Package ‘shazam’

February 6, 2020

Type Package
Version 0.2.3
Date 2020-02-05
Title Immunoglobulin Somatic Hypermutation Analysis
Description Provides a computational framework for analyzing mutations in immunoglobulin (Ig) sequences. Includes methods for Bayesian estimation of antigen-driven selection pressure, mutational load quantification, building of somatic hypermutation (SHM) models, and model-dependent distance calculations. Also includes empirically derived models of SHM for both mice and humans. Citations:
License CC BY-SA 4.0
URL http://shazam.readthedocs.io
BugReports https://bitbucket.org/kleinstein/shazam/issues
LazyData true
BuildVignettes true
VignetteBuilder knitr
Encoding UTF-8
Depends R (>= 3.1.2), ggplot2 (>= 2.0.0), stringi (>= 1.1.3)
Imports alakazam (>= 0.3.0), ape, diptest, doParallel, dplyr (>=
0.8.1), foreach, graphics, grid, igraph, iterators, kedd,
KernSmooth, lazyeval, MASS, methods, parallel, progress, rlang,
scales, seqinr, stats, tidyr, utils
Suggests knitr, rmarkdown, testthat
Collate 'Shazam.R' 'RegionDefinitions.R' 'Baseline.R' 'Core.R'
'DistToNearest.R' 'MutationDefinitions.R' 'MutationProfiling.R'
'Shmulate.R' 'TargetingModels.R'
RoxygenNote 7.0.2

NeedsCompilation no

Author Mohamed Uduman [aut],
Gur Yaari [aut],
Namita Gupta [aut],
Jason Vander Heiden [aut, cre],
Ang Cui [ctb],
Susanna Marquez [ctb],
Julian Zhou [ctb],
Nima Nouri [ctb],
Steven Kleinstein [aut, cph]

Maintainer Jason Vander Heiden <jason.vanderheiden@yale.edu>

Repository CRAN

Date/Publication 2020-02-06 06:20:06 UTC

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**Description**

Baseline defines a common data structure the results of selection analysis using the BASELINe method.

**Usage**

```r
## S4 method for signature 'Baseline,character'
plot(x, y, ...)

## S4 method for signature 'Baseline'
summary(object, nproc = 1)
```
Arguments

- `x`: Baseline object.
- `y`: name of the column in the db slot of baseline containing primary identifiers.
- `...`: arguments to pass to `plotBaselineDensity`.
- `object`: Baseline object.
- `nproc`: number of cores to distribute the operation over.

Slots

- `description`: character providing general information regarding the sequences, selection analysis and/or object.
- `db`: data.frame containing annotation information about the sequences and selection results.
- `regionDefinition`: RegionDefinition object defining the regions and boundaries of the Ig sequences.
- `testStatistic`: character indicating the statistical framework used to test for selection. For example, "local" or "focused".
- `regions`: character vector defining the regions the BASELINE analysis was carried out on. For "CDR" and "FWR" or "CDR1", "CDR2", "CDR3", etc.
- `numbOfSeqs`: matrix of dimensions r x c containing the number of sequences or PDFs in each region, where:
  - r = number of rows = number of groups or sequences.
  - c = number of columns = number of regions.
- `binomK`: matrix of dimensions r x c containing the number of successes in the binomial trials in each region, where:
  - r = number of rows = number of groups or sequences.
  - c = number of columns = number of regions.
- `binomN`: matrix of dimensions r x c containing the total number of trials in the binomial in each region, where:
  - r = number of rows = number of groups or sequences.
  - c = number of columns = number of regions.
- `binomP`: matrix of dimensions r x c containing the probability of success in one binomial trial in each region, where:
  - r = number of rows = number of groups or sequences.
  - c = number of columns = number of regions.
- `pdfs`: list of matrices containing PDFs with one item for each defined region (e.g. "CDR" and "FWR"). Matrices have dimensions r x c dementions, where:
  - r = number of rows = number of sequences or groups.
  - c = number of columns = length of the PDF (default 4001).
- `stats`: data.frame of BASELINE statistics, including: mean selection strength (mean Sigma), 95% confidence intervals, and p-values with positive signs for the presence of positive selection and/or p-values with negative signs for the presence of negative selection.

See Also

See `summarizeBaseline` for more information on `@stats`. 
Description

calcBaseline calculates the BASELINe posterior probability density functions (PDFs) for sequences in the given Change-O data.frame.

Usage

calcBaseline(
  db,
  sequenceColumn = "CLONAL_SEQUENCE",
  germlineColumn = "CLONAL_GERMLINE",
  testStatistic = c("local", "focused", "imbalanced"),
  regionDefinition = NULL,
  targetingModel = HH_S5F,
  mutationDefinition = NULL,
  calcStats = FALSE,
  nproc = 1
)

Arguments

db      data.frame containing sequence data and annotations.
sequenceColumn  character name of the column in db containing input sequences.
germlineColumn  character name of the column in db containing germline sequences.
testStatistic  character indicating the statistical framework used to test for selection. One of c("local","focused","imbalanced").
regionDefinition  RegionDefinition object defining the regions and boundaries of the Ig sequences.
targetingModel  TargetingModel object. Default is HH_S5F.
mutationDefinition  MutationDefinition object defining replacement and silent mutation criteria. If NULL then replacement and silent are determined by exact amino acid identity. Note, if the input data.frame already contains observed and expected mutation frequency columns then mutations will not be recalculated and this argument will be ignored.
calcStats  logical indicating whether or not to calculate the summary statistics data.frame stored in the stats slot of a Baseline object.
nproc  number of cores to distribute the operation over. If nproc=0 then the cluster has already been set and will not be reset.
Details

Calculates the BASELINe posterior probability density function (PDF) for sequences in the provided db.

Note: Individual sequences within clonal groups are not, strictly speaking, independent events and it is generally appropriate to only analyze selection pressures on an effective sequence for each clonal group. For this reason, it is strongly recommended that the input db contains one effective sequence per clone. Effective clonal sequences can be obtained by calling the collapseClones function.

If the db does not contain the required columns to calculate the PDFs (namely MU_COUNT & MU_EXPECTED) then the function will:

1. Calculate the numbers of observed mutations.
2. Calculate the expected frequencies of mutations and modify the provided db. The modified db will be included as part of the returned Baseline object.

The testStatistic indicates the statistical framework used to test for selection. E.g.

- local = CDR_R / (CDR_R + CDR_S).
- focused = CDR_R / (CDR_R + CDR_S + FWR_S).
- imbalanced = CDR_R + CDR_S / (CDR_R + CDR_S + FWR_S + FRW_R).

For focused the regionDefinition must only contain two regions. If more than two regions are defined the local test statistic will be used. For further information on the frame of these tests see Uduman et al. (2011).

Value

A Baseline object containing the modified db and BASELINe posterior probability density functions (PDF) for each of the sequences.

References


See Also

See Baseline for the return object. See groupBaseline and summarizeBaseline for further processing. See plotBaselineSummary and plotBaselineDensity for plotting results.
calcExpectedMutations

Examples

# Load and subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgG" & SAMPLE == "+7d")

# Collapse clones
db <- collapseClones(db, sequenceColumn="SEQUENCE_IMGT",
                germlineColumn="GERMLINE_IMGT_D_MASK",
                method="thresholdedFreq", minimumFrequency=0.6,
                includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

# Calculate BASELINe
baseline <- calcBaseline(db,
                sequenceColumn="CLONAL_SEQUENCE",
                germlineColumn="CLONAL_GERMLINE",
                testStatistic="focused",
                regionDefinition=IMGT_V,
                targetingModel=HH_S5F,
                nproc=1)

calcExpectedMutations  Calculate expected mutation frequencies of a sequence

Description

calcExpectedMutations calculates the expected mutation frequencies of a given sequence. This is primarily a helper function for expectedMutations.

Usage

calcExpectedMutations(
    germlineSeq,
    inputSeq = NULL,
    targetingModel = HH_S5F,
    regionDefinition = NULL,
    mutationDefinition = NULL
)

Arguments

  germlineSeq  germline (reference) sequence.
  inputSeq    input (observed) sequence. If this is not NULL, then germlineSeq will be processed to be the same same length as inputSeq and positions in germlineSeq corresponding to positions with Ns in inputSeq will also be assigned an N.
  targetingModel  TargetingModel object. Default is HH_S5F.
  regionDefinition  RegionDefinition object defining the regions and boundaries of the Ig sequences.
calcExpectedMutations

**mutationDefinition**

*MutationDefinition* object defining replacement and silent mutation criteria. If NULL then replacement and silent are determined by exact amino acid identity.

**Details**

calcExpectedMutations calculates the expected mutation frequencies of a given sequence and its germline.

Note, only the part of the sequences defined in *regionDefinition* are analyzed. For example, when using the default IMGT_V definition, mutations in positions beyond 312 will be ignored.

**Value**

A numeric vector of the expected frequencies of mutations in the regions in the *regionDefinition*. For example, when using the default IMGT_V definition, which defines positions for CDR and FWR, the following columns are calculated:

- **MU_EXPECTED_CDR_R**: number of replacement mutations in CDR1 and CDR2 of the V-segment.
- **MU_EXPECTED_CDR_S**: number of silent mutations in CDR1 and CDR2 of the V-segment.
- **MU_EXPECTED_FWR_R**: number of replacement mutations in FWR1, FWR2 and FWR3 of the V-segment.
- **MU_EXPECTED_FWR_S**: number of silent mutations in FWR1, FWR2 and FWR3 of the V-segment.

**See Also**

expectedMutations calls this function. To create a custom targetingModel see createTargetingModel. See calcObservedMutations for getting observed mutation counts.

**Examples**

```r
# Load example data
data(ExampleDb, package="alakazam")

# Use first entry in the exampled data for input and germline sequence
in_seq <- ExampleDb[["SEQUENCE_IMGT"]][1]
germ_seq <- ExampleDb[["GERMLINE_IMGT_D_MASK"]][1]

# Identify all mutations in the sequence
calcExpectedMutations(in_seq, germ_seq)

# Identify only mutations the V segment minus CDR3
calcExpectedMutations(in_seq, germ_seq, regionDefinition=IMGT_V)

# Define mutations based on hydropathy
calcExpectedMutations(in_seq, germ_seq, regionDefinition=IMGT_V,
  mutationDefinition=HYDROPATHY_MUTATIONS)
```
**calcObservedMutations**  
*Count the number of observed mutations in a sequence.*

**Description**

`calcObservedMutations` determines all the mutations in a given input sequence compared to its germline sequence.

