "~/study2/VDJ/S1/
VDJ.out.directory.list[[3]] <- "~/study2/VDJ/S2/
GEX.out.directory.list <- list()
GEX.out.directory.list[[1]] <- "~/study1/GEX/S1/
GEX.out.directory.list[[2]] <- "~/study2/GEX/S1/
GEX.out.directory.list[[2]] <- "~/study2/GEX/S2/
FB.out.directory.list[[1]] <- "PLACEHOLDER" # Study 1 does not contain FB data
FB.out.directory.list[[2]] <- "~/study2/FB/S1/
FB.out.directory.list[[3]] <- "~/study2/FB/S2/
VGM <- VDJ_GEX_matrix(
  VDJ.out.directory.list = VDJ.out.directory.list,
  GEX.out.directory.list = GEX.out.directory.list,
  FB.out.directory.list = FB.out.directory.list,
  FB.ratio.threshold = 2)

## End (Not run)

VDJ.GEX_overlay_clones
Overlay clones on GEX projection
**Description**

Highlights the cells belonging to any number of top clonotypes or of specifically selected clonotypes from one or more samples or groups in a GEX dimensional reduction.

**Usage**

```r
VDJ_GEX_overlay_clones(
  GEX,
  reduction,
  n.clones,
  clones.to.plot,
  by.sample,
  by.other.group,
  ncol.facet,
  pt.size,
  clone.colors,
  others.color,
  split.plot.and.legend,
  platypus.version
)
```

**Arguments**

- **GEX**
  A single seurat object from VDJ_GEX_matrix, which also includes VDJ information in the metadata (set integrate.VDJ.to.GEX to TRUE in the VDJ_GEX_matrix function) (VDJ_GEX_matrix.output[[2]])! Clone ids and frequencies are drawn from the columns "clonotype_id" and "clonotype_frequency"

- **reduction**
  Character. Defaults to "umap". Name of the reduction to overlay clones on. Can be "pca", "umap", "tsne"

- **n.clones**
  Integer. Defaults to 5. To PLOT TOP N CLONES. Number of Top clones to plot. If either by.sample or by.group is TRUE, n.clones clones from each sample or group will be overlayed

- **clones.to.plot**
  Character. Alternative to n.clones. TO PLOT SPECIFIC CLONES. Must reference a column in the GEX@meta.data filled with TRUE and FALSE. Entries with TRUE label are plotted. Such a column may be generated using GEX@metadata$clones_to_plot_column <- GEX@metadata$Some_cell_identifier == "Interesting"

- **by.sample**
  Boolean. Defaults to FALSE. Whether to overlay clones by sample. If set to TRUE this will generate a facet_wrap plot with as many facets as samples.

- **by.other.group**
  Character string. Defaults to "none". Must be a valid column name of the metadata of the input seurat object. If so, this will generate a facet_wrap plot with as many facets unique entries in the specified column. This may be useful to plot cell type specific clones

- **ncol.facet**
  Integer. Defaults to 2. Number of columns in the facet_wrap plot if by.sample or by.group is TRUE

- **pt.size**
  Numeric. Defaults to 1. Size of points in DimPlot. Passed to Seurat::DimPlot
clone.colors  Character vector. Defaults to rainbow(n.clones). Colors to use for individual clones. One can provide either a vector of length n.clones or a of length Nr. of samples/groups \* n.clones. In case that a vector of length n.clones is provided and by.group or by.sample is TRUE, colors are repeated for each sample/group.

others.color  Character. Color for cells that are not selected i.e. not part of the overlayed clonotypes. Defaults to "grey80". To hide the rest of the umap set to "white"

split.plot.and.legend  Boolean. Defaults to FALSE. Whether to return the plot and the legend separately as a list. This can be useful if legends get large and distort the actual plots. The packages gridExtra and cowplot are required for this. If set to TRUE a list is returned where out[[1]] is the plot which can be printed just by executing out[[1]]; out[[2]] is the legend, which can be printed either using plot(out[[2]]) or grid.arrange(out[[2]])

platypus.version  Character. At the moment this function runs only on the output of the VDJ_GEX_matrix function meaning that it is exclusively part of Platypus "v3". With further updates the functionality will be extended.

Value

A ggplot object or a list of a ggplot and a gtable legend (if split.plot.and.legend \=\= TRUE). Theme, colors etc. may be changed directly by adding new elements to this output (e.g. out \+ theme_minimal())

Examples

#To return a single plot with top clones across samples
overlay_clones_plot <- VDJ_GEX_overlay_clones(
  GEX = Platypus::small_vgm[[2]], reduction = "umap"
  ,n.clones = 5, by.sample = FALSE
  ,by.other.group = "none", pt.size = 1,split.plot.and.legend = FALSE)

#To return a facet plot with top clones for each sample
overlay_clones_plot <- VDJ_GEX_overlay_clones(
  GEX = Platypus::small_vgm[[2]], reduction = "umap"
  ,n.clones = 5, by.sample = TRUE, by.other.group = "none"
  ,pt.size = 1,ncol.facet = 2, split.plot.and.legend = FALSE)

#To return a facet plot and the legend separately with top clones for each group
overlay_clones_plot <- VDJ_GEX_overlay_clones(
  GEX = Platypus::small_vgm[[2]], reduction = "umap"
  ,n.clones = 5, by.sample = TRUE, by.other.group = "group_id", pt.size = 1
  ,ncol.facet = 2, split.plot.and.legend = TRUE)

#To print both:
#overlay_clones_plot[[1]] #Plot
#gridExtra::grid.arrange(overlay_clones_plot[[2]]) #Legend
#To save, ggsave() is applicable to both
#To return a single plot with selected clones
#add a clonotype_to_plot column
#GEX$meta.data$clonotype_to_plot <- GEX$VJ_vgene == "TRAV5-1"
#Column with TRUE for all clones with a particular V gene
#overlay_clones_plot <- VDJ_GEX_overlay_clones(GEX = GEX, reduction = "umap"
#, clones.to.plot = "clonotype_to_plot", by.sample = TRUE, by.other.group = "none"
#, split.plot.and.legend = FALSE, pt.size = 1.5)

**VDJ_GEX_stats**

*Standalone VDJ and GEX statistics.*

**Description**

Gives stats on number and quality of reads. This function is integrated into the VDJ_GEX_matrix. Before running, please check list element [[3]] of VDJ_GEX_matrix output for already generated statistics.

