Package ‘scoper’

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Title Spectral Clustering-Based Method for Identifying B Cell Clones
Description Provides a computational framework for identification of B cell clones from Adaptive Immune Receptor Repertoire sequencing (AIRR-Seq) data. Three main functions are included (identicalClones, hierarchicalClones, and spectralClones) that perform clustering among sequences of BCRs/IGs (B cell receptors/immunoglobulins) which share the same V gene, J gene and junction length.
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ExampleDb

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**ExampleDb**  

**Example database**

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**Description**

A small example database subset from Laserson and Vigneault et al, 2014.

**Usage**

ExampleDb

**Format**

A data.frame with the following columns:

- sequence_id: Sequence identifier
- sequence_alignment: IMGT-gapped observed sequence.
- germline_alignment: IMGT-gapped germline sequence.
- germline_alignment_d_mask: IMGT-gapped germline sequence with N, P and D regions masked.
- v_call: V region allele assignments.
- v_call_genotyped: TiGER corrected V region allele assignment.
- d_call: D region allele assignments.
hierarchicalClones

- j_call: J region allele assignments.
- junction: Junction region sequence.
- junction_length: Length of the junction region in nucleotides.
- np1_length: Number of nucleotides between V and D segments
- np2_length: Number of nucleotides between D and J segments
- sample_id: Sample identifier
- c_call: C region assignment.
- duplicate_count: Copy number of the sequence
- locus: Locus of the receptor

References

hierarchicalClones  
Hierarchical clustering method for clonal partitioning

Description
hierarchicalClones provides a hierarchical agglomerative clustering approach to infer clonal relationships in high-throughput Adaptive Immune Receptor Repertoire sequencing (AIRR-seq) data. This approach clusters B or T cell receptor sequences based on junction region sequence similarity within partitions that share the same V gene, J gene, and junction length, allowing for ambiguous V or J gene annotations.

Usage
hierarchicalClones(
  db,
  threshold,
  method = c("nt", "aa"),
  linkage = c("single", "average", "complete"),
  normalize = c("len", "none"),
  junction = "junction",
  v_call = "v_call",
  j_call = "j_call",
  clone = "clone_id",
  fields = NULL,
  cell_id = NULL,
  locus = "locus",
  only_heavy = TRUE,
  split_light = TRUE,
  first = FALSE,
  cdr3 = FALSE,
hierarchicalClones

mod3 = FALSE,
max_n = 0,
nproc = 1,
verbose = FALSE,
log = NULL,
summarize_clones = TRUE
)

Arguments
db data.frame containing sequence data.
threshold numeric scalar where the tree should be cut (the distance threshold for clonal grouping).
method one of the "nt" for nucleotide based clustering or "aa" for amino acid based clustering.
linkage available linkage are "single", "average", and "complete".
normalize method of normalization. The default is "len", which divides the distance by the length of the sequence group. If "none" then no normalization if performed.
junction character name of the column containing junction sequences. Also used to determine sequence length for grouping.
v_call name of the column containing the V-segment allele calls.
j_call name of the column containing the J-segment allele calls.
clone output column name containing the clonal cluster identifiers.
fields character vector of additional columns to use for grouping. Sequences with disjoint values in the specified fields will be classified as separate clones.
cell_id name of the column containing cell identifiers or barcodes. If specified, grouping will be performed in single-cell mode with the behavior governed by the locus and only_heavy arguments. If set to NULL then the bulk sequencing data is assumed.
locus name of the column containing locus information. Only applicable to single-cell data. Ignored if cell_id=NULL.
only_heavy use only the IGH (BCR) or TRB/TRD (TCR) sequences for grouping. Only applicable to single-cell data. Ignored if cell_id=NULL.
split_light split clones by light chains. Ignored if cell_id=NULL.
first specifies how to handle multiple V(D)J assignments for initial grouping. If TRUE only the first call of the gene assignments is used. If FALSE the union of ambiguous gene assignments is used to group all sequences with any overlapping gene calls.
cdr3 if TRUE removes 3 nucleotides from both ends of "junction" prior to clustering (converts IMGT junction to CDR3 region). If TRUE this will also remove records with a junction length less than 7 nucleotides.
mod3 if TRUE removes records with a junction length that is not divisible by 3 in nucleotide space.
max_n  The maximum number of degenerate characters to permit in the junction sequence before excluding the record from clonal assignment. Note, with linkage="single" non-informative positions can create artifactual links between unrelated sequences. Use with caution. Default is set to be zero. Set it as "NULL" for no action.