**Usage**

```r
calcObservedMutations(
  inputSeq,  
germlineSeq,  
  regionDefinition = NULL,  
mutationDefinition = NULL,  
  ambiguousMode = c("eitherOr", "and"),  
  returnRaw = FALSE,  
  frequency = FALSE
)
```

**Arguments**

- **inputSeq** input sequence. IUPAC ambiguous characters for DNA are supported.
- **germlineSeq** germline sequence. IUPAC ambiguous characters for DNA are supported.
- **regionDefinition** `RegionDefinition` object defining the regions and boundaries of the Ig sequences. Note, only the part of sequences defined in `regionDefinition` are analyzed. If NULL, mutations are counted for entire sequence.
- **mutationDefinition** `MutationDefinition` object defining replacement and silent mutation criteria. If NULL then replacement and silent are determined by exact amino acid identity.
- **ambiguousMode** whether to consider ambiguous characters as "either or" or "and" when determining and counting the type(s) of mutations. Applicable only if `inputSeq` and/or `germlineSeq` contain(s) ambiguous characters. One of c("eitherOr","and"). Default is "eitherOr".
- **returnRaw** return the positions of point mutations and their corresponding mutation types, as opposed to counts of mutations across positions. Also returns the number of bases used as the denominator when calculating frequency. Default is FALSE.
- **frequency** logical indicating whether or not to calculate mutation frequencies. The denominator used is the number of bases that are not one of "N", ",", or ":" in either the input or the germline sequences. If set, this overwrites `returnRaw`. Default is FALSE.
Details

Each mutation is considered independently in the germline context. For illustration, consider the case where the germline is TGG and the observed is TAC. When determining the mutation type at position 2, which sees a change from G to A, we compare the codon TGG (germline) to TAG (mutation at position 2 independent of other mutations in the germline context). Similarly, when determining the mutation type at position 3, which sees a change from G to C, we compare the codon TGG (germline) to TGC (mutation at position 3 independent of other mutations in the germline context).

If specified, only the part of inputSeq defined in regionDefinition is analyzed. For example, when using the default IMGT_V definition, then mutations in positions beyond 312 will be ignored. Additionally, non-triplet overhang at the sequence end is ignored.

Only replacement (R) and silent (S) mutations are included in the results. Excluded are:

- Stop mutations
  E.g.: the case where TAGTGG is observed for the germline TGGTGG.
- Mutations occurring in codons where one or both of the observed and the germline involve(s) one or more of "N", ".", or ":".
  E.g.: the case where TTG is observed for the germline being any one of TNG, .TG, or ~TG. Similarly, the case where any one of TTN, TT., or TT~ is observed for the germline TTG.

In other words, a result that is NA or zero indicates absence of R and S mutations, not necessarily all types of mutations, such as the excluded ones mentioned above. NA is also returned if inputSeq or germlineSeq is shorter than 3 nucleotides.

Value

For returnRaw=FALSE, an array with the numbers of replacement (R) and silent (S) mutations.

For returnRaw=TRUE, a list containing

- $pos: A data frame whose columns (position, R, S, and region) indicate, respectively, the nucleotide position, the number of R mutations at that position, the number of S mutations at that position, and the region in which that nucleotide is in.
- $nonN: A vector indicating the number of bases in regions defined by regionDefinition (excluding non-triplet overhang, if any) that are not one of "N", ",", or ":" in either the inputSeq or germlineSeq.

For frequency=TRUE, regardless of returnRaw, an array with the frequencies of replacement (R) and silent (S) mutations.

Ambiguous characters

When there are ambiguous characters present, the user could choose how mutations involving ambiguous characters are counted through ambiguousMode. The two available modes are "eitherOr" and "and".

- With "eitherOr", ambiguous characters are each expanded but only 1 mutation is recorded. When determining the type of mutation, the priority for different types of mutations, in decreasing order, is as follows: no mutation, replacement mutation, silent mutation, and stop mutation.
When counting the number of non-N, non-dash, and non-dot positions, each position is counted only once, regardless of the presence of ambiguous characters.

As an example, consider the case where `germlineSeq` is "TST" and `inputSeq` is "THT". Expanding "H" at position 2 in `inputSeq` into "A", "C", and "T", as well as expanding "S" at position 2 in `germlineSeq` into "C" and "G", one gets:

- "TCT" (germline) to "TAT" (observed): replacement
- "TCT" (germline) to "TCT" (observed): no mutation
- "TCT" (germline) to "TTT" (observed): replacement
- "TGT" (germline) to "TAT" (observed): replacement
- "TGT" (germline) to "TCT" (observed): replacement
- "TGT" (germline) to "TTT" (observed): replacement

Because "no mutation" takes priority over replacement mutation, the final mutation count returned for this example is NA (recall that only R and S mutations are returned). The number of non-N, non-dash, and non-dot positions is 3.

• With "and", ambiguous characters are each expanded and mutation(s) from all expansions are recorded.

When counting the number of non-N, non-dash, and non-dot positions, if a position contains ambiguous character(s) in `inputSeq` and/or `germlineSeq`, the count at that position is taken to be the total number of combinations of germline and observed codons after expansion.

Using the same example from above, the final result returned for this example is that there are 5 R mutations at position 2. The number of non-N, non-dash, and non-dot positions is 8, since there are 6 combinations stemming from position 2 after expanding the germline codon ("TST") and the observed codon ("THT").

See Also

See `observedMutations` for counting the number of observed mutations in a data.frame.

Examples

```r
# Use an entry in the example data for input and germline sequence
data(ExampleDb, package="alakazam")
in_seq <- ExampleDb["SEQUENCE_IMGT"][[100]]
germ_seq <- ExampleDb["GERMLINE_IMGT_D_MASK"][[100]]

# Identify all mutations in the sequence
ex1_raw <- calcObservedMutations(in_seq, germ_seq, returnRaw=TRUE)

# Count all mutations in the sequence
ex1_count <- calcObservedMutations(in_seq, germ_seq, returnRaw=FALSE)
ex1_freq <- calcObservedMutations(in_seq, germ_seq, returnRaw=FALSE, frequency=TRUE)

# Compare this with ex1_count
table(ex1_raw$pos$region, ex1_raw$pos$R)[, "1"]/ex1_raw$nonN

# Compare this with ex1_freq
table(ex1_raw$pos$region, ex1_raw$pos$R)[, "1"]/ex1_raw$nonN

# Identify only mutations the V segment minus CDR3
```
calcTargetingDistance <- function(model, places = 2) {
  distance_matrix <- calcTargetingDistance(model, places = places)
  return(distance_matrix)
}

Description

calcTargetingDistance converts either the targeting rates in a TargetingModel model to a matrix of 5-mer to single-nucleotide mutation distances, or the substitution rates in a 1-mer substitution model to a symmetric distance matrix.
calcTargetingDistance

Arguments

- **model**: TargetingModel object with mutation likelihood information, or a 4x4 1-mer substitution matrix normalized by row with rownames and colnames consisting of "A", "T", "G", and "C".

- **places**: decimal places to round distances to.

Details

The targeting model is transformed into a distance matrix by:

1. Converting the likelihood of being mutated \( p = \text{mutability} \times \text{substitution} \) to distance \( d = -\log_{10}(p) \).
2. Dividing this distance by the mean of the distances.
3. Converting all infinite, no change (e.g., A->A), and NA distances to zero.

The 1-mer substitution matrix is transformed into a distance matrix by:

1. Symmetrize the 1-mer substitution matrix.
2. Converting the rates to distance \( d = -\log_{10}(p) \).
3. Dividing this distance by the mean of the distances.
4. Converting all infinite, no change (e.g., A -> A), and NA distances to zero.

Value

For input of TargetingModel, a matrix of distances for each 5-mer motif with rows names defining the center nucleotide and column names defining the 5-mer nucleotide sequence. For input of 1-mer substitution matrix, a 4x4 symmetric distance matrix.

See Also

See TargetingModel for this class of objects and createTargetingModel for building one.

Examples

```r
# Calculate targeting distance of HH_S5F
dist <- calcTargetingDistance(HH_S5F)

# Calculate targeting distance of HH_S1F
dist <- calcTargetingDistance(HH_S1F)
```
**calculateMutability**  
*Calculate total mutability*

**Description**

`calculateMutability` calculates the total (summed) mutability for a set of sequences based on a 5-mer nucleotide mutability model.

**Usage**

```r
calculateMutability(sequences, model = HH_S5F, progress = FALSE)
```

**Arguments**

- `sequences` character vector of sequences.
- `model` `TargetingModel` object with mutation likelihood information.
- `progress` if TRUE print a progress bar.

**Value**

Numeric vector with a total mutability score for each sequence.

**Examples**

```r
# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Calculate mutability of germline sequences using \link{HH_S5F} model
mutability <- calculateMutability(sequences=db$GERMLINE_IMGT_D_MASK, model=HH_S5F)
```

---

**collapseClones**  
*Constructs effective clonal sequences for all clones*

**Description**

`collapseClones` creates effective input and germline sequences for each clonal group and appends columns containing the consensus sequences to the input data.frame.
collapseClones

db, cloneColumn = "CLONE", sequenceColumn = "SEQUENCE IMGT", germlineColumn = "GERMLINE IMGT_D_MASK", muFreqColumn = NULL,
regionDefinition = NULL, method = c("mostCommon", "thresholdedFreq", "catchAll", "mostMutated", "leastMutated"),
minimumFrequency = NULL, includeAmbiguous = FALSE,
breakTiesStochastic = FALSE, breakTiesByColumns = NULL,
expandedDb = FALSE, nproc = 1
)

Arguments

db data.frame containing sequence data. Required.
clonelColumn character name of the column containing clonal identifiers. Required.
sequenceColumn character name of the column containing input sequences. Required. The length of each input sequence should match that of its corresponding germline sequence.
germlineColumn character name of the column containing germline sequences. Required. The length of each germline sequence should match that of its corresponding input sequence.
muFreqColumn character name of the column containing mutation frequency. Optional. Applicable to the "mostMutated" and "leastMutated" methods. If not supplied, mutation frequency is computed by calling observedMutations. Default is NULL. See Cautions for note on usage.
regionDefinition RegionDefinition object defining the regions and boundaries of the Ig sequences. Optional. Default is NULL.
method method for calculating input consensus sequence. Required. One of "thresholdedFreq", "mostCommon", "catchAll", "mostMutated", or "leastMutated". See "Methods" for details.
minimumFrequency frequency threshold for calculating input consensus sequence. Applicable to and required for the "thresholdedFreq" method. A canonical choice is 0.6. Default is NULL.
includeAmbiguous whether to use ambiguous characters to represent positions at which there are multiple characters with frequencies that are at least minimumFrequency or that are maximal (i.e. ties). Applicable to and required for the "thresholdedFreq" and "mostCommon" methods. Default is FALSE. See "Choosing ambiguous characters" for rules on choosing ambiguous characters.
collapseClones

breakTiesStochastic
In case of ties, whether to randomly pick a sequence from sequences that fulfill the criteria as consensus. Applicable to and required for all methods except for "catchAll". Default is FALSE. See "Methods" for details.

breakTiesByColumns
A list of the form list(c(col_1,col_2,...),c(fun_1,fun_2,...)), where col_i is a character name of a column in db, and fun_i is a function to be applied on that column. Currently, only max and min are supported. Note that the two c()’s in list() are essential (i.e. if there is only 1 column, the list should be of the form list(c(col_1),c(func_1)). Applicable to and optional for the "mostMutated" and "leastMutated" methods. If supplied, fun_i’s are applied on col_i’s to help break ties. Default is NULL. See "Methods" for details.

expandedDb logical indicating whether or not to return the expanded db, containing all the sequences (as opposed to returning just one sequence per clone).

nproc Number of cores to distribute the operation over. If the cluster has already been set earlier, then pass the cluster. This will ensure that it is not reset.

Value
A modified db with the following additional columns:

- CLONAL_SEQUENCE: effective sequence for the clone.
- CLONAL_GERMLINE: germline sequence for the clone.
- CLONAL_SEQUENCE_MUFREQ: mutation frequency of CLONAL_SEQUENCE; only added for the "mostMutated" and "leastMutated" methods.