**Usage**

```r
VDJ_GEX_stats(
    VDJ.out.directory,
    GEX.out.directory,
    sample.names,
    metrics10x,
    save.csv,
    filename
)
```

**Arguments**

- **VDJ.out.directory** List of paths with each element containing the path to the output of cellranger VDJ runs. This pipeline assumes that the output file names have not been changed from the default 10x settings in the /outs/ folder. This is compatible with B and T cell repertoires (both separately and simultaneously).
- **GEX.out.directory** OPTIONAL list of paths with each element containing the path to the output of cellranger GEX runs. This pipeline assumes that the output file names have not been changed from the default 10x settings in the /outs/ folder. This is compatible with B and T cell repertoires (both separately and simultaneously).
- **sample.names** OPTIONAL: an array of the same length as the input VDJ.out.directory list with custom names for each sample. If not provided samples will be numbered by processing order
- **metrics10x** Whether to append metrics_summary.csv information provided by Cellranger for both VDJ and GEX. Defaults to T
- **save.csv** Boolean. Defaults to TRUE. Whether to directly save the results as a comma delimited .csv file in the current working directory.
- **filename** Character ending in .csv. Filename to save .csv as.
Value

returns a single matrix where the rows are individual cells and the columns are repertoire features.

Examples

```r
## Not run:
stats <- VDJ_GEX_stats(VDJ.out.directory = VDJ.out.directory.list
, GEX.out.directory = GEX.out.directory.list, sample.names = c(1:4)
, metrics10x = TRUE, save.csv = TRUE , filename = "stats.csv")
## End(Not run)
```

### VDJ_isotypes_per_clone

*Platypus V2 clonal utility*

**Description**

Only for Platypus v2 Clonal frequency plot displaying the isotype usage of each clone. For platypus v3 use VDJ_clonal_expansion

**Usage**

```r
VDJ_isotypes_per_clone(
    VDJ_clonotype_output,
    VDJ_per_clone_output,
    clones,
    subtypes,
    species,
    sample.names,
    treat.incomplete.clones,
    treat.incomplete.cells,
    platypus.version,
    VDJ.matrix
)
```

**Arguments**

- **VDJ_clonotype_output**
  
  list of dataframes based on the VDJ_clonotype function output.

- **VDJ_per_clone_output**
  
  list of dataframes based on the VDJ_per_clone function output.

- **clones**
  
  numeric value indicating the number of clones to be displayed on the clonal expansion plot. Can take values between 1-50. Default value is 50.

- **subtypes**
  
  Logical indicating whether to display isotype subtypes or not.
species  Character indicating whether the samples are from mouse or human. Default is set to human. # @param sample.names  Character vector with the same length of the VDJ.GEX.matrix.out list. If a VDJ table is provided, length of samples names must be one. These names are used as references to the output and as title for the plots

description

treat.incomplete.clones  Character indicating how to proceed with clonotypes lacking a VDJC (in other words, no cell within the clonotype has a VDJC). "exclude" removes these clonotypes from the analysis. This may result in a different frequency ranking of clonotypes than in the output of the VDJ_analyse function with filter.HC.LC = FALSE. "include" keeps these clonotypes in the analysis. In the plot they will appear as having an unknown isotype.

treat.incomplete.cells  Character indicating how to proceed with cells assigned to a clonotype but missing a VDJC. "proportional" to fill in the VDJ isotype according to the proportions present in of clonotype (in case present proportions are not replicable in the total number of cells e.g. 1/3 in 10 cells, values are rounded to the next full integer and if the new counts exceed the total number of cells, 1 is subtracted from the isotype of highest frequency. If the number is below the number of cell, 1 is added to the isotype with lowest frequency to preserve diversity), "exclude" to exclude them from analysis and rank clonotypes only by the number of actual contigs of there heavy chain. This ranking may deviate from the frequency column in the clonotype table. CAVE: if treat.incomplete_cells is set to "exclude", clonotypes lacking a VDJC entirely will be removed from the analysis. This results in a similar but not identical output as when treat_incomplete_clones is set to true. The two parameters are thereby non-redundant.

platypus.version  Defaults to "v3". For a more flexible analysis in v3 use VDJ_clonalExpansion()

VDJ.matrix  The VDJ table output of the VDJ_GEX_matrix function. (VDJ_GEX_matrix.output[[1]])

Value

returns a list containing plots with the percentages of isotypes for each clone on the cell level.

Examples

## Not run:
VDJ.isotype.per.clone <- VDJ_isotypes_per_clone(
  VDJ_clonotype_output = VDJ.analyze.output
  VDJ_per_clone_output = VDJ.per.clone.output, clones = 30)

## End(Not run)
vdj_length_prob A list dataframe specifying lengths and probabilities of bases deleted or inserted at each junction site of VDJ recombination event.

v3_deletion length and probability of deleted bases at 3’ end of V segment
d5_deletion length and probability of deleted bases at 5’ end of D segment
d3_deletion length and probability of deleted bases at 3’ end of D segment
ej5_deletion length and probability of deleted bases at 5’ end of J segment
dj_insertion length and probability of inserted bases between D-J segment
vj_insertion length and probability of inserted bases between V-J segment for light or alpha chains

Description

vdj_length_prob A list dataframe specifying lengths and probabilities of bases deleted or inserted at each junction site of VDJ recombination event.

v3_deletion length and probability of deleted bases at 3’ end of V segment
d5_deletion length and probability of deleted bases at 5’ end of D segment
d3_deletion length and probability of deleted bases at 3’ end of D segment
ej5_deletion length and probability of deleted bases at 5’ end of J segment
dj_insertion length and probability of inserted bases between D-J segment
vj_insertion length and probability of inserted bases between V-J segment for light or alpha chains