nproc  number of cores to distribute the function over.

verbose  if TRUE prints out a summary of each step cloning process. if FALSE (default) process cloning silently.

log  output path and filename to save the verbose log. The input file directory is used if path is not specified. The default is NULL for no action.

summarize_clones  if TRUE performs a series of analysis to assess the clonal landscape and returns a ScoperClones object. If FALSE then a modified input db is returned. When grouping by fields, summarize_clones should be FALSE.

Value

If summarize_clones=TRUE (default) a ScoperClones object is returned that includes the clonal assignment summary information and a modified input db in the db slot that contains clonal identifiers in the specified clone column. If summarize_clones=FALSE modified data.frame is returned with clone identifiers in the specified clone column.

Single-cell data

To invoke single-cell mode the cell_id argument must be specified and the locus column must be correct. Otherwise, clustering will be performed with bulk sequencing assumptions, using all input sequences regardless of the values in the locus column.

Values in the locus column must be one of c("IGH", "IGI", "IGK", "IGL") for BCR or c("TRA", "TRB", "TRD", "TRG") for TCR sequences. Otherwise, the operation will exit and return an error message.

Under single-cell mode with paired-chain sequences, there is a choice of whether grouping should be done by (a) using IGH (BCR) or TRB/TRD (TCR) sequences only or (b) using IGH plus IGK/IGL (BCR) or TRB/TRD plus TRA/TRG (TCR) sequences. This is governed by the only_heavy argument. There is also choice as to whether inferred clones should be split by the light/short chain (IGK, IGL, TRA, TRG) following heavy/long chain clustering, which is governed by the split_light argument.

In single-cell mode, clonal clustering will not be performed on data where cells are assigned multiple heavy/long chain sequences (IGH, TRB, TRD). If observed, the operation will exit and return an error message. Cells that lack a heavy/long chain sequence (i.e., cells with light/short chains only) will be assigned a clone_id of NA.

See Also

See plotCloneSummary for plotting summary results. See groupGenes for more details about grouping requirements.
**Examples**

```r
# Find clonal groups
results <- hierarchicalClones(ExampleDb, threshold=0.15)

# Retrieve modified input data with clonal clustering identifiers
df <- as.data.frame(results)

# Plot clonal summaries
plot(results, binwidth=0.02)
```

---

**identicalClones**

Sequence identity method for clonal partitioning

**Description**

identicalClones provides a simple sequence identity based partitioning approach for inferring clonal relationships in high-throughput Adaptive Immune Receptor Repertoire sequencing (AIRR-seq) data. This approach partitions B or T cell receptor sequences into clonal groups based on junction region sequence identity within partitions that share the same V gene, J gene, and junction length, allowing for ambiguous V or J gene annotations.