CLONAL_SEQUENCE is generated with the method of choice indicated by method, and CLONAL_GERMLINE is generated with the "mostCommon" method, along with, where applicable, user-defined parameters such as minimumFrequency, includeAmbiguous, breakTiesStochastic, and breakTiesByColumns.

Consensus lengths
For each clone, CLONAL_SEQUENCE and CLONAL_GERMLINE have the same length.

- For the "thresholdedFreq", "mostCommon", and "catchAll" methods:
  The length of the consensus sequences is determined by the longest possible consensus sequence (based on inputSeq and germlineSeq) and regionDefinition@seqLength (if supplied), whichever is shorter.
  Given a set of sequences of potentially varying lengths, the longest possible length of their consensus sequence is taken to be the longest length along which there is information contained at every nucleotide position across majority of the sequences. Majority is defined to be greater than floor(n/2), where n is the number of sequences. If the longest possible consensus length is 0, there will be a warning and an empty string ("") will be returned.
  If a length limit is defined by supplying a regionDefinition via regionDefinition@seqLength, the consensus length will be further restricted to the shorter of the longest possible length and regionDefinition@seqLength.
• For the "mostMutated" and "leastMutated" methods:
The length of the consensus sequences depends on that of the most/least mutated input sequence, and, if supplied, the length limit defined by regionDefinition@seqLength, whichever is shorter. If the germline consensus computed using the "mostCommon" method is longer than the most/least mutated input sequence, the germline consensus is trimmed to be of the same length as the input consensus.

Methods

The descriptions below use "sequences" as a generalization of input sequences and germline sequences.

• method="thresholdedFreq"
  A threshold must be supplied to the argument minimumFrequency. At each position along the length of the consensus sequence, the frequency of each nucleotide/character across sequences is tabulated. The nucleotide/character whose frequency is at least (i.e. $\geq$) minimumFrequency becomes the consensus; if there is none, the consensus nucleotide will be "N". When there are ties (frequencies of multiple nucleotides/characters are at least minimumFrequency), this method can be deterministic or stochastic, depending on additional parameters.
  – With includeAmbiguous=TRUE, ties are resolved deterministically by representing ties using ambiguous characters. See "Choosing ambiguous characters" for how ambiguous characters are chosen.
  – With breakTiesStochastic=TRUE, ties are resolved stochastically by randomly picking a character amongst the ties.
  – When both TRUE, includeAmbiguous takes precedence over breakTiesStochastic.
  – When both FALSE, the first character from the ties is taken to be the consensus following the order of "A", "T", "G", "C", "N", ".", and "-".

Below are some examples looking at a single position based on 5 sequences with minimumFrequency=0.6, includeAmbiguous=FALSE, and breakTiesStochastic=FALSE:
  – If the sequences have "A", "A", "A", "T", "C", the consensus will be "A", because "A" has frequency 0.6, which is at least minimumFrequency.
  – If the sequences have "A", "A", "T", "T", "C", the consensus will be "N", because none of "A", "T", or "C" has frequency that is at least minimumFrequency.

• method="mostCommon"
The most frequent nucleotide/character across sequences at each position along the length of the consensus sequence makes up the consensus.
  When there are ties (multiple nucleotides/characters with equally maximal frequencies), this method can be deterministic or stochastic, depending on additional parameters. The same rules for breaking ties for method="thresholdedFreq" apply.

Below are some examples looking at a single position based on 5 sequences with includeAmbiguous=FALSE, and breakTiesStochastic=FALSE:
  – If the sequences have "A", "A", "T", "A", "C", the consensus will be "A".
  – If the sequences have "T", "T", "C", "C", "G", the consensus will be "T", because "T" is before "C" in the order of "A", "T", "G", "C", "N", ".", and "-".
• method="catchAll"
This method returns a consensus sequence capturing most of the information contained in the sequences. Ambiguous characters are used where applicable. See "Choosing ambiguous characters" for how ambiguous characters are chosen. This method is deterministic and does not involve breaking ties.

Below are some examples for method="catchAll" looking at a single position based on 5 sequences:

– If the sequences have "N", "N", "N", "N", "N", the consensus will be "N".
– If the sequences have "N", "A", "A", "A", "A", the consensus will be "A".
– If the sequences have "N", "A", "G", "A", "A", the consensus will be "R".
– If the sequences have ",", ",", ",", ",", ",", the consensus will be ",".
– If the sequences have ",", ",", ",", ",", "-", the consensus will be ",-".
– If the sequences have ",", ",", ",", ",", ",", "-", the consensus will be "-".

• method="mostMutated" and method="leastMutated"
These methods return the most/least mutated sequence as the consensus sequence.
When there are ties (multiple sequences have the maximal/minimal mutation frequency), this method can be deterministic or stochastic, depending on additional parameters.

– With breakTiesStochastic=TRUE, ties are resolved stochastically by randomly picking a sequence out of sequences with the maximal/minimal mutation frequency.
– When breakTiesByColumns is supplied, ties are resolved deterministically. Column by column, a function is applied on the column and sequences with column value matching the functional value are retained, until ties are resolved or columns run out. In the latter case, the first remaining sequence is taken as the consensus.
– When breakTiesStochastic=TRUE and breakTiesByColumns is supplied, breakTiesStochastic takes precedence over breakTiesByColumns.
– When breakTiesStochastic=FALSE and breakTiesByColumns is not supplied (i.e. NULL), the sequence that appears first amongst the ties is taken as the consensus.

Choosing ambiguous characters

Ambiguous characters may be present in the returned consensuses when using the "catchAll" method and when using the "thresholdedFreq" or "mostCommon" methods with includeAmbiguous=TRUE.

The rules on choosing ambiguous characters are as follows:

• If a position contains only "N" across sequences, the consensus at that position is "N".
• If a position contains one or more of "A", "T", "G", or "C", the consensus will be an IUPAC character representing all of the characters present, regardless of whether "N", ",", or "," is present.
• If a position contains only ",-" and "," across sequences, the consensus at that position is taken to be ",-".
• If a position contains only one of "-" or "," across sequences, the consensus at that position is taken to be the character present.
Cautions

• Note that this function does not perform multiple sequence alignment. As a prerequisite, it is assumed that the sequences in sequenceColumn and germlineColumn have been aligned somehow. In the case of immunoglobulin repertoire analysis, this usually means that the sequences are IMGT-gapped.

• When using the "mostMutated" and "leastMutated" methods, if you supply both muFreqColumn and regionDefinition, it is your responsibility to ensure that the mutation frequency in muFreqColumn was calculated with sequence lengths restricted to the same regionDefinition you are supplying. Otherwise, the "most/least mutated" sequence you obtain might not be the most/least mutated given the regionDefinition supplied, because your mutation frequency was based on a regionDefinition different from the one supplied.

• If you intend to run collapseClones before building a 5-mer targeting model, you **must** choose parameters such that your collapsed clonal consensuses do not include ambiguous characters. This is because the targeting model functions do NOT support ambiguous characters in their inputs.

See Also

See [IMGT_SCHEMES](#) for a set of predefined RegionDefinition objects.

Examples

```r
# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE %in% c("IgA", "IgG") & SAMPLE == "+7d" & CLONE %in% c("3100", "3141", "3184"))

# thresholdedFreq method, resolving ties deterministically without using ambiguous characters
clones <- collapseClones(db, method="thresholdedFreq", minimumFrequency=0.6, includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

# mostCommon method, resolving ties deterministically using ambiguous characters
clones <- collapseClones(db, method="mostCommon", includeAmbiguous=TRUE, breakTiesStochastic=FALSE)

# Make a copy of db that has a mutation frequency column
db2 <- observedMutations(db, frequency=TRUE, combine=TRUE)

# mostMutated method, resolving ties stochastically
clones <- collapseClones(db2, method="mostMutated", muFreqColumn="MU_FREQ", breakTiesStochastic=TRUE, breakTiesByColumns=NULL)

# mostMutated method, resolving ties deterministically using additional columns
clones <- collapseClones(db2, method="mostMutated", muFreqColumn="MU_FREQ", breakTiesStochastic=FALSE, breakTiesByColumns=list(c("DUPCOUNT"), c(max)))

# Build consensus for V segment only
# Capture all nucleotide variations using ambiguous characters
clones <- collapseClones(db, method="catchAll", regionDefinition=IMGT_V)
```
# Return the same number of rows as the input
clones <- collapseClones(db, method="mostCommon", expandedDb=TRUE)

## consensusSequence

### Construct a consensus sequence

#### Description

Construct a consensus sequence

#### Usage

```r
consensusSequence(
  sequences,
  db = NULL,
  method = c("mostCommon", "thresholdedFreq", "catchAll", "mostMutated", "leastMutated"),
  minFreq = NULL,
  muFreqColumn = NULL,
  lenLimit = NULL,
  includeAmbiguous = FALSE,
  breakTiesStochastic = FALSE,
  breakTiesByColumns = NULL
)
```

#### Arguments

- **sequences**: character vector of sequences.
- **db**: data.frame containing sequence data for a single clone. Applicable to and required for the "mostMutated" and "leastMutated" methods. Default is NULL.
- **method**: method to calculate consensus sequence. One of "thresholdedFreq", "mostCommon", "catchAll", "mostMutated", or "leastMutated". See "Methods" under collapseClones for details.
- **minFreq**: frequency threshold for calculating input consensus sequence. Applicable to and required for the "thresholdedFreq" method. A canonical choice is 0.6. Default is NULL.
- **muFreqColumn**: character name of the column in db containing mutation frequency. Applicable to and required for the "mostMutated" and "leastMutated" methods. Default is NULL.
- **lenLimit**: limit on consensus length. if NULL then no length limit is set.
- **includeAmbiguous**: whether to use ambiguous characters to represent positions at which there are multiple characters with frequencies that are at least minimumFrequency or that are maximal (i.e. ties). Applicable to and required for the "thresholdedFreq" and "mostCommon" methods. Default is FALSE. See "Choosing ambiguous characters" under collapseClones for rules on choosing ambiguous characters.
**breakTiesStochastic**
In case of ties, whether to randomly pick a sequence from sequences that fulfill the criteria as consensus. Applicable to and required for all methods except for "catchAll". Default is FALSE. See "Methods" under collapseClones for details.

**breakTiesByColumns**
A list of the form list(c(col_1, col_2,...), c(fun_1, fun_2,...)), where col_i is a character name of a column in db, and fun_i is a function to be applied on that column. Currently, only max and min are supported. Note that the two c()'s in list() are essential (i.e. if there is only 1 column, the list should be of the form list(c(col_1), c(func_1)). Applicable to and optional for the "mostMutated" and "leastMutated" methods. If supplied, fun_i's are applied on col_i's to help break ties. Default is NULL. See "Methods" under collapseClones for details.

**Details**
See collapseClones for detailed documentation on methods and additional parameters.

**Value**
A list containing cons, which is a character string that is the consensus sequence for sequences; and muFreq, which is the maximal/minimal mutation frequency of the consensus sequence for the "mostMutated" and "leastMutated" methods, or NULL for all other methods.