Usage

data("vdj_length_prob")

Format

An object of class list of length 7.
**VDJ_logoplot_vector**

Flexible logoplot wrapper

**Description**

Plots a logoplot of the CDR3 aminoacid region

**Usage**

VDJ_logoplot_vector(cdr3.vector, length_cdr3, seq_type)

**Arguments**

- **cdr3.vector**
  A character vector of aa sequences. This is to increase flexibility of this function. Such a sequence vector may be retrieved from the VDJ_analyse function output on a clonotype level or from the VDJ_GEX_matrix function output on a per cell level. Additionally, any length of sequence may be used (e.g. HCDR3 only or H and LCDR3 pasted together)

- **length_cdr3**
  Integer or character. Defaults to "auto". Sets the length of the CDR3 regions that are selected to be plotted. If set to auto, the most frequently appearing length in the vector will be used

- **seq_type**
  passed to ggseqlogo. Can be set to "aa", "dna", "rna" or "other"

**Value**

Returns the logo plot.

**Examples**

VDJ_logoplot_vector(
  cdr3.vector = Platypus::small_vgm[[1]]$VDJ_cdr3s_aa,
  length_cdr3 = "auto", seq_type = "auto")

**VDJ_network**

Similarity networks based on CDR3 regions

**Description**

Creates a similarity network where clones with similar CDR3s are connected.
Usage

VDJ_network(
    VDJ,
    distance.cutoff,
    per.sample,
    platypus.version,
    known.binders,
    hcdr3.only,
    is.bulk
)

Arguments

VDJ Either (for platypus version "v2") output from VDJ_analyse function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire, OR (for platypus version "v3") the VDJ matrix output of the VDJ_GEX_matrix() function (VDJ.GEX.matrix.output[[1]])

distance.cutoff The threshold Levenshtein distance for which two nodes will be connected on the similarity network.

per.sample logical value indicating if a single networks should be produced for each mouse.

platypus.version Character. Defaults to "v3". Can be "v2" or "v3" dependent on the input format

known.binders Either a character vector with cdr3s of known binders or a data frame with cdr3s in the first and the corresponding specificity in the second column. If this parameter is defined, the output will be a network with only edges between known binders and the repertoire nodes and edges between the known binders that have at least one edge to a repertoire node

hcdr3.only logical value indicating if the network is based on heavy chain cdr3s (hcdr3.only = T) or pasted heavy and light chain cdr3s (hcdr3.only = F), works for platypus version 3 only

is.bulk logical value indicating whether the VDJ input was generated from bulk-sequencing data using the bulk_to_vgm function. If is.bulk = T, the VDJ_network function is compatible for use with bulk data. Defaults to False (F).

Value

returns a list containing networks and network information. If per.sample is set to TRUE then the result will be a network for each repertoire. If per.sample == F, output[[1]] <- will contain the network, output[[2]] will contain the dataframe with information on each node, such as frequency, mouse origin etc. output[[3]] will contain the connected index - these numbers indicate that the nodes are connected to at least one other node. output[[4]] contains the paired graph - so the graph where only the connected nodes are drawn.

Examples

#Platypus v2
VDJ_overlap_heatmap

Wrapper to determine and plot overlap between VDJ features across groups

Description

Yields overlap heatmap and datatable of features or combined features for different samples or groups

Usage

VDJ_overlap_heatmap(
  VDJ,  
  feature.columns,  
  grouping.column,  
  jaccard,  
  plot.type,  
  pvalues.label.size,  
  axis.label.size,  
  add.barcode.table
)

Arguments

VDJ 
VDJ output of the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]])

feature.columns 
A character array of column names of which the overlap should be displayed. The content of these columns is pasted together (separated by "/"). E.g. if the overlap in cells germline gene usage is desired, the input could be c("VDJ_jgene","VDJ_dgene","VDJ_vgene"). These columns would be pasted and compared across the grouping variable.

grouping.column 
A column which acts as a grouping variable. If repertoires are to be compared use the sample_id column.

jaccard 
Boolean. Defaults to FALSE. If set to TRUE, the overlap will be reported as jaccard index. If set to FALSE the overlap will be reported as absolute counts

plot.type 
Character. Either "ggplot" or "pheatmap". Defaults to Pheatmap

pvalues.label.size 
Numeric. Defaults to 4. Is passed on to ggplot theme

axis.label.size 
Numeric. Defaults to 4. Is passed on to ggplot theme
add.barcode.table

Boolean. Defaults to T. Whether to generate a dataframe with frequencies and barcodes of cells with overlapping features. This is useful to e.g. analyze differentially expressed genes between cells of two samples or groups expressing the same VDJ or VJ chain

Value

A list of a ggplot (out[[1]]), the source table or matrix for the plot out[[2]] and a table containing additional information in case that add.barcode.table was set to TRUE (out[[3]])

Examples

#To test the overlap of CDR3s between multiple samples
overlap <- VDJ_overlap_heatmap(VDJ = Platypus::small_vgm[[1]],
feature.columns = c("VDJ_cdr3s_aa"),
grouping.column = "sample_id", axis.label.size = 15,
plot.type = "ggplot")

VDJ_per_clone

Description

only Platypus v2 Analyzes and processes the repertoire sequencing data from cellranger vdj. This provides information on the single-cell level for each clone, as opposed to the output from VDJ_analyze.