**Usage**

```r
identicalClones(
  db,
  method = c("nt", "aa"),
  junction = "junction",
  v_call = "v_call",
  j_call = "j_call",
  clone = "clone_id",
  fields = NULL,
  cell_id = NULL,
  locus = "locus",
  only_heavy = TRUE,
  split_light = TRUE,
  first = FALSE,
  cdr3 = FALSE,
  mod3 = FALSE,
  max_n = 0,
  nproc = 1,
  verbose = FALSE,
  log = NULL,
  summarize_clones = TRUE
)
```
**Arguments**

- **db**: data.frame containing sequence data.
- **method**: one of the "nt" for nucleotide based clustering or "aa" for amino acid based clustering.
- **junction**: character name of the column containing junction sequences. Also used to determine sequence length for grouping.
- **v_call**: name of the column containing the V-segment allele calls.
- **j_call**: name of the column containing the J-segment allele calls.
- **clone**: output column name containing the clonal cluster identifiers.
- **fields**: character vector of additional columns to use for grouping. Sequences with disjoint values in the specified fields will be classified as separate clones.
- **cell_id**: name of the column containing cell identifiers or barcodes. If specified, grouping will be performed in single-cell mode with the behavior governed by the **locus** and **only_heavy** arguments. If set to NULL then the bulk sequencing data is assumed.
- **locus**: name of the column containing locus information. Only applicable to single-cell data. Ignored if cell_id=NULL.
- **only_heavy**: use only the IGH (BCR) or TRB/TRD (TCR) sequences for grouping. Only applicable to single-cell data. Ignored if cell_id=NULL.
- **split_light**: split clones by light chains. Ignored if cell_id=NULL.
- **first**: specifies how to handle multiple V(D)J assignments for initial grouping. If TRUE only the first call of the gene assignments is used. If FALSE the union of ambiguous gene assignments is used to group all sequences with any overlapping gene calls.
- **cdr3**: if TRUE removes 3 nucleotides from both ends of "junction" prior to clustering (converts IMGT junction to CDR3 region). If TRUE this will also remove records with a junction length less than 7 nucleotides.
- **mod3**: if TRUE removes records with a junction length that is not divisible by 3 in nucleotide space.
- **max_n**: The maximum number of degenerate characters to permit in the junction sequence before excluding the record from clonal assignment. Default is set to be zero. Set it as "NULL" for no action.
- **nproc**: number of cores to distribute the function over.
- **verbose**: if TRUE prints out a summary of each step cloning process. if FALSE (default) process cloning silently.
- **log**: output path and filename to save the verbose log. The input file directory is used if path is not specified. The default is NULL for no action.
- **summarize_clones**: if TRUE performs a series of analysis to assess the clonal landscape and returns a ScoperClones object. If FALSE then a modified input db is returned. When grouping by fields, summarize_clones should be FALSE.
Value

If `summarize_clones=TRUE` (default) a `ScoperClones` object is returned that includes the clonal assignment summary information and a modified input `db` in the `db` slot that contains clonal identifiers in the specified `clone` column. If `summarize_clones=FALSE` modified data.frame is returned with clone identifiers in the specified clone column.

Single-cell data

To invoke single-cell mode the `cell_id` argument must be specified and the `locus` column must be correct. Otherwise, clustering will be performed with bulk sequencing assumptions, using all input sequences regardless of the values in the `locus` column.

Values in the `locus` column must be one of c("IGH", "IGI", "IGK", "IGL") for BCR or c("TRA", "TRB", "TRD", "TRG") for TCR sequences. Otherwise, the operation will exit and return an error message.

Under single-cell mode with paired-chain sequences, there is a choice of whether grouping should be done by (a) using IGH (BCR) or TRB/TRD (TCR) sequences only or (b) using IGH plus IGK/IGL (BCR) or TRB/TRD plus TRA/TRG (TCR) sequences. This is governed by the `only_heavy` argument. There is also choice as to whether inferred clones should be split by the light/short chain (IGK, IGL, TRA, TRG) following heavy/long chain clustering, which is governed by the `split_light` argument.

In single-cell mode, clonal clustering will not be performed on data where cells are assigned multiple heavy/long chain sequences (IGH, TRB, TRD). If observed, the operation will exit and return an error message. Cells that lack a heavy/long chain sequence (i.e., cells with light/short chains only) will be assigned a `clone_id` of NA.