**Examples**
# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE %in% c("IgA", "IgG") & SAMPLE == "+7d")
close <- subset(db, CLONE == "3192")

# First compute mutation frequency for most/leastMutated methods
close <- observedMutations(clone, frequency=TRUE, combine=TRUE)

# Manually create a tie
close <- rbind(close, clone[which.max(close$MU_FREQ), ])

# ThresholdedFreq method,
# Resolve ties deterministically without using ambiguous characters
cons1 <- consensusSequence(close$SEQUENCE_IMGT,
method="thresholdedFreq", minFreq=0.3,
includeAmbiguous=FALSE,
brakeTiesStochastic=FALSE)

cons1$cons
createBaseline

Creates a Baseline object

Description

createBaseline creates and initialize a Baseline object.

Usage

createBaseline(
  description = "",
  db = data.frame(),
  regionDefinition = createRegionDefinition(),
  testStatistic = "",
  regions = NULL,
  numbOfSeqs = matrix(),
  binomK = matrix(),
  binomN = matrix(),
  binomP = matrix(),
  pdfs = list(),
  stats = data.frame()
)

Arguments

description character providing general information regarding the sequences, selection analysis and/or object.

db data.frame containing annotation information about the sequences and selection results.

regionDefinition RegionDefinition object defining the regions and boundaries of the Ig sequences.

testStatistic character indicating the statistical framework used to test for selection. For example, "local" or "focused" or "imbalanced".

regions character vector defining the regions the BASELINE analysis was carried out on. For "CDR" and "FWR" or "CDR1", "CDR2", "CDR3", etc. If NULL then regions will be determined automatically from regionDefinition.

numbOfSeqs matrix of dimensions r x c containing the number of sequences or PDFs in each region, where:
r = number of rows = number of groups or sequences.
c = number of columns = number of regions.

binomK matrix of dimensions r x c containing the number of successes in the binomial trials in each region, where:
r = number of rows = number of groups or sequences.
c = number of columns = number of regions.
createBaseline

- **binomN**: matrix of dimensions $r \times c$ containing the total number of trials in the binomial in each region, where:
  - $r$ = number of rows = number of groups or sequences.
  - $c$ = number of columns = number of regions.

- **binomP**: matrix of dimensions $r \times c$ containing the probability of success in one binomial trial in each region, where:
  - $r$ = number of rows = number of groups or sequences.
  - $c$ = number of columns = number of regions.

- **pdfs**: list of matrices containing PDFs with one item for each defined region (e.g. "CDR" and "FWR"). Matrices have dimensions $r \times c$ dementions, where:
  - $r$ = number of rows = number of sequences or groups.
  - $c$ = number of columns = length of the PDF (default 4001).

- **stats**: data.frame of BASELINE statistics, including: mean selection strength (mean Sigma), 95% confidence intervals, and p-values with positive signs for the presence of positive selection and/or p-values with negative signs for the presence of negative selection.

**Details**

Create and initialize a Baseline object.

The `testStatistic` indicates the statistical framework used to test for selection. For example,

- local = $\frac{CDR_R}{CDR_R + CDR_S}$.
- focused = $\frac{CDR_R}{CDR_R + CDR_S + FWR_S}$.
- imbalance = $\frac{CDR_R + CDR_s}{CDR_R + CDR_S + FWR_S + FWR_R}$

For focused the `regionDefinition` must only contain two regions. If more than two regions are defined, then the `local` test statistic will be used. For further information on the frame of these tests see Uduman et al. (2011).

**Value**

A Baseline object.

**References**


**See Also**

See Baseline for the return object.
createMutabilityMatrix

*Builds a mutability model*

**Description**

`createMutabilityMatrix` builds a 5-mer nucleotide mutability model by counting the number of mutations occurring in the center position for all 5-mer motifs.

**Usage**

```r
createMutabilityMatrix(
  db,
  substitutionModel,
  model = c("S", "RS"),
  sequenceColumn = "SEQUENCE_IMGT",
  germlineColumn = "GERMLINE_IMGT_D_MASK",
  vCallColumn = "V_CALL",
  multipleMutation = c("independent", "ignore"),
  minNumSeqMutations = 500,
  numSeqMutationsOnly = FALSE,
  returnSource = FALSE
)
```

**Arguments**

- `db`: data.frame containing sequence data.
- `substitutionModel`: matrix of 5-mer substitution rates built by `createSubstitutionMatrix`. Note, this model will only impact mutability scores when `model="S"` (using only silent mutations).
- `model`: type of model to create. The default model, "S", builds a model by counting only silent mutations. `model="S"` should be used for data that includes functional sequences. Setting `model="RS"` creates a model by counting both replacement and silent mutations and may be used on fully non-functional sequence data sets.
- `sequenceColumn`: name of the column containing IMGT-gapped sample sequences.
- `germlineColumn`: name of the column containing IMGT-gapped germline sequences.
- `vCallColumn`: name of the column containing the V-segment allele call.
multipleMutation

string specifying how to handle multiple mutations occurring within the same 5-mer. If "independent" then multiple mutations within the same 5-mer are counted independently. If "ignore" then 5-mers with multiple mutations are excluded from the total mutation tally.

minNumSeqMutations

minimum number of mutations in sequences containing each 5-mer to compute the mutability rates. If the number is smaller than this threshold, the mutability for the 5-mer will be inferred. Default is 500. Not required if numSeqMutationsOnly=TRUE.

numSeqMutationsOnly

when TRUE, return only a vector counting the number of observed mutations in sequences containing each 5-mer. This option can be used for parameter tuning for minNumSeqMutations during preliminary analysis using minNumSeqMutationsTune. Default is FALSE.

returnSource

return the sources of 5-mer mutabilities (measured vs. inferred). Default is FALSE.

Details

Caution: The targeting model functions do NOT support ambiguous characters in their inputs. You MUST make sure that your input and germline sequences do NOT contain ambiguous characters (especially if they are clonal consensuses returned from collapseClones).

Value

When numSeqMutationsOnly is FALSE, a named numeric vector of 1024 normalized mutability rates for each 5-mer motif with names defining the 5-mer nucleotide sequence.

When numSeqMutationsOnly is TRUE, a named numeric vector of length 1024 counting the number of observed mutations in sequences containing each 5-mer.

References


See Also

extendMutabilityMatrix, createSubstitutionMatrix, createTargetingMatrix, createTargetingModel, minNumSeqMutationsTune

Examples

# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")
# Create model using only silent mutations
sub_model <- createSubstitutionMatrix(db, model="S")
mut_model <- createMutabilityMatrix(db, sub_model, model="S",
minNumSeqMutations=200,
numSeqMutationsOnly=FALSE)

# Count the number of mutations in sequences containing each 5-mer
mut_count <- createMutabilityMatrix(db, sub_model, model="S",
numSeqMutationsOnly=TRUE)

createMutationDefinition

*Creates a MutationDefinition*

**Description**
createMutationDefinition creates a MutationDefinition.

**Usage**
createMutationDefinition(name, classes, description = "", citation = "")

**Arguments**

- **name**
  name of the mutation definition.

- **classes**
  named character vectors with single-letter amino acid codes as names and amino acid classes as values, with NA assigned to set of characters c("X","*","-","."). Replacement (R) is be defined as a change in amino acid class and silent (S) as no change in class.

- **description**
  description of the mutation definition and its source data.

- **citation**
  publication source.

**Value**
A MutationDefinition object.

**See Also**
See MutationDefinition for the return object.
Examples

# Define hydropathy classes
library(alakazam)
hydropathy <- list(hydrophobic=c("A", "I", "L", "M", "F", "W", "V"),
                  hydrophilic=c("R", "N", "D", "C", "Q", "E", "K"),
                  neutral=c("G", "H", "P", "S", "T", "Y"))
chars <- unlist(hydropathy, use.names=FALSE)
classes <- setNames(translateStrings(chars, hydropathy), chars)

# Create hydropathy mutation definition
md <- createMutationDefinition("Hydropathy", classes)

createRegionDefinition

Creates a RegionDefinition

Description

createRegionDefinition creates a RegionDefinition.

Usage

createRegionDefinition(
  name = "",
  boundaries = factor(),
  description = "",
  citation = ""
)

Arguments

name name of the region definition.
boundaries factor defining the region boundaries of the sequence. The levels and values of boundaries determine the number of regions (e.g. CDR and FWR).
description description of the region definition and its source data.
citation publication source.

Value

A RegionDefinition object.

See Also

See RegionDefinition for the return object.
createSubstitutionMatrix

Builds a substitution model

Description

createSubstitutionMatrix builds a 5-mer nucleotide substitution model by counting the number of substitution mutations occurring in the center position for all 5-mer motifs.

Usage

createSubstitutionMatrix(
  db,
  model = c("S", "RS"),
  sequenceColumn = "SEQUENCE_IMGT",
  germlineColumn = "GERMLINE_IMGT_D_MASK",
  vCallColumn = "V_CALL",
  multipleMutation = c("independent", "ignore"),
  returnModel = c("5mer", "1mer", "1mer_raw"),
  minNumMutations = 50,
  numMutationsOnly = FALSE
)

Arguments

db data.frame containing sequence data.
model type of model to create. The default model, "S", builds a model by counting only silent mutations. model="S" should be used for data that includes functional sequences. Setting model="RS" creates a model by counting both replacement and silent mutations and may be used on fully non-functional sequence data sets.
sequenceColumn name of the column containing IMGT-gapped sample sequences.
germlineColumn name of the column containing IMGT-gapped germline sequences.
vCallColumn name of the column containing the V-segment allele call.
multipleMutation string specifying how to handle multiple mutations occurring within the same 5-mer. If "independent" then multiple mutations within the same 5-mer are counted independently. If "ignore" then 5-mers with multiple mutations are excluded from the total mutation tally.
createSubstitutionMatrix

returnModel  
string specifying what type of model to return; one of c("5mer", "1mer", "1mer_raw").  
If "5mer" (the default) then a 5-mer nucleotide context model is returned. If "1mer" or "1mer_raw" then a single nucleotide substitution matrix (no context) is returned; where "1mer_raw" is the unnormalized version of the "1mer" model. Note, neither 1-mer model may be used as input to createMutabilityMatrix.

minNumMutations  
minimum number of mutations required to compute the 5-mer substitution rates.  
If the number of mutations for a 5-mer is below this threshold, its substitution rates will be estimated from neighboring 5-mers. Default is 50. Not required if numMutationsOnly=TRUE.

numMutationsOnly  
when TRUE, return counting information on the number of mutations for each 5-mer, instead of building a substitution matrix. This option can be used for parameter tuning for minNumMutations during preliminary analysis. Default is FALSE. Only applies when returnModel is set to "5mer". The data.frame returned when this argument is TRUE can serve as the input for minNumMutationsTune.

Details

Caution: The targeting model functions do NOT support ambiguous characters in their inputs. You MUST make sure that your input and germline sequences do NOT contain ambiguous characters (especially if they are clonal consensuses returned from collapseClones).

Value

For returnModel = "5mer":

When numMutationsOnly is FALSE, a 4x1024 matrix of column normalized substitution rates for each 5-mer motif with row names defining the center nucleotide, one of c("A", "C", "G", "T"), and column names defining the 5-mer nucleotide sequence.

When numMutationsOnly is TRUE, a 1024x4 data frame with each row providing information on counting the number of mutations for a 5-mer. Columns are named fivemer.total, fivemer.every, inner3.total, and inner3.every, corresponding to, respectively, the total number of mutations when counted as a 5-mer, whether there is mutation to every other base when counted as a 5-mer, the total number of mutations when counted as an inner 3-mer, and whether there is mutation to every other base when counted as an inner 3-mer.

For returnModel = "1mer" or "1mer_raw": a 4x4 normalized or un-normalized 1-mer substitution matrix respectively.