Usage

VDJ_per_clone(
  clonotype.list,
  VDJ.out.directory,
  contig.list,
  fasta.list,
  reference.list,
  filtered.contigs,
  annotations.json,
  JSON
)

Arguments

clonotype.list Output from either VDJ_analyze or VDJ_clonotype functions. This list should correspond to a single GEX.list object, in which each list element in clonotype.list is found in the GEX.object. Furthermore, the i'th entry in the directory supplied to GEX.list should correspond to the i'th element in the clonotype.list object.
VDJ_out.directory

Character vector with each element containing the path to the output of cellranger vdj runs. This corresponds to the same object used for the VDJ_analyze function. Multiple repertoires to be integrated in a single transcriptome should be supplied as multiple elements of the character vector. This can be left blank if supplying the clonotypes and contig files directly as input. This pipeline assumes that the output file names have not been changed from the default 10x settings in the /outs/ folder. This is compatible with B and T cell repertoires (both separately and simultaneously).

contig.list

List of dataframe based on the all_contigs.csv file from cellranger vdj output. If 10x sequencing was not used then this object should be formatted with the same columns as the 10x object.

fasta.list

Contains the full-length sequence information in the same format as filtered_contig.fasta file from the output of cellranger.

reference.list

Contains the reference sequence information in the same format as concat_ref.fasta file from the output of cellranger.

filtered.contigs

Logical indicating if the filtered contigs file should be used. TRUE will read VDJ information from only the filtered output of cellranger. FALSE will read the all contigs file from cellranger. Default set to TRUE (filtered output)

annotations.json

Optional input from loaded all_contig_annotations.json. Will be read in automatically if not provided

JSON

Boolean. Defaults to FALSE. Whether to load all_contig_annotations.json

Details

Platypus V2 data frame utility

Value

Returns a list of dataframes containing

Examples

```r
# Not run:
VDJ_per_clone_out <- VDJ_per_clone(clonotype.list = output.from.VDJ_analyze,
VDJ.out.directory = "path/to/cellranger/outs/"
)
```

```r
# End(Not run)
```
Description

Creates phylogenetic trees as tidytree dataframes from an input VDJ dataframe. The resulting phylogenetic trees can be plotted using VDJ_phylogenetic_trees_plot. Both of these functions require the tidytree and ggtree packages.

Usage

```r
VDJ_phylogenetic_trees(
  VDJ,
  sequence.type,
  as.nucleotide,
  trimmed,
  include.germline,
  global.clonotype,
  VDJ.VJ.1chain,
  additional.feature.columns,
  filter.na.columns,
  maximum.lineages,
  minimum.sequences,
  maximum.sequences,
  tree.algorithm,
  tree.level,
  no.trees.combined,
  germline.scale.factor,
  output.format,
  parallel
)
```

Arguments

- **VDJ**: VDJ or VDJ.GEX.matrix[[1]] object, as obtained from the VDJ_GEX_matrix function in Platypus.
- **sequence.type**: string - sequences which will be used when creating the phylogenetic trees. 'cdr3' for CDR3s of both VDJs and VJs, 'cdrh3' for VDJ CDR3s, 'VDJ.VJ' for pasted full sequences of both VDJ and VJ, 'VDJ' for full VDJ sequences, 'VJ' for full VJ.
- **as.nucleotide**: boolean - if T, will only consider the DNA sequences specified by sequence.type, else it will consider the amino acid ones.
- **trimmed**: boolean - in the case of full VDJ or VJ nt sequences, if the trimmed sequences should be consider (trimmed=T), or raw ones. You need to call MIXCR first on the VDJ dataframe using VDJ_call_MIXCR().
include.germline
boolean - if T, a germline sequence will be included in the trees (root), obtained by pasting the VDJ_trimmed_ref and VJ_trimmed_ref sequences. You need to call MIXCR first on the VDJ dataframe using VDJ_call_MIXCR().

global.clonotype
boolean - if T, will ignore samples from the sample_id column, creating global clonotypes.

VDJ.VJ.1chain
boolean - if T, will remove aberrant cells from the VDJ matrix.

additional.feature.columns
list of strings or NULL - VDJ column names which will comprise the per-sequence features to be included in the tidytree dataframe, which will be used to label nodes/determines their color/size etc. See also the VDJ_phylogenetic_trees_plot function.

filter.na.columns
list of strings - VDJ columns names: if a phylogenetic tree/tidytree dataframe has all elements = NA in that feature, that tree will be completely removed.

maximum.lineages
integer or 'all' - maximum number of clonotypes to create trees for. If 'all', will create trees for all clonotypes.

minimum.sequences
integer - lower bound of sequences for a tree. Defaults to 3. Trees with a lower number will be automatically removed.

maximum.sequences
integer - upper bound of sequences for a tree. Additional sequences will be removed, after being ordered by their total frequency.

tree.algorithm
string - the algorithm used when constructing the phylogenetic trees. 'nj' for Neighbour-Joining, 'bionj', 'fastme.bal', and 'fastme.ols'

tree.level
string - level at which to build phylogenetic trees. 'intraclonal' - tree per clonotype, per sample, 'global.clonotype' - global clonotype trees (include.germline must be F), irrespective of sample, 'combine.first.trees' will combine the trees for the most expanded clonotypes, per sample (include.germline must be F).

no.trees.combined
integer - number of trees to combine if tree.level='combine.first.trees'.

germline.scale.factor
numeric - as germlines are incredibly distant from their closest neighbours (in the tree), this controls the scale factor for the germline tree branch length for more intelligible downstream plotting.

output.format
string - 'tree.df.list' returns a nested list of tidytree dataframes, per clonotype and per sample; 'lineage.df.list' returns a list of lineage dataframes - unique sequences per clonotype,

parallel
string - parallelization method to be used to accelerate computations, 'none', 'mclapply', or 'parlapply'.

Value

Nested list of tidytree dataframes or lineage dataframes.
Examples

```r
## Not run:
VDJ_phylogenetic_trees(VDJ=VDJ, sequence.type='VDJ.VJ',
trimmed=TRUE, as.nucleotide=TRUE, include.germline=TRUE,
additional.feature.columns=NULL, tree.level='intraclonal',
output.format='tree.df.list')

## End(Not run)
```

---

**VDJ_phylogenetic_trees_plot**

*Phylogenetic tree plotting*

**Description**

Function to plot phylogenetic trees obtained from VDJ_phylogenetic_trees.