See Also

See `plotCloneSummary` for plotting summary results. See `groupGenes` for more details about grouping requirements.

Examples

```r
# Find clonal groups
results <- identicalClones(ExampleDb)

# Retrieve modified input data with clonal clustering identifiers
df <- as.data.frame(results)

# Plot clonal summaries
plot(results, binwidth=0.02)
```
plotCloneSummary

Description

plotCloneSummary plots the results in a ScoperClones object returned by spectralClones, identicalClones or hierarchicalClones. Includes the minimum inter (between) and maximum intra (within) clonal distances and the calculated effective threshold.

Usage

plotCloneSummary(
  data,
  xmin = NULL,
  xmax = NULL,
  breaks = NULL,
  binwidth = NULL,
  title = NULL,
  size = 0.75,
  silent = FALSE,
  ...
)

Arguments

data
  ScoperClones object output by the spectralClones, identicalClones or hierarchicalClones.

xmin
  minimum limit for plotting the x-axis. If NULL the limit will be set automatically.

xmax
  maximum limit for plotting the x-axis. If NULL the limit will be set automatically.

breaks
  number of breaks to show on the x-axis. If NULL the breaks will be set automatically.

binwidth
  binwidth for the histogram. If NULL the binwidth will be set automatically.

title
  string defining the plot title.

size
  numeric value for lines in the plot.

silent
  if TRUE do not draw the plot and just return the ggplot2 object; if FALSE draw the plot.

...
  additional arguments to pass to ggplot2::theme.

Value

A ggplot object defining the plot.

See Also

See ScoperClones for the the input object definition. See spectralClones, identicalClones and hierarchicalClones for generating the input object.
Examples

```r
# Find clones
results <- hierarchicalClones(ExampleDb, threshold=0.15)

# Plot clonal summaries
plot(results, binwidth=0.02)
```

---

**scoper**  
*The SCOPer package*

**Description**

scoper is a member of the Immcantation framework and provides computational approaches for the identification of B cell clones from adaptive immune receptor repertoire sequencing (AIRR-Seq) datasets. It includes methods for assigning clonal identifiers using sequence identity, hierarchical clustering, and spectral clustering.

**Clonal clustering**

- `identicalClones`: Clonal assignment using sequence identity partitioning.
- `hierarchicalClones`: Hierarchical clustering approach to clonal assignment.
- `spectralClones`: Spectral clustering approach to clonal assignment.

**Visualization**

- `plotCloneSummary`: Visualize inter- and intra-clone distances.

**References**

ScoperClones-class

S4 class containing clonal assignments and summary data

Description

ScoperClones stores output from identicalClones, hierarchicalClones and spectralClones functions.

Usage

## S4 method for signature 'ScoperClones'
print(x)

## S4 method for signature 'ScoperClones'
summary(object)

## S4 method for signature 'ScoperClones,missing'
plot(x, y, ...)

## S4 method for signature 'ScoperClones'
as.data.frame(x)

Arguments

x ScoperClones object
object ScoperClones object
y ignored.
... arguments to pass to plotCloneSummary.

Slots

db data.frame of repertoire data including with clonal identifiers in the column specified during processing.
vjl_groups data.frame of clonal summary, including sequence count, V gene, J gene, junction length, and clone counts.
inter_intra data.frame containing minimum inter (between) and maximum intra (within) clonal distances.
eff_threshold effective cut-off separating the inter (between) and intra (within) clonal distances.

See Also

identicalClones, hierarchicalClones and spectralClones
spectralClones provides an unsupervised spectral clustering approach to infer clonal relationships in high-throughput Adaptive Immune Receptor Repertoire sequencing (AIRR-seq) data. This approach clusters B or T cell receptor sequences based on junction region sequence similarity and shared mutations within partitions that share the same V gene, J gene, and junction length, allowing for ambiguous V or J gene annotations.