References

createTargetingMatrix

Calculates a targeting rate matrix

description

createTargetingMatrix calculates the targeting model matrix as the combined probability of mutability and substitution.

usage

createTargetingMatrix(substitutionModel, mutabilityModel)

arguments

substitutionModel
matrix of 5-mers substitution rates built by createSubstitutionMatrix or extendSubstitutionMatrix.

mutabilityModel
vector of 5-mers mutability rates built by createMutabilityMatrix or extendMutabilityMatrix.

details

Targeting rates are calculated by multiplying the normalized mutability rate by the normalized substitution rates for each individual 5-mer.
Value

A matrix with the same dimensions as the input substitutionModel containing normalized targeting probabilities for each 5-mer motif with row names defining the center nucleotide and column names defining the 5-mer nucleotide sequence.

References


See Also

createSubstitutionMatrix, extendSubstitutionMatrix, createMutabilityMatrix, extendMutabilityMatrix, createTargetingModel

Examples

# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Create 4x1024 models using only silent mutations
sub_model <- createSubstitutionMatrix(db, model="S")
mut_model <- createMutabilityMatrix(db, sub_model, model="S")

# Extend substitution and mutability to including Ns (5x3125 model)
sub_model <- extendSubstitutionMatrix(sub_model)
mut_model <- extendMutabilityMatrix(mut_model)

# Create targeting model from substitution and mutability
tar_model <- createTargetingMatrix(sub_model, mut_model)
sequenceColumn = "SEQUENCE_IMGT",
germlineColumn = "GERMLINE_IMGT_D_MASK",
vCallColumn = "V_CALL",
multipleMutation = c("independent", "ignore"),
minNumMutations = 50,
minNumSeqMutations = 500,
modelName = "",
modelDescription = "",
modelSpecies = "",
modelCitation = "",
modelDate = NULL
)

Arguments

**db**
data.frame containing sequence data.

**model**
type of model to create. The default model, "S", builds a model by counting only silent mutations. model="S" should be used for data that includes functional sequences. Setting model="RS" creates a model by counting both replacement and silent mutations and may be used on fully non-functional sequence data sets.

**sequenceColumn**
name of the column containing IMGT-gapped sample sequences.

**germlineColumn**
name of the column containing IMGT-gapped germline sequences.

**vCallColumn**
name of the column containing the V-segment allele calls.

**multipleMutation**
string specifying how to handle multiple mutations occurring within the same 5-mer. If "independent" then multiple mutations within the same 5-mer are counted independently. If "ignore" then 5-mers with multiple mutations are excluded from the total mutation tally.

**minNumMutations**
minimum number of mutations required to compute the 5-mer substitution rates. If the number of mutations for a 5-mer is below this threshold, its substitution rates will be estimated from neighboring 5-mers. Default is 50.

**minNumSeqMutations**
minimum number of mutations in sequences containing each 5-mer to compute the mutability rates. If the number is smaller than this threshold, the mutability for the 5-mer will be inferred. Default is 500.

**modelName**
name of the model.

**modelDescription**
description of the model and its source data.

**modelSpecies**
genus and species of the source sequencing data.

**modelCitation**
publication source.

**modelDate**
date the model was built. If NULL the current date will be used.

Details

**Caution**: The targeting model functions do NOT support ambiguous characters in their inputs. You MUST make sure that your input and germline sequences do NOT contain ambiguous characters (especially if they are clonal consensuses returned from collapseClones).
DensityThreshold-class

Value

A TargetingModel object.

References


See Also

See TargetingModel for the return object. See plotMutability plotting a mutability model. See createSubstitutionMatrix, extendSubstitutionMatrix, createMutabilityMatrix, extendMutabilityMatrix and createTargetingMatrix for component steps in building a model.

Examples

# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Create model using only silent mutations and ignore multiple mutations
model <- createTargetingModel(db, model="S", multipleMutation="ignore")

DensityThreshold-class

Output of the dens method of findThreshold

Description

DensityThreshold contains output from the dens method findThreshold.

Usage

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

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

Arguments

x DensityThreshold object
y ignored.
... arguments to pass to plotDensityThreshold.
Slots

- x: input distance vector with NA or infinite values removed.
- bandwidth: bandwidth value fit during density estimation.
- xdens: x-axis (distance value) vector for smoothed density estimate.
- ydens: y-axis (density) vector for smoothed density estimate.
- threshold: distance threshold that separates two modes of the input distribution.

See Also

findThreshold

distToNearest

Distance to nearest neighbor

Description

Get non-zero distance of every heavy chain (IGH) sequence (as defined by sequenceColumn) to its nearest sequence in a partition of heavy chains sharing the same V gene, J gene, and junction length (VJL), or in a partition of single cells with heavy chains sharing the same heavy chain VJL combination, or of single cells with heavy and light chains sharing the same heavy chain VJL and light chain VJL combinations.

Usage

```r
distToNearest(
  db,
  sequenceColumn = "JUNCTION",
  vCallColumn = "V_CALL",
  jCallColumn = "J_CALL",
  model = c("ham", "aa", "hh_s1f", "hh_s5f", "mk_rs1nf", "mk_rs5nf", "m1n_compat", "hs1f_compat"),
  normalize = c("len", "none"),
  symmetry = c("avg", "min"),
  first = TRUE,
  VJthenLen = TRUE,
  nproc = 1,
  fields = NULL,
  cross = NULL,
  mst = FALSE,
  subsample = NULL,
  progress = FALSE,
  cellIdColumn = NULL,
  locusColumn = NULL,
  groupUsingOnlyIGH = TRUE,
  keepVJLgroup = TRUE
)
```
distToNearest

**Arguments**

- **db**
  - data.frame containing sequence data.
- **sequenceColumn**
  - name of the column containing the junction for grouping and for calculating nearest neighbor distances. Note that while both heavy and light chain junctions may be used for VJL grouping, only the heavy chain junction is used to calculate distances.
- **vCallColumn**
  - name of the column containing the V-segment allele calls.
- **jCallColumn**
  - name of the column containing the J-segment allele calls.
- **model**
  - underlying SHM model, which must be one of \texttt{c("ham","aa","hh\_s1f","hh\_s5f","mk\_rs1nf","hs1f")}.
  - See Details for further information.
- **normalize**
  - method of normalization. The default is \texttt{"len"}, which divides the distance by the length of the sequence group. If \texttt{"none"} then no normalization if performed.
- **symmetry**
  - if model is \texttt{hs5f}, distance between seq1 and seq2 is either the average (avg) of seq1->seq2 and seq2->seq1 or the minimum (min).
- **first**
  - if \texttt{TRUE} only the first call of the gene assignments is used. if \texttt{FALSE} the union of ambiguous gene assignments is used to group all sequences with any overlapping gene calls.
- **VJthenLen**
  - a Boolean value specifying whether to perform partitioning as a 2-stage process. If \texttt{TRUE}, partitions are made first based on V and J annotations, and then further split based on junction lengths corresponding to \texttt{sequenceColumn}. If \texttt{FALSE}, perform partition as a 1-stage process during which V annotation, J annotation, and junction length are used to create partitions simultaneously. Defaults to \texttt{TRUE}.
- **nproc**
  - number of cores to distribute the function over.
- **fields**
  - additional fields to use for grouping.
- **cross**
  - character vector of column names to use for grouping to calculate distances across groups. Meaning the columns that define self versus others.
- **mst**
  - if \texttt{TRUE}, return comma-separated branch lengths from minimum spanning tree.
- **subsample**
  - number of sequences to subsample for speeding up pairwise-distance-matrix calculation. Subsampling is performed without replacement in each VJL group of heavy chain sequences. If \texttt{subsample} is larger than the unique number of heavy chain sequences in each VJL group, then the subsampling process is ignored for that group. For each heavy chain sequence in \texttt{db}, the reported DIST_NEAREST is the distance to the closest heavy chain sequence in the subsampled set for the VJL group. If \texttt{NULL} no subsampling is performed.
- **progress**
  - if \texttt{TRUE} print a progress bar.
- **cellIdColumn**
  - name of the column containing cell IDs. Only applicable and required for single-cell mode.
- **locusColumn**
  - name of the column containing locus information. Only applicable and required for single-cell mode.
- **groupUsingOnlyIGH**
  - use only heavy chain (IGH) sequences for VJL grouping, disregarding light chains. Only applicable and required for single-cell mode. Default is \texttt{TRUE}. Also see \texttt{groupGenes}.

Also see \texttt{groupGenes}.
keepVJLgroup  a Boolean value specifying whether to keep in the output the the column column indicating grouping based on VJL combinations. Only applicable for 1-stage partitioning (i.e. VJthenLen=FALSE). Also see groupGenes.

Details

To invoke single-cell mode, both cellIdColumn and locusColumn must be supplied. Otherwise, the function will run under non-single-cell mode.

Under single-cell mode, only heavy chain sequences will be used for calculating nearest neighbor distances. Under non-single-cell mode, all input sequences will be used for calculating nearest neighbor distances, regardless of the values in the locusColumn field (if present).

For single-cell mode, the input format is the same as that for groupGenes. Namely, each row represents a sequence/chain. Sequences/chains from the same cell are linked by a cell ID in the cellIdColumn field. Under this mode, there is a choice of whether grouping should be done using only heavy chain (IGH) sequences only, or using both heavy chain (IGH) and light chain (IGK, IGL) sequences. This is governed by groupUsingOnlyIGH.

If used, values in the locusColumn column must be one of "IGH", "IGK", and "IGL".

Note that for distToNearest, a cell with multiple heavy chains is not allowed.

The distance to nearest (heavy chain) neighbor can be used to estimate a threshold for assigning Ig sequences to clonal groups. A histogram of the resulting vector is often bimodal, with the ideal threshold being a value that separates the two modes.

The following distance measures are accepted by the model parameter.

- "ham": Single nucleotide Hamming distance matrix from getDNAMatrix with gaps assigned zero distance.
- "aa": Single amino acid Hamming distance matrix from getAAMatrix.
- "hh_s1f": Human single nucleotide distance matrix derived from HH_S1F with calcTargetingDistance.
- "hh_s5f": Human 5-mer nucleotide context distance matrix derived from HH_S5F with calcTargetingDistance.
- "mk_rs1nf": Mouse single nucleotide distance matrix derived from MK_RS1NF with calcTargetingDistance.
- "mk_rs5nf": Mouse 5-mer nucleotide context distance matrix derived from MK_RS1NF with calcTargetingDistance.
- "hs1f_compat": Backwards compatible human single nucleotide distance matrix used in SHazaM v0.1.4 and Change-O v0.3.3.
- "m1n_compat": Backwards compatible mouse single nucleotide distance matrix used in SHazaM v0.1.4 and Change-O v0.3.3.

Note on NAs: if, for a given combination of V gene, J gene, and sequence length, there is only 1 heavy chain sequence (as defined by sequenceColumn), NA is returned instead of a distance (since it has no heavy chain neighbor). If for a given combination there are multiple heavy chain sequences but only 1 unique one, (in which case every heavy chain sequence in this group is the de facto nearest neighbor to each other, thus giving rise to distances of 0), NAs are returned instead of zero-distances.
**distToNearest**

Note on subsample: Subsampling is performed independently in each VJL group for heavy chain sequences. If subsample is larger than number of heavy chain sequences in the group, it is ignored. In other words, subsampling is performed only on groups in which the number of heavy chain sequences is equal to or greater than subsample. DIST_NEAREST has values calculated using all heavy chain sequences in the group for groups with fewer than subsample heavy chain sequences, and values calculated using a subset of heavy chain sequences for the larger groups. To select a value of subsample, it can be useful to explore the group sizes in db (and the number of heavy chain sequences in those groups).