*Requires the ggtree package to be loaded! Plots trees from function VDJ_phylogenetic_trees*

**Usage**

```r
VDJ_phylogenetic_trees_plot(
  tree.dfs,
  color.by,
  size.by,
  shape.by,
  specific.leaf.colors
)
```

**Arguments**

- `tree.dfs` nested list of tidytree dataframes obtained from VDJ_phylogenetic_trees with output.format='tree.df.list'. tree.dfs[[1]][[2]] represent a tree dataframe for the first sample, second clonotype.
- `color.by` string - VDJ or tree df column name which will be used to color the tree nodes.
- `size.by` string or NULL - VDJ or tree df column name which determines the node size. If NULL, node sizes will be equal.
- `shape.by` string or NULL - VDJ or tree df column name which determines the node shape. If NULL, node sizes will be equal.
- `specific.leaf.colors` named list or NULL - if NULL, colors will be automatically selected for each node according to its color.by value.

**Value**

nested list of ggtree plot objects for each sample and each clonotype.
Examples

```r
## Not run:
VDJ_phylogenetic_trees_plot(tree.dfs,color.by='clonotype_id', size.by='sequence_frequency')
## End(Not run)
```

## Not run:

### Description

Plots for SHM based on MIXCR output generated using the VDJ_call_MIXCR function and appended to the VDJ.GEX.matrix.output

### Usage

```r
VDJ_plot_SHM(
  VDJ.mixcr.matrix,
  group.by, quantile.label, point.size, mean.line.color,
  stats.to.console, platypus.version
)
```

### Arguments

- **VDJ.mixcr.matrix**: Output dataframe from the VDJ_call_MIXCR function or a dataframe generated using the VDJ_GEX_matrix function and supplemented with MIXCR information
- **group.by**: Character. Defaults to "sample_id". Column name of VDJ.matrix to split VDJ.matrix by. For each unique entry in that column a set of plots will be generated. This can be useful to plot SHM by expansion or by transcriptomics-derived clusters
- **quantile.label**: Numeric. Defaults to 0.9. Which points to label in the SHM scatterplot. If set to 0.9, the top 10% of cells by SHM number will be labelled. If ggrepel throws a warning, concerning overlap it is recommended to attempt to label less points to avoid cluttering
- **point.size**: Size of points in plots. Passed to geom_jitter()
- **mean.line.color**: Color of mean bar in dotplots. Passed to geom_errorbar()
- **stats.to.console**: Boolean. Defaults to FALSE. Prints basic statistics (AOV + post hoc test) to console
- **platypus.version**: Character. Only "v3" available.
Value

Returns a list of ggplot objects. out[[1]] is a boxplot comparing SHM by group.by. out[[2]] to out[[n]] are plots for each group that visualize VDJ and VJ SHM distribution for each group. Data for any plot can be accessed via out[[any]]$data

Examples

#Simulating SHM data
small_vgm <- Platypus::small_vgm
small_vgm[[1]]$VDJ_SHM <- as.integer(rnorm(nrow(small_vgm[[1]]), mean = 5, sd = 3))
small_vgm[[1]]$VJ_SHM <- as.integer(rnorm(nrow(small_vgm[[1]]), mean = 5, sd = 3))

#Standard plots
SHM_plots <- VDJ_plot_SHM(VDJ = small_vgm[[1]],
                          group.by = "sample_id", quantile.label = 0.9)

#Group by transcriptional cluster and label only top 1%
SHM_plots <- VDJ_plot_SHM(VDJ = small_vgm[[1]],
                          group.by = "seurat_clusters", quantile.label = 0.99)

VDJ_reclonotype_list_arrange

Platypus V2 dataframe utility

Description

Only Platypus v2 Organizes the top N genes that define each Seurat cluster and converts them into a single dataframe. This can be useful for obtaining insight into cluster-specific phenotypes.

Usage

VDJ_reclonotype_list_arrange(
  VDJ_clonotype.output,
  VDJ_analyze.output,
  Platypus_list.object
)

Arguments

VDJ_clonotype.output
  The output object from the VDJ_clonotype function. The column of the merged nucleotide clonotype IDs will be used to rearrange the new object.

VDJ_analyze.output
  The output from the initial VDJ_analyze, containing clonotype information based on nucleotide sequence.
Platypus_list.object

The new list object from one of Platypus functions (for example, clonal lineages, VDJ_per_clne, etc) that should be merged based on the VDJ_clonotype output structure. nested list structure, where outer list corresponds to repertoire and the inner list corresponds to clones (on the nucleotide level).

Value

Returns a dataframe in which the top N genes defining each cluster based on differential expression are selected.

Examples

```r
## Not run:
checking_vdj_reclono <- VDJ_reclonotype_list_arrange(
  VDJ_clonotype.output = repertoire_reclonotype
, VDJ_analyze.output = repertoire_list
, Platypus_list.object = repertoire_vdj_per_clone)
## End(Not run)
```

VDJ_tree

Platypus V2 phylogenetic trees.

### Description

Please refer tp VDJ_phylogenetic_tree for Platypus V3. Produces neighbor joining phylogenetic trees from the output of VDJ_clonal_lineages

### Usage

```r
VDJ_tree(
  clonal.lineages,
  with.germline,
  min.sequences,
  max.sequences,
  normalize.germline.length,
  unique.sequences
)
```

### Arguments

- **clonal.lineages**
  
  Output from VDJ_clonal_lineages. This should be nested list, with the outer list element corresponding to the individual repertoire and the inner list corresponding to individual clonal lineages based on the initial clonotyping strategy in the form of a dataframe with either Seq or Name.

- **with.germline**
  
  Logical specifying if the germline should be set as outgroup. Default is set to TRUE.
min.sequences  integer value specifying the minimum number of sequences to be allowed for clonal lineages. Default is 3.
max.sequences  integer value specifying the maximum number of sequences to be allowed for clonal lineages. Default is 500
normalize.germline.length  Logical determining whether or not the branch length separating the germline from the first internal node should be normalized. Potentially useful for visualization if the remainder tips are far from the root. Default is TRUE.
unique.sequences  Logical indicating if those cells containing identical VDJRegion sequences should be merged into single nodes and have their variant added as the tip label. Default is TRUE.

Value

Returns a nested list of phylogenetic trees. The output[[i]][[j]] corresponds to the j' th clone in the i' th input repertoire. plot(output[[i]][[j]]) should display the phylogenetic tree if the ape package is loaded.