Usage

```r
spectralClones(
  db,
  method = c("novj", "vj"),
  germline = "germline_alignment",
  sequence = "sequence_alignment",
  junction = "junction",
  v_call = "v_call",
  j_call = "j_call",
  clone = "clone_id",
  fields = NULL,
  cell_id = NULL,
  locus = "locus",
  only_heavy = TRUE,
  split_light = TRUE,
  targeting_model = NULL,
  len_limit = NULL,
  first = FALSE,
  cdr3 = FALSE,
  mod3 = FALSE,
  max_n = 0,
  threshold = NULL,
  base_sim = 0.95,
  iter_max = 1000,
  nstart = 1000,
  nproc = 1,
  verbose = FALSE,
  log = NULL,
  summarize_clones = TRUE
)
```

Arguments

db data.frame containing sequence data.
method one of the "novj" or "vj". See Details for description.
**spectralClones**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>germline</td>
<td>character name of the column containing the germline or reference sequence.</td>
</tr>
<tr>
<td>sequence</td>
<td>character name of the column containing input sequences.</td>
</tr>
<tr>
<td>junction</td>
<td>character name of the column containing junction sequences. Also used to determine sequence length for grouping.</td>
</tr>
<tr>
<td>v_call</td>
<td>name of the column containing the V-segment allele calls.</td>
</tr>
<tr>
<td>j_call</td>
<td>name of the column containing the J-segment allele calls.</td>
</tr>
<tr>
<td>clone</td>
<td>output column name containing the clonal cluster identifiers.</td>
</tr>
<tr>
<td>fields</td>
<td>character vector of additional columns to use for grouping. Sequences with disjoint values in the specified fields will be classified as separate clones.</td>
</tr>
<tr>
<td>cell_id</td>
<td>name of the column containing cell identifiers or barcodes. If specified, grouping will be performed in single-cell mode with the behavior governed by the <code>locus</code> and <code>only_heavy</code> arguments. If set to <code>NULL</code> then the bulk sequencing data is assumed.</td>
</tr>
<tr>
<td>locus</td>
<td>name of the column containing locus information. Only applicable to single-cell data. Ignored if <code>cell_id=NULL</code>.</td>
</tr>
<tr>
<td>only_heavy</td>
<td>use only the IGH (BCR) or TRB/TRD (TCR) sequences for grouping. Only applicable to single-cell data. Ignored if <code>cell_id=NULL</code>.</td>
</tr>
<tr>
<td>split_light</td>
<td>split clones by light chains. Ignored if <code>cell_id=NULL</code>.</td>
</tr>
<tr>
<td>targeting_model</td>
<td><code>TargetingModel</code> object. Only applicable if <code>method=&quot;vj&quot;</code>. See Details for description.</td>
</tr>
<tr>
<td>len_limit</td>
<td><code>IMGT_V</code> object defining the regions and boundaries of the Ig sequences. If <code>NULL</code>, mutations are counted for entire sequence. Only applicable if <code>method = &quot;vj&quot;</code>.</td>
</tr>
<tr>
<td>first</td>
<td>specifies how to handle multiple V(D)J assignments for initial grouping. If <code>TRUE</code> only the first call of the gene assignments is used. If <code>FALSE</code> the union of ambiguous gene assignments is used to group all sequences with any overlapping gene calls.</td>
</tr>
<tr>
<td>cdr3</td>
<td>if <code>TRUE</code> removes 3 nucleotides from both ends of &quot;junction&quot; prior to clustering (converts IMGT junction to CDR3 region). If <code>TRUE</code> this will also remove records with a junction length less than 7 nucleotides.</td>
</tr>
<tr>
<td>mod3</td>
<td>if <code>TRUE</code> removes records with a junction length that is not divisible by 3 in nucleotide space.</td>
</tr>
<tr>
<td>max_n</td>
<td>the maximum number of degenerate characters to permit in the junction sequence before excluding the record from clonal assignment. Default is set to be zero. Set it as &quot;NULL&quot; for no action.</td>
</tr>
<tr>
<td>threshold</td>
<td>the supervising cut-off to enforce an upper-limit distance for clonal grouping. A numeric value between (0,1).</td>
</tr>
<tr>
<td>base_sim</td>
<td>required similarity cut-off for sequences in equal distances from each other.</td>
</tr>
<tr>
<td>iter_max</td>
<td>the maximum number of iterations allowed for kmean clustering step.</td>
</tr>
<tr>
<td>nstart</td>
<td>the number of random sets chosen for kmean clustering initialization.</td>
</tr>
<tr>
<td>nproc</td>
<td>number of cores to distribute the function over.</td>
</tr>
</tbody>
</table>
verbose if TRUE prints out a summary of each step cloning process. if FALSE (default) process cloning silently.