**Value**

Returns a modified db data.frame with nearest neighbor distances between heavy chain sequences in the DIST_NEAREST column if cross=NULL. If cross was specified, distances will be added as the CROSS_DIST_NEAREST column.

Note that distances between light chain sequences are not calculated, even if light chains were used for VJL grouping via groupUsingOnlyIGH=FALSE. Light chain sequences, if any, will have NA in the DIST_NEAREST field.

**References**


**See Also**

See calcTargetingDistance for generating nucleotide distance matrices from a TargetingModel object. See HH_S5F, HH_S1F, MK_RS1NF, getDNAMatrix, and getAAMatrix for individual model details.

**Examples**

```r
# Subset example data to one sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, SAMPLE == "-1h")

# Use genotyped V assignments, Hamming distance, and normalize by junction length
# First partition based on V and J assignments, then by junction length
# Take into consideration ambiguous V and J annotations
dist <- distToNearest(db, vCallColumn="V_CALL_GENOTYPED", model="ham",
                      first=FALSE, VJthenLen=TRUE, normalize="len")

# Plot histogram of non-NA distances
p1 <- ggplot(data=subset(dist, !is.na(DIST_NEAREST))) +
```
theme_bw() +
ggtitle("Distance to nearest: Hamming") +
  xlab("distance") +
  geom_histogram(aes(x=DIST_NEAREST), binwidth=0.025,
  fill="steelblue", color="white")

plot(p1)

---

### editBaseline

>Edit the Baseline object

**Description**

editBaseline edits a field in a Baseline object.

**Usage**

```r
editBaseline(baseline, field, value)
```

**Arguments**

- `baseline` Baseline object to be edited.
- `field` name of the field in the Baseline object to be edited.
- `value` value to set the field.

**Value**

A Baseline object with the field of choice updated.

**See Also**

See Baseline for the input and return object.

**Examples**

```r
# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgG" & SAMPLE == "+7d")

# Make Baseline object
baseline <- calcBaseline(db,
  sequenceColumn="SEQUENCE_IMGT",
  germlineColumn="GERMLINE_IMGT_D_MASK",
  testStatistic="focused",
  regionDefinition=IMGT_V,
  targetingModel=HH_S5F,
  nproc=1)
```
# Edit the field "description"
baseline <- editBaseline(baseline, field="description",
value="+7d IgG")

## expectedMutations

### Description

expectedMutations calculates the expected mutation frequencies for each sequence in the input data.frame.

### Usage

```r
expectedMutations(
  db,
  sequenceColumn = "SEQUENCE_IMGT",
  germlineColumn = "GERMLINE_IMGT_D_MASK",
  targetingModel = HH_S5F,
  regionDefinition = NULL,
  mutationDefinition = NULL,
  nproc = 1
)
```

### Arguments

- **db**: data.frame containing sequence data.
- **sequenceColumn**: character name of the column containing input sequences.
- **germlineColumn**: character name of the column containing the germline or reference sequence.
- **targetingModel**: TargetingModel object. Default is HH_S5F.
- **regionDefinition**: RegionDefinition object defining the regions and boundaries of the Ig sequences.
- **mutationDefinition**: MutationDefinition object defining replacement and silent mutation criteria. If NULL then replacement and silent are determined by exact amino acid identity.
- **nproc**: numeric number of cores to distribute the operation over. If the cluster has already been set the call function with nproc = 0 to not reset or reinitialize. Default is nproc = 1.

### Details

Only the part of the sequences defined in regionDefinition are analyzed. For example, when using the IMGT_V definition, mutations in positions beyond 312 will be ignored.
extendMutabilityMatrix

Value

A modified db data.frame with expected mutation frequencies for each region defined in regionDefinition. The column names are dynamically created based on the regions in regionDefinition. For example, when using the IMGT_V definition, which defines positions for CDR and FWR, the following columns are added:

- MUEXPECTED_CDR_R: number of replacement mutations in CDR1 and CDR2 of the V-segment.
- MUEXPECTED_CDR_S: number of silent mutations in CDR1 and CDR2 of the V-segment.
- MUEXPECTED_FWR_R: number of replacement mutations in FWR1, FWR2 and FWR3 of the V-segment.
- MUEXPECTED_FWR_S: number of silent mutations in FWR1, FWR2 and FWR3 of the V-segment.

See Also

calcExpectedMutations is called by this function to calculate the expected mutation frequencies. See observedMutations for getting observed mutation counts. See IMGT_SCHEMES for a set of predefined RegionDefinition objects.

Examples

```r
# Subset example data
data(ExampleDb, package="alakazam")

db <- subset(ExampleDb, ISOTYPE %in% c("IgA", "IgG") & SAMPLE == "+7d")

# Calculate expected mutations over V region
db_exp <- expectedMutations(db,
  sequenceColumn="SEQUENCE_IMGT",
  germlineColumn="GERMLINE_IMGT_D_MASK",
  regionDefinition=IMGT_V,
  nproc=1)

# Calculate hydropathy expected mutations over V region
db_exp <- expectedMutations(db,
  sequenceColumn="SEQUENCE_IMGT",
  germlineColumn="GERMLINE_IMGT_D_MASK",
  regionDefinition=IMGT_V,
  mutationDefinition=HYDROPATHY_MUTATIONS,
  nproc=1)
```

---

extendMutabilityMatrix

*Extends a mutability model to include Ns.*
**extendSubstitutionMatrix**

*Extends a substitution model to include Ns.*

**Description**

`extendSubstitutionMatrix` extends a 5-mer nucleotide substitution model with 5-mers that include Ns by averaging over all corresponding 5-mers without Ns.

**Usage**

```
extendSubstitutionMatrix(substitutionModel)
```

**Arguments**

- `substitutionModel`
  
  vector of 5-mer substitution rates built by `createSubstitutionMatrix`.

**Value**

A 3125 vector of normalized substitution rates for each 5-mer motif with names defining the 5-mer nucleotide sequence. Note that "normalized" means that the substitution rates for the 1024 5-mers that contain no "N" at any position sums up to 1 (as opposed to the entire vector summing up to 1).

**See Also**

- `createSubstitutionMatrix`
- `createMutabilityMatrix`
- `extendMutabilityMatrix`

**Examples**

```r
# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Create model using only silent mutations and ignore multiple mutations
sub_model <- createSubstitutionMatrix(db, model="S")
mut_model <- createMutabilityMatrix(db, sub_model, model="S")
ext_model <- extendMutabilityMatrix(mut_model)
```

```r
extendSubstitutionMatrix(substitutionModel)
```

---

**extendMutabilityMatrix**

*Extends a 5-mer nucleotide mutability model with 5-mers that include Ns by averaging over all corresponding 5-mers without Ns.*

**Description**

`extendMutabilityMatrix` extends a 5-mer nucleotide mutability model with 5-mers that include Ns by averaging over all corresponding 5-mers without Ns.

**Usage**

```
extendMutabilityMatrix(mutabilityModel)
```

**Arguments**

- `mutabilityModel`
  
  vector of 5-mer mutability rates built by `createMutabilityMatrix`.

**Value**

A 3125 vector of normalized mutability rates for each 5-mer motif with names defining the 5-mer nucleotide sequence. Note that "normalized" means that the mutability rates for the 1024 5-mers that contain no "N" at any position sums up to 1 (as opposed to the entire vector summing up to 1).

**See Also**

- `createMutabilityMatrix`
- `extendSubstitutionMatrix`
findThreshold

Description

findThreshold automatically determines an optimal threshold for clonal assignment of Ig sequences using a vector of nearest neighbor distances. It provides two alternative methods using either a Gamma/Gaussian Mixture Model fit (method="gmm") or kernel density fit (method="density").

Usage

```r
findThreshold(
  distances,
  method = c("density", "gmm"),
  edge = 0.9,
  cross = NULL,
  subsample = NULL,
  model = c("gamma-gamma", "gamma-norm", "norm-gamma", "norm-norm"),
  cutoff = c("optimal", "intersect", "user"),
  sen = NULL,
  spc = NULL,
  progress = FALSE
)
```

Arguments

substitutionModel
matrix of 5-mers substitution counts built by `createSubstitutionMatrix`.

Value

A 5x3125 matrix of normalized substitution rate for each 5-mer motif with rows names defining the center nucleotide, one of c("A", "C", "G", "T", "N"), and column names defining the 5-mer nucleotide sequence.

See Also

createSubstitutionMatrix, extendMutabilityMatrix

Examples

```r
# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Create model using only silent mutations
sub_model <- createSubstitutionMatrix(db, model="S")
ext_model <- extendSubstitutionMatrix(sub_model)
```

---

### findThreshold

Find distance threshold

Description

findThreshold automatically determines an optimal threshold for clonal assignment of Ig sequences using a vector of nearest neighbor distances. It provides two alternative methods using either a Gamma/Gaussian Mixture Model fit (method="gmm") or kernel density fit (method="density").

Usage

```r
findThreshold(
  distances,
  method = c("density", "gmm"),
  edge = 0.9,
  cross = NULL,
  subsample = NULL,
  model = c("gamma-gamma", "gamma-norm", "norm-gamma", "norm-norm"),
  cutoff = c("optimal", "intersect", "user"),
  sen = NULL,
  spc = NULL,
  progress = FALSE
)
```
findThreshold

Arguments

- **distances**: numeric vector containing nearest neighbor distances.
- **method**: string defining the method to use for determining the optimal threshold. One of "gmm" or "density". See Details for methodological descriptions.
- **edge**: upper range as a fraction of the data density to rule initialization of Gaussian fit parameters. Default value is 90. Applies only when method="density".
- **cross**: supplementary nearest neighbor distance vector output from distToNearest for initialization of the Gaussian fit parameters. Applies only when method="gmm".
- **subsample**: maximum number of distances to subsample to before threshold detection.
- **model**: allows the user to choose among four possible combinations of fitting curves: "norm-norm", "norm-gamma", "gamma-norm", and "gamma-gamma". Applies only when method="gmm".
- **cutoff**: method to use for threshold selection: the optimal threshold "opt", the intersection point of the two fitted curves "intersect", or a value defined by user for one of the sensitivity or specificity "user". Applies only when method="gmm".
- **sen**: sensitivity required. Applies only when method="gmm" and cutoff="user".
- **spc**: specificity required. Applies only when method="gmm" and cutoff="user".
- **progress**: if TRUE print a progress bar.

Details

- "gmm": Performs a maximum-likelihood fitting procedure, for learning the parameters of two mixture univariate, either Gamma or Gaussian, distributions which fit the bimodal distribution entries. Retrieving the fit parameters, it then calculates the optimum threshold method="optimal", where the average of the sensitivity plus specificity reaches its maximum. In addition, the findThreshold function is also able to calculate the intersection point (method="intersect") of the two fitted curves and allows the user to invoke its value as the cut-off point, instead of optimal point.
- "density": Fits a binned approximation to the ordinary kernel density estimate to the nearest neighbor distances after determining the optimal bandwidth for the density estimate via least-squares cross-validation of the 4th derivative of the kernel density estimator. The optimal threshold is set as the minimum value in the valley in the density estimate between the two modes of the distribution.