Examples

```r
## Not run:
vdj.tree <- VDJ_tree(clonal.lineages = VDJ.clonal.lineage.output
,with.germline=TRUE,min.sequences = 5
,max.sequences = 30,unique.sequences = TRUE)
## End(Not run)
```

VDJ_variants_per_clone

Wrapper for variant analysis by clone

Description

Returns statistics and plots to examine diversity of any sequence or metadata item within clones on a by sample level or global level

Usage

```r
VDJ_variants_per_clone(
  VDJ,
  variants.of,
  clonotypes.col,
  stringDist.method,
  split.by,
  platypus.version
)
```
Argument

**VDJ**
VDJ output of the VDJ.GEX_matrix (VDJ.GEX_matrix.output[[1]]). VDJ matrix supplemented with with MIXCR information is also valid.

**variants.of**
Character vector. Defaults to c("VDJ_cdr3s_aa", "VJ_cdr3s_aa"). Column name(s) of VDJ to examine variants of. If more than one name is given, these columns will be pasted together. The default will therefore return statistics on the number of variants of VDJ and VJ cdr3s in every clone.

**clonotypes.col**
Column name of the VDJ column containing clonotype information. Defaults to "clonotype_id_10x". This is useful if alternative clonotyping strategies have been used and are stored in other columns.

**stringDist.method**
Character. Passed to Biostrings::strinDist. Method to calculate distance between variants of a clone. Defaults to "levenshtein". Other options are "hamming", "quality". If "hamming" variants of a clone will be shortened from the end to the shortest variant to make all input sequences the same length.

**split.by**
Character. Defaults to "sample_id". Column name of VDJ to split the analysis by. This is necessary, if clonotyping was done on a per sample level (e.g. "clonotype1" in sample 1 is not the same sequence as "clonotype1" in sample 2). If clonotyping was done across samples and no splitting is necessary input "none".

**platypus.version**
Character. Only "v3" available.

Value

Returns a list of dataframes. Each dataframe contains the statistics of one split.by element (by default: one sample).

Examples

```r
variants_per_clone <- VDJ_variants_per_clone(VDJ = Platypus::small_vgm[[1]],
                                           variants.of = c("VDJ_cdr3s_aa", "VJ_cdr3s_aa"),
                                           stringDist.method = "levenshtein", split.by = "sample_id")
```

---

**VDJ_Vgene_usage**

*V(D)J gene usage stacked barplots*

Description

Produces a matrix counting the number of occurrences for each VDJ and VJ Vgene combinations for each list entry in VDJ.clonotype.output or for each sample_id in VDJ.matrix.

Usage

```r
VDJ_Vgene_usage(VDJ, group.by, platypus.version)
```
Arguments

**VDJ**

For platypus.version = "v2" output from VDJ_analyze function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire. For platypus.version = "v3" output VDJ dataframe from VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]])

**group.by**

Character. Defaults to "sample_id". Column name of VDJ to group plot by.

**platypus.version**

Character. Defaults to "v3". Can be "v2" or "v3" dependent on the input format

Value

Returns a list of matrices containing the number of Vgene heavy/light chain combinations per repertoire.

Examples

```r
example.vdj.vgene_usage <- VDJ_Vgene_usage(VDJ = Platypus::small_vgm[[1]], platypus.version = "v3")
```

VDJ_Vgene_usage_barplot

**V(D)J gene usage barplots**

Description

Produces a barplot with the most frequently used IgH and IgK/L Vgenes.

Usage

```r
VDJ_Vgene_usage_barplot(
  VDJ,
  group.by,
  HC.gene.number,
  LC.Vgene,
  LC.gene.number,
  platypus.version,
  is.bulk
)
```

Arguments

**VDJ**

Either (for platypus version "v2") output from VDJ_analyze function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire, OR (for platypus version "v3") the the VDJ matrix output of the VDJ_GEX_matrix() function (VDJ.GEX.matrix.output[[1]])
**VDJ_Vgene_usage_stacked_barplot**

### Description

Produces a stacked barplot with the fraction of the most frequently used IgH and IgK/L V genes. This function can be used in combination with the `VDJ_Vgene_usage_barplot` to visualize V gene usage per sample and among samples.

### Usage

```
VDJ_Vgene_usage_stacked_barplot(
  VDJ,  
  group.by,  
  HC.gene.number,  
  Fraction.HC,  
  LC.Vgene,  
  LC.gene.number,  
  platypus.version
)
```

### Arguments

- **group.by**: Character. Defaults to "sample_id". Column name of VDJ to group plot by.

- **HC.gene.number**: Numeric value indicating the top genes to be displayed. If this number is higher than the total number of unique HC V genes in the VDJ repertoire, then this number is equal to the number of unique HC V genes.

- **LC.Vgene**: Logical indicating whether to make a barplot of the LC V genes distribution. Default is set to FALSE.

- **LC.gene.number**: Numeric value indicating the top genes to be displayed. If this number is higher than the total number of unique LC V genes in the VDJ repertoire, then this number is equal to the number of unique LC V genes.

- **platypus.version**: Character. Defaults to "v3". Can be "v2" or "v3" dependent on the input format.

- **is.bulk**: logical value indicating whether the VDJ input was generated from bulk-sequencing data using the bulk_to_vgm function. If is.bulk = T, the VDJ_Vgene_usage_barplot function is compatible for use with bulk data. Defaults to False (F).

### Value

Returns a list of ggplot objects which show the distribution of IgH and IgK/L V genes for the most used V genes.

### Examples

```
## Not run:
VDJ_Vgene_usage_barplot(VDJ = Platypus::small_vgm[[1]],
                        HC.gene.number = 2, platypus.version = "v3")

## End(Not run)
```
Arguments

**VDJ**
Either (for platypus version "v2") output from VDJ_analyze function. This should be a list of clonotype dataframes, with each list element corresponding to a single VDJ repertoire, OR (for platypus version "v3") the the VDJ matrix output of the VDJ_GEX_matrix() function (normally VDJ.GEX.matrix.output[[1]])

**group.by**
Character. Defaults to "sample_id". Column name of VDJ to group plot by.

**HC.gene.number**
Numeric value indicating the top genes to be displayed. If this number is higher than the total number of unique HC V genes in the VDJ repertoire, then this number is equal to the number of unique HC V genes.