log output path and filename to save the verbose log. The input file directory is used if path is not specified. The default is NULL for no action.

summarize_clones if TRUE performs a series of analysis to assess the clonal landscape and returns a `ScoperClones` object. If FALSE then a modified input db is returned. When grouping by fields, summarize_clones should be FALSE.

Details

If method="novj", then clonal relationships are inferred using an adaptive threshold that indicates the level of similarity among junction sequences in a local neighborhood.

If method="vj", then clonal relationships are inferred not only on junction region homology, but also taking into account the mutation profiles in the V and J segments. Mutation counts are determined by comparing the input sequences (in the column specified by sequence) to the effective germline sequence (IUPAC representation of sequences in the column specified by germline).

While not mandatory, the influence of SHM hot-cold-spot biases in the clonal inference process will be noted if a SHM targeting model is provided through the targeting_model argument. See TargetingModel for more technical details.

If the threshold argument is specified, then an upper limit for clonal grouping will be imposed to prevent sequences with dissimilarity above the threshold from grouping together. Any sequence with a distance greater than the threshold value from the other sequences, will be assigned to a singleton group.

Value

If summarize_clones=TRUE (default) a `ScoperClones` object is returned that includes the clonal assignment summary information and a modified input db in the db slot that contains clonal identifiers in the specified clone column. If summarize_clones=FALSE modified data.frame is returned with clone identifiers in the specified clone column.

Single-cell data

To invoke single-cell mode the cell_id argument must be specified and the locus column must be correct. Otherwise, clustering will be performed with bulk sequencing assumptions, using all input sequences regardless of the values in the locus column.

Values in the locus column must be one of c("IGH", "IGI", "IGK", "IGL") for BCR or c("TRA", "TRB", "TRD", "TRG") for TCR sequences. Otherwise, the operation will exit and return an error message.

Under single-cell mode with paired-chain sequences, there is a choice of whether grouping should be done by (a) using IGH (BCR) or TRB/TRD (TCR) sequences only or (b) using IGH plus IGK/IGL (BCR) or TRB/TRD plus TRA/TRG (TCR) sequences. This is governed by the only_heavy argument. There is also choice as to whether inferred clones should be split by the light/short chain (IGK, IGL, TRA, TRG) following heavy/long chain clustering, which is governed by the split_light argument.
In single-cell mode, clonal clustering will not be performed on data were cells are assigned multiple heavy/long chain sequences (IGH, TRB, TRD). If observed, the operation will exit and return an error message. Cells that lack a heavy/long chain sequence (i.e., cells with light/short chains only) will be assigned a clone_id of NA.

See Also

See `plotCloneSummary` for plotting summary results. See `groupGenes` for more details about grouping requirements.

Examples

```r
# Subset example data
db <- subset(ExampleDb, c_call == "IGHG")

# Find clonal groups
results <- spectralClones(db, method="novj", germline="germline_alignment_d_mask")

# Retrieve modified input data with clonal clustering identifiers
df <- as.data.frame(results)

# Plot clonal summaries
plot(results, binwidth=0.02)
```
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