Value

- "gmm" method: Returns a GmmThreshold object including the threshold and the function fit parameters, i.e. mixing weight, mean, and standard deviation of a Normal distribution, or mixing weight, shape and scale of a Gamma distribution.
- "density" method: Returns a DensityThreshold object including the optimum threshold and the density fit parameters.
*Note*

Visually inspecting the resulting distribution fits is strongly recommended when using either fitting method. Empirical observations imply that the bimodality of the distance-to-nearest distribution is detectable for a minimum of 1,000 distances. Larger numbers of distances will improve the fitting procedure, although this can come at the expense of higher computational demands.

*See Also*

See `distToNearest` for generating the nearest neighbor distance vectors. See `plotGmmThreshold` and `plotDensityThreshold` for plotting output.

*Examples*

```r
# Subset example data to one sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, SAMPLE == "-1h")

# Use nucleotide Hamming distance and normalize by junction length
db <- distToNearest(db, model="ham", normalize="len", nproc=1)

# Find threshold using the "gmm" method with optimal threshold
output <- findThreshold(db$DIST_NEAREST, method="gmm", model="gamma-gamma", cutoff="opt")
plot(output, binwidth=0.02, title=paste0(output@model, " loglk=", output@loglk))
print(output)

# Find threshold using the "gmm" method with user defined specificity
output <- findThreshold(db$DIST_NEAREST, method="gmm", model="gamma-gamma",
                        cutoff="user", spc=0.99)
plot(output, binwidth=0.02, title=paste0(output@model, " loglk=", output@loglk))
print(output)

# Find threshold using the "density" method and plot the results
output <- findThreshold(db$DIST_NEAREST, method="density")
plot(output)
print(output)
```

---

**GmmThreshold-class**

Output of the `gmm` method of `findThreshold`

**Description**

`GmmThreshold` contains output from the `gmm` method `findThreshold`. It includes parameters of two Gaussian fits and threshold cut.
GmmThreshold-class

Usage

```r
## S4 method for signature 'GmmThreshold'
print(x)

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

Arguments

- `x`: GmmThreshold object
- `y`: ignored.
- `...`: arguments to pass to `plotGmmThreshold`.

Slots

- `x`: input distance vector with NA or infinite values removed.
- `model`: first-second fit functions.
- `cutoff`: type of threshold cut.
- `a1`: mixing weight of the first curve.
- `b1`: second parameter of the first curve. Either the mean of a Normal distribution or shape of a Gamma distribution.
- `c1`: third parameter of the first curve. Either the standard deviation of a Normal distribution or scale of a Gamma distribution.
- `a2`: mixing weight of the second curve.
- `b2`: second parameter of the second curve. Either the mean of a Normal distribution or shape of a Gamma distribution.
- `c2`: third parameter of the second curve. Either the standard deviation of a Normal distribution or scale of a Gamma distribution.
- `loglk`: log-likelihood of the fit.
- `threshold`: threshold.
- `sensitivity`: sensitivity.
- `specificity`: specificity.
- `pvalue`: p-value from Hartigans' dip statistic (HDS) test. Values less than 0.05 indicate significant bimodality.

See Also

- `findThreshold`
groupBaseline

*Group BASELINe PDFs*

**Description**

groupBaseline convolves groups of BASELINe posterior probability density functions (PDFs) to get combined PDFs for each group.

**Usage**

groupBaseline(baseline, groupBy, nproc = 1)

**Arguments**

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>baseline</td>
<td>Baseline object containing the db and the BASELINe posterior probability density functions (PDF) for each of the sequences, as returned by calcBaseline.</td>
</tr>
<tr>
<td>groupBy</td>
<td>The columns in the db slot of the Baseline object by which to group the sequence PDFs.</td>
</tr>
<tr>
<td>nproc</td>
<td>number of cores to distribute the operation over. If nproc = 0 then the cluster has already been set and will not be reset.</td>
</tr>
</tbody>
</table>

**Details**

While the selection strengths predicted by BASELINe perform well on average, the estimates for individual sequences can be highly variable, especially when the number of mutations is small.

To overcome this, PDFs from sequences grouped by biological or experimental relevance, are convolved to from a single PDF for the selection strength. For example, sequences from each sample may be combined together, allowing you to compare selection across samples. This is accomplished through a fast numerical convolution technique.

**Value**

A Baseline object, containing the modified db and the BASELINe posterior probability density functions (PDF) for each of the groups.

**References**


**See Also**

To generate the Baseline object see calcBaseline. To calculate BASELINe statistics, such as the mean selection strength and the 95% confidence interval, see summarizeBaseline.
Examples

# Subset example data from alakazam
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE %in% c("IgM", "IgG"))

# Collapse clones
db <- collapseClones(db, sequenceColumn="SEQUENCE_IMGT",
                     germlineColumn="GERMLINE_IMGT_D_MASK",
                     method="thresholdedFreq", minimumFrequency=0.6,
                     includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

# Calculate BASELINE
baseline <- calcBaseline(db,
                         sequenceColumn="CLONAL_SEQUENCE",
                         germlineColumn="CLONAL_GERMLINE",
                         testStatistic="focused",
                         regionDefinition=IMGT_V,
                         targetingModel=HH_S5F,
                         nproc=1)

# Group PDFs by sample
grouped1 <- groupBaseline(baseline, groupBy="SAMPLE")
sample_colors <- c("-1h"="steelblue", "+7d"="firebrick")
plotBaselineDensity(grouped1, idColumn="SAMPLE", colorValues=sample_colors,
                     sigmaLimits=c(-1, 1))

# Group PDFs by both sample (between variable) and isotype (within variable)
grouped2 <- groupBaseline(baseline, groupBy=c("SAMPLE", "ISOTYPE"))
isotype_colors <- c("IgM"="darkorchid", "IgD"="firebrick",
                    "IgG"="seagreen", "IgA"="steelblue")
plotBaselineDensity(grouped2, idColumn="SAMPLE", groupColumn="ISOTYPE",
                    colorElement="group", colorValues=isotype_colors,
                    sigmaLimits=c(-1, 1))

# Collapse previous isotype (within variable) grouped PDFs into sample PDFs
grouped3 <- groupBaseline(grouped2, groupBy="SAMPLE")
sample_colors <- c("-1h"="steelblue", "+7d"="firebrick")
plotBaselineDensity(grouped3, idColumn="SAMPLE", colorValues=sample_colors,
                     sigmaLimits=c(-1, 1))

HH_S1F

Human heavy chain, silent, 1-mer, functional substitution model.

Description

1-mer substitution model of somatic hypermutation based on analysis of silent mutations in functional heavy chain Ig sequences from Homo sapiens.
Usage

HH_S1F

Format

A 4x4 matrix of nucleotide substitution rates. The rates are normalized, therefore each row sums up to 1.

Note

HH_S1F replaces HS1Distance in versions of SHazaM prior to 0.1.5.

References


See Also

See HKL_S1F for the human light chain 1-mer substitution model and MK_RS1NF for the mouse light chain 1-mer substitution model.

HH_S5F

Human heavy chain, silent, 5-mer, functional targeting model.

Description

5-mer model of somatic hypermutation targeting based on analysis of silent mutations in functional heavy chain Ig sequences from Homo sapiens.

Usage

HH_S5F

Format

A TargetingModel object.

References

See Also

See HH_S1F for the 1-mer substitution matrix from the same publication; HKL_S5F for the human light chain 5-mer targeting model; MK_RS5NF for the mouse 5-mer targeting model; and U5N for the uniform 5-mer null targeting model.

| HKL_S1F |
|---------|------------------|
|         | *Human kappa and lambda chain, silent, 1-mer, functional substitution model.* |

Description

1-mer substitution model of somatic hypermutation based on analysis of silent mutations in functional kappa and lambda light chain Ig sequences from Homo sapiens.

Usage

HKL_S1F

Format

A 4x4 matrix of nucleotide substitution rates. The rates are normalized, therefore each row sums up to 1.

Note

Reported in Table III in Cui et al, 2016.

References


See Also

See HH_S1F for the human heavy chain 1-mer substitution model and MK_RS1NF for the mouse light chain 1-mer substitution model.
HKL_S5F  Human kappa and lambda light chain, silent, 5-mer, functional targeting model.

Description

5-mer model of somatic hypermutation targeting based on analysis of silent mutations in functional kappa and lambda light chain Ig sequences from Homo sapiens.

Usage

HKL_S5F

Format

A TargetingModel object.

References


See Also

See HH_S5F for the human heavy chain 5-mer targeting model; MK_RS5NF for the mouse kappa light chain 5-mer targeting model; and U5N for the uniform 5-mer null targeting model.

IMGT_SCHEMES  IMGT unique numbering schemes

Description

Sequence region definitions according to the IMGT unique numbering scheme.

Format

A RegionDefinition object defining:

- IMGT_V: The IMGT numbered V segment up to position nucleotide 312. This definition combines the CDR1 and CDR2 into a single CDR region, and FWR1, FWR2 and FWR3 into a single FWR region. CDR3 and FWR4 are excluded as they are downstream of nucleotide 312.
- IMGT_V_BY_CODONS: The IMGT numbered V segment up to position nucleotide 312. This definition treats each codon, from codon 1 to codon 104, as a distinct region.
- IMGT_V_BY_REGIONS: The IMGT numbered V segment up to position nucleotide 312. This defines separate regions for each of CDR1, CDR2, FWR1, FWR2 and FWR3. CDR3 and FWR4 are excluded as they are downstream of nucleotide 312.
- IMGT_V_BY_SEGMENTS: The IMGT numbered V segment up to position nucleotide 312. This definition has no subdivisions and treats the entire V segment as a single region.

References


---

**makeAverage1merMut**

Make a 1-mer mutability model by averaging over a 5-mer mutability model

**Description**

makeAverage1merMut averages mutability rates in a 5-mer mutability model to derive a 1-mer mutability model.

**Usage**

```r
makeAverage1merMut(mut5mer)
```

**Arguments**

- `mut5mer` a named vector of length 1024 such as that returned by `createMutabilityMatrix` and that returned by `makeDegenerate5merMut` with extended=FALSE. Names should correspond to 5-mers made up of "A", "T", "G", and "C" (case-insensitive). NA values are allowed.

**Details**

For example, the mutability rate of "A" in the resultant 1-mer model is derived by averaging the mutability rates of all the 5-mers that have an "A" as their central 1-mer, followed by normalization.

**Value**

A named vector of length 4 containing normalized mutability rates.

**See Also**

See `makeDegenerate5merMut` for making a degenerate 5-mer mutability model based on a 1-mer mutability model.
Examples

# Make a degenerate 5-mer model (length of 1024) based on a 1-mer model
example1merMut <- c(A=0.2, T=0.1, C=0.4, G=0.3)
degenerate5merMut <- makeDegenerate5merMut(mut1mer = example1merMut)

# Now make a 1-mer model by averaging over the degenerate 5-mer model
# Expected to get back example1merMut
makeAverage1merMut(mut5mer = degenerate5merMut)

makeAverage1merSub

Make a 1-mer substitution model by averaging over a 5-mer substitution model

Description

makeAverage1merSub averages substitution rates in a 5-mer substitution model to derive a 1-mer substitution model.

Usage

makeAverage1merSub(sub5mer)

Arguments

sub5mer  a 4x1024 matrix such as that returned by createSubstitutionMatrix and that returned by makeDegenerate5merSub with extended=FALSE. Column names should correspond to 5-mers containing the central 1-mer to mutate from. Row names should correspond to nucleotides to mutate into. Nucleotides should include "A", "T", "G", and "C" (case-insensitive).

Details

For example, the substitution rate from "A" to "T" in the resultant 1-mer model is derived by averaging the substitution rates into a "T" of all the 5-mers that have an "A" as their central 1-mer.

Value

A 4x4 matrix with row names representing nucleotides to mutate from and column names representing nucleotides to mutate into. Rates are normalized by row.

See Also

See makeDegenerate5merSub for making a degenerate 5-mer substitution model based on a 1-mer substitution model.
Examples

# Make a degenerate 5-mer model (4x1024) based on HKL_S1F (4x4)
degenerate5merSub <- makeDegenerate5merSub(sub1mer = HKL_S1F)

# Now make a 1-mer model by averaging over the degenerate 5-mer model
# Expected to get back HKL_S1F
makeAverage1merSub(sub5mer = degenerate5merSub)

makeDegenerate5merMut Make a degenerate 5-mer mutability model based on a 1-mer mutability model

Description

makeDegenerate5merMut populates mutability rates from a 1-mer mutability model into 5-mers with corresponding central 1-mers.

Usage

makeDegenerate5merMut(mut1mer, extended = FALSE)

Arguments

mut1mer a named vector of length 4 containing (normalized) mutability rates. Names should correspond to nucleotides, which should include "A", "T", "G", and "C" (case-insensitive).

extended whether to return the unextended (extended=FALSE) or extended (extended=TRUE) 5-mer mutability model. Default is FALSE.

Details

As a concrete example, consider a 1-mer mutability model in which mutability rates of "A", "T", "G", and "C" are, respectively, 0.14, 0.23, 0.31, and 0.32. In the resultant degenerate 5-mer mutability model, all the 5-mers that have an "A" as their central 1-mer would have mutability rate of 0.14/256, where 256 is the number of such 5-mers.

When extended=TRUE, extendMutabilityMatrix is called to extend the mutability vector of length 1024 into a vector of length 3125.

Value

For extended=FALSE, a vector of length 1024. The vector returned is normalized. For extended=TRUE, a vector of length 3125.

See Also

See makeAverage1merMut for making a 1-mer mutability model by taking the average of a 5-mer mutability model. See extendMutabilityMatrix for extending the mutability vector.
Examples

# Make a degenerate 5-mer model (length of 1024) based on a 1-mer model
example1merMut <- c(A=0.2, T=0.1, C=0.4, G=0.3)
degenerate5merMut <- makeDegenerate5merSub(mut1mer = example1merMut)

# Look at a few 5-mers
degenerate5merMut[c("AAAAT", "AACAT", "AAGAT", "AATAT")]

# Normalized
sum(degenerate5merMut)

makeDegenerate5merSub  Make a degenerate 5-mer substitution model based on a 1-mer substitution model

Description

makeDegenerate5merSub populates substitution rates from a 1-mer substitution model into 5-mers with corresponding central 1-mers.

Usage

makeDegenerate5merSub(sub1mer, extended = FALSE)

Arguments

sub1mer a 4x4 matrix containing (normalized) substitution rates. Row names should correspond to nucleotides to mutate from. Column names should correspond to nucleotides to mutate into. Nucleotides should include "A", "T", "G", and "C" (case-insensitive).

extended whether to return the unextended (extended=FALSE) or extended (extended=TRUE) 5-mer substitution model. Default is FALSE.

Details

As a concrete example, consider a 1-mer substitution model in which substitution rates from "A" to "T", "G", and "C" are, respectively, 0.1, 0.6, and 0.3. In the resultant degenerate 5-mer substitution model, all the 5-mers (columns) that have an "A" as their central 1-mer would have substitution rates (rows) of 0.1, 0.6, and 0.3 to "T", "G", and "C" respectively.

When extended=TRUE, extendSubstitutionMatrix is called to extend the 4x1024 substitution matrix.

Value

For extended=FALSE, a 4x1024 matrix. For extended=TRUE, a 5x3125 matrix.
See Also

See `makeAverage1merSub` for making a 1-mer substitution model by taking the average of a 5-mer substitution model. See `extendSubstitutionMatrix` for extending the substitution matrix.

Examples

```r
# Make a degenerate 5-mer model (4x1024) based on HKL_S1F (4x4)
# Note: not to be confused with HKL_S5F@substitution, which is non-degenerate
degenerate5merSub <- makeDegenerate5merSub(sub1mer = HKL_S1F)

# Look at a few 5-mers
degenerate5merSub[, c('AAAAT', 'AACAT', 'AAGAT', 'AATAT')]
```

**minNumMutationsTune Parameter tuning for minNumMutations**

Description

`minNumMutationsTune` helps with picking a threshold value for `minNumMutations` in `createSubstitutionMatrix` by tabulating the number of 5-mers for which substitution rates would be computed directly or inferred at various threshold values.

Usage

```r
minNumMutationsTune(subCount, minNumMutationsRange)
```

Arguments

- `subCount` data.frame returned by `createSubstitutionMatrix` with `numMutationsOnly=TRUE`.
- `minNumMutationsRange` a number or a vector indicating the value or range of values of `minNumMutations` to try.

Details

At a given threshold value of `minNumMutations`, for a given 5-mer, if the total number of mutations is greater than the threshold and there are mutations to every other base, substitution rates are computed directly for the 5-mer using its mutations. Otherwise, mutations from 5-mers with the same inner 3-mer as the 5-mer of interest are aggregated. If the number of such mutations is greater than the threshold and there are mutations to every other base, these mutations are used for inferring the substitution rates for the 5-mer of interest; if not, mutations from all 5-mers with the same center nucleotide are aggregated and used for inferring the substitution rates for the 5-mer of interest (i.e. the 1-mer model).
Value

A 3xn matrix, where n is the number of trial values of minNumMutations supplied in minNumMutationsRange. Each column corresponds to a value in minNumMutationsRange. The rows correspond to the number of 5-mers for which substitution rates would be computed directly using the 5-mer itself ("5mer"), using its inner 3-mer ("3mer"), and using the central 1-mer ("1mer"), respectively.

References


See Also

See argument numMutationsOnly in createSubstitutionMatrix for generating the required input data.frame subCount. See argument minNumMutations in createSubstitutionMatrix for what it does.

Examples

# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Count the number of mutations per 5-mer
subCount <- createSubstitutionMatrix(db, model="S", multipleMutation="independent",
                                      returnModel="5mer", numMutationsOnly=TRUE)

# Tune minNumMutations
minNumMutationsTune(subCount, seq(from=10, to=100, by=10))
Arguments

mutCount a vector of length 1024 returned by createMutabilityMatrix with numSeqMutationsOnly=TRUE.

minNumSeqMutationsRange

a number or a vector indicating the value or the range of values of minNumSeqMutations to try.

Details

At a given threshold value of minNumSeqMutations, for a given 5-mer, if the total number of mutations is greater than the threshold, mutability is computed directly. Otherwise, mutability is inferred.

Value

A 2xn matrix, where n is the number of trial values of minNumSeqMutations supplied in minNumSeqMutationsRange. Each column corresponds to a value in minNumSeqMutationsRange. The rows correspond to the number of 5-mers for which mutability would be computed directly ("measured") and inferred ("inferred"), respectively.

References


See Also

See argument numSeqMutationsOnly in createMutabilityMatrix for generating the required input vector mutCount. See argument minNumSeqMutations in createMutabilityMatrix for what it does.

Examples

```r
# Subset example data to one isotype and sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA" & SAMPLE == "-1h")

# Create model using only silent mutations
sub <- createSubstitutionMatrix(db, model="S", multipleMutation="independent",
                                 returnModel="5mer", numMutationsOnly=FALSE,
                                 minNumMutations=20)

# Count the number of mutations in sequences containing each 5-mer
mutCount <- createMutabilityMatrix(db, substitutionModel = sub,
                                    model="S", multipleMutation="independent",
                                    numSeqMutationsOnly=TRUE)

# Tune minNumSeqMutations
minNumSeqMutationsTune(mutCount, seq(from=100, to=300, by=50))
```
**MK_RS1NF**

*Mouse kappa chain, replacement and silent, 1-mer, non-functional substitution model.*

**Description**

1-mer substitution model of somatic hypermutation based on analysis of replacement and silent mutations in non-functional kappa light chain Ig sequences from NP-immunized Mus musculus.

**Usage**

MK_RS1NF

**Format**

A 4x4 matrix of nucleotide substitution rates. The rates are normalized, therefore each row sums up to 1.

**Note**

MK_RS1NF replaces M1NDistance from versions of SHazaM prior to 0.1.5.

**References**


**See Also**

See HH_S1F for the human heavy chain 1-mer substitution model and HKL_S1F for the human light chain 1-mer substitution model.

---

**MK_RS5NF**

*Mouse kappa light chain, replacement and silent, 5-mer, non-functional targeting model.*

**Description**

5-mer model of somatic hypermutation targeting based on analysis of replacement and silent mutations in non-functional kappa light chain Ig sequences from NP-immunized Mus musculus.

**Usage**

MK_RS5NF
MutationDefinition-class

Format

TargetingModel object.

References


See Also

See MK_RS1NF for the 1-mer substitution matrix from the same publication; HH_S5F for the human heavy chain silent 5-mer functional targeting model; HKL_S5F for the human light chain silent 5-mer functional targeting model; and U5N for the uniform 5-mer null targeting model.

S4 class defining replacement and silent mutation definitions

Description

MutationDefinition defines a common data structure for defining the whether a mutation is annotated as a replacement or silent mutation.

Slots

name name of the MutationDefinition.

description description of the model and its source.

classes named character vectors with single-letter amino acid codes as names and amino acid classes as values, with NA assigned to set of characters c("X","*","-","."). Replacement (R) is be defined as a change in amino acid class and silent (S) as no change in class.

codonTable matrix of codons (columns) and substitutions (rows).

citation publication source.

See Also

See MUTATION_SCHEMES for a set of predefined MutationDefinition objects.
MUTATION_SCHEMES

Amino acid mutation definitions

Description

Definitions of replacement (R) and silent (S) mutations for different amino acid physicochemical classes.

Format

A MutationDefinition object defining:

- **CHARGE_MUTATIONS**: Amino acid mutations are defined by changes in side chain charge class.
- **HYDROPATHY_MUTATIONS**: Amino acid mutations are defined by changes in side chain hydrophobicity class.
- **POLARITY_MUTATIONS**: Amino acid mutations are defined by changes in side chain polarity class.
- **VOLUME_MUTATIONS**: Amino acid mutations are defined by changes in side chain volume class.

References

1. [http://www.imgt.org/IMGTeducation/Aide-memoire/_UK/aminoacids/IMGTclasses.html](http://www.imgt.org/IMGTeducation/Aide-memoire/_UK/aminoacids/IMGTclasses.html)

observedMutations

Calculate observed numbers of mutations

Description

observedMutations calculates the observed number of mutations for each sequence in the input data.frame.

Usage

```r
observedMutations(
  db,
  sequenceColumn = "SEQUENCE_IMGT",
  germlineColumn = "GERMLINE_IMGT_D_MASK",
  regionDefinition