**Fraction.HC**
Numeric value indicating the minimum fraction of clones expressing a particular HC V gene. If the usage of a particular gene is below this value, then this gene is excluded. If the usage of a particular gene is above this value even in one sample, then this gene is included in the analysis. Default value is set to 0, thus all genes are selected.

**LC.Vgene**
Logical indicating whether to make a barplot of the LC V gene distribution. Default is set to FALSE.

**LC.gene.number**
Numeric value indicating the top genes to be displayed. If this number is higher than the total number of unique LC V genes in the VDJ repertoire, then this number is equal to the number of unique LC V genes.

**Fraction.LC**
Numeric value indicating the minimum fraction of clones expressing a particular LC V gene. If the usage of a particular gene is below this value, then this gene is excluded. If the usage of a particular gene is above this value even in one sample, then this gene is included in the analysis. Default value is set to 0, thus all genes are selected.

**platypus.version**
Set according to input format to either "v2" or "v3". Defaults to "v3"

**is.bulk**
logical value indicating whether the VDJ input was generated from bulk-sequencing data using the bulk_to_vgm function. If is.bulk = T, the VDJ_Vgene_usage_stacked_barplot function is compatible for use with bulk data. Defaults to False (F).

Value

Returns a list of ggplot objects which show the stacked distribution of IgH and IgK/L V genes for the most used V genes. Returns an empty plot if the Fraction.HC or Fraction.LC that were selected were too high, resulting in the exclusion of all the genes.

Examples

```r
#Platypus v3
eexample.vdj.vgene_usage <- VDJ_Vgene_usage_stacked_barplot(
```
VDJ_VJ_usage_circos

vdj = Platypus::small_vgm[[1]], LC.Vgene = TRUE, HC.gene.number = 15, Fraction.HC = 1, platypus.version = "v3"

VDJ_VJ_usage_circos  Makes a Circos plot from the VDJ_analyze output. Connects the V gene with the corresponding J gene for each clonotype.

Description

Makes a Circos plot from the VDJ_analyze output. Connects the V gene with the corresponding J gene for each clonotype.

Usage

VDJ_VJ_usage_circos(  
VDJ,  
A.or.B,  
label.threshold,  
cell.level,  
c.threshold,  
clonotype.per.gene.threshold,  
c.count,  
platypus.version,  
filter1H1L  )

Arguments

VDJ  The output of the VDJ_GEX_integrate function (Platypus platypus.version v2). A list of data frames for each sample containing the clonotype information and cluster membership information. For Platypus platypus.version v3, the VDJ output of the VDJ_GEX_matrix function (VDJ_GEX_matrix.output[[1]]) has to be supplied.

A.or.B  Determines whether to plot the V J gene pairing of the alpha or beta chain. "A", "B" or "both" as possible inputs. Default: "both".

label.threshold  Minimal amount of clonotypes per gene necessary to add a gene label to the sector. Default: 0.

cell.level  Logical, defines whether weight of connection should be based on number of clonotypes or number of cells. Default: number of clonotypes.

c.threshold  Only clonotypes are considered with a frequency higher than c.threshold. Allows to filter for only highly expanded clonotypes.

clonotype.per.gene.threshold  How many clonotypes are required to plot a sector for a gene. Filters the rows and columns of the final adjacency matrix.

filter1H1L  Any other arguments passed will be removed from the final adjacency matrix.
c.count                  Show clonotype or cell count on Circos plot. Default = T.
platypus.version        Which platypus.version of platypus is being used. Default = v3. Set to v3 if
                        VDJ_GEX_matrix.output[[1]] is used
filter1H1L               Whether to filter the input VDJ in "v3" to only include cells with 1 VDJ and 1
                        VJ chain. Defaults to TRUE

Value
Returns list of plots. The first n elements contain the circos plot of the n datasets from the VDJ.analyze
function. The n+1 element contains a list of the n adjacency matrices for each dataset.

Examples
## Not run:
plots <- VDJ_VJ_usage_circos(VDJ = Platypus::small_vgm[[1]], platypus.version = "v3",
cell.level = TRUE)
## End(Not run)

VGM_expanded_clones      VDJ utility for T/F column for clonal expansion

Description
Adds discrete columns containing TRUE / FALSE on whether a given cell is part of a expanded or
not-expanded clonotype. Threshold frequency can be set.

Usage
VGM_expanded_clones(VGM, add.to.VDJ, add.to.GEX, expansion.threshold)

Arguments
VGM                     Output object from the VDJ_GEX_matrix function (VDJ_GEX_matrix.output)
add.to.VDJ              Boolean. Whether to add expanded columns to VDJ matrix. Defaults to TRUE
add.to.GEX              Boolean. Whether to add expanded columns to GEX matrix. Defaults to TRUE
expansion.threshold      Integer. Defaults to 1. Cells in clonotypes above this threshold will be marked
                        as expanded = TRUE.

Value
An output object from the VDJ_GEX_matrix function with added columns containing TRUE / FALSE values based on clonotype frequency.
Examples

```r
#Add info to whole VGM object
VGM <- VGM_expanded_clones(
  VGM = Platypus::small_vgm, add.to.VDJ = TRUE, add.to.GEX = TRUE,
  expansion.threshold = 1)
```

**Description**

The `VGM_expand_featurebarcodes` function can be used to trace back the cell origin of each sample after using cell hashing for single-cell sequencing. Replaces the original sample_id column of a vgm object with a pasted version of the original sample_id and the last digits of the feature barcode.

The original sample_id is stored in a new column called `original_sample_id`. Additionally, a second new column is created containing final barcode assignment information. Those barcodes match the origin `FB_assignment` if `by.majority.barcodes` is set to `FALSE` (default). However, if this input parameter is set to `TRUE`, the majority barcode assignment in stored in this column.

Note: The majority barcode of a cell is the feature barcode which is most frequently assigned to the cells clonotype (10x default clonotype). The majority barcode assignment can be used under the assumption that all cells which are assigned to the same clonotype (within one sample), originate from the same donor organ or at least the same donor depending on the experimental setup.

For example: The original sample_id of a cell is "s1", the cell belongs to "clonotype1" and the feature barcode assigned to it is "i1-TotalSeq-C0953". If `by.majority.barcodes` default (`FALSE`) is used, the resulting new sample_id would be "s1_0953". However, if majority barcode assignment is used AND "i1-TotalSeq-C0953" is not the most frequently occurring barcode in "clonotype1" but rather barcode "i1-TotalSeq-C0951", the new sample_id would be "s1_0951". -> e.g., if 15 cells belong to clonotype1: 3 cells have no assigned barcode, 2 are assigned to "i1-TotalSeq-C0953" and 10 are assigned to "i1-TotalSeq-C0951" -> all 15 cells will have the new sample_id "s1_0951".

**Usage**

```r
VGM_expand_featurebarcodes(
  vgm,
  by.majority.barcodes,
  integrate.in.gex,
  vdj.only,
  platypus.version
)
```
**Arguments**

- **vgm**: VGM output of VDJ_GEX_matrix function (Platypus V3)
- **by.majority.barcodes**: Logical. Default is FALSE. Indicated whether strict barcode assignment or majority barcode assignment should be used to create the new sample_id. If TRUE, for each clonotype the most frequent feature barcode will be chosen and assigned to each cell, even if that cell itself does not have this particular barcode assigned.
- **integrate.in.gex**: Logical. Default is FALSE. If TRUE, the newly created sample_id’s are integrated into gex component as well. Not recommended if no further gex analysis is done due to much longer computational time.
- **vdj.only**: Logical. Defines if only vdj information is provided as input. Default is set to FALSE. If set to TRUE a vdj dataframe has to be provided as input (vgm = vdj). Also, integrate.in.gex is automatically set to FALSE since no gex (vgm[2]) information is provided.
- **platypus.version**: This function works with "v3" only, there is no need to set this parameter.

**Value**

This function returns a vgm with new sample_id’s in case vdj.only is set to FALSE (default). If vdj.only is set to true only the vdj dataframe with new sample_id’s is returned. Note: If vdj.only is set to default (FALSE), VDJ information in the metadata of the GEX object is necessary. For this set integrate.VDJ.to.GEX to TRUE in the VDJ_GEX_matrix function.

**Examples**

#For Platypus version 3

# 1. If only vdj data (vgm[1]) and strict feature barcode assignment is used:
vgm_expanded_fb <- VGM_expand_featurebarcodes(
  vgm = small_vgm[1],
  by.majority.barcodes = FALSE,
  integrate.in.gex=FALSE, vdj.only= TRUE)

# 2. If whole vgm and strict fb assignment is used
#(gex and vdj - necessary if gene expression analysis # of sub-samples is desired):
vgm_expanded_fb <- VGM_expand_featurebarcodes(
  vgm = small_vgm,
  by.majority.barcodes = FALSE,
  integrate.in.gex=TRUE, vdj.only= FALSE)

# 3. If whole vgm and majority barcode assignment is used
#(gex and vdj) - necessary if gene expression analysis # of sub-samples is desired):
vgm_expanded_fb <- VGM_expand_featurebarcodes(vgm = small_vgm,
by.majority.barcodes = TRUE,
integrate.in.gex=TRUE, vdj.only= FALSE)

#Note: Majority barcode assignment is recommended
#if the assumption that all cells within one clonotype
#originate from the same sample sub-group is feasible.

---

**VGM_integrate**  
*Utility for VDJ GEX matrix to integrated VDJ and GEX objects after addition of data to either*

---

**Description**

(Re)-intergrated VDJ and GEX of one or two separate VGM objects. This can be used as a simple “updating” utility function, if metadata has been added to the VDJ dataframe and is also needed in the GEX matrix or the reverse. Entries are integrated by barcode. If barcodes have been altered (barcode column in VDJ and cell names in GEX), the function will not yield results

**Usage**

```r
VGM_integrate(VGM, columns.to.transfer, genes.to.VDJ, seurat.slot)
```

**Arguments**

- **VGM**: Output object from the VDJ_GEX_matrix function (VDJ_GEX_matrix.output)
- **columns.to.transfer**: Optional. Character Vector. Column names of either the VDJ matrix or GEX meta.data that should be transferred to the corresponding other matrix. If not provided all columns missing from one will be integrated into the other matrix.
- **genes.to.VDJ**: Character vector of gene names in GEX. In many cases it is useful to extract expression values for a gene to metadata. This is done via SeuratObject::FetchData(vars = genes, slot = seurat.slot) function. The VGM integrate takes gene ids, extracts these and adds them to the VDJ dataframe. If provided, no other columns are integrated between VDJ and GEX and columns.to.transfer is ignored.
- **seurat.slot**: GEX object data slot to pull from. Can be 'counts', 'data', or 'scale.data'

**Value**

An output object from the VDJ_GEX_matrix function with added columns in VDJ or GEX

**Examples**

```r
#Adding a new clonotyping method to VDJ
small_vgm[[1]] <- VDJ_clonotype_v3(VDJ=Platypus::small_vgm[[1]],
clonestrategy="cdr3.nt",
hierarchical = "single.chains", global.clonotype = TRUE)
```
small_vgm <- VGM_integrate(
  VGM = small_vgm,
  columns.to.transfer = NULL)  # transfer all new columns
# and update clonotype_id and clonotype_frequency column
# (as does VDJ_clonotype_v3 in VDJ)

small_vgm <- VGM_integrate(
  VGM = small_vgm,
  columns.to.transfer = c("global_clonotype_id_cdr3.nt"))
# transfer only selected columns

# Pull genes from GEX and add as metadata column to VDJ
small_vgm <- VGM_integrate(
  small_vgm, genes.to.VDJ = c("CD19", "CD24A"), seurat.slot = "counts")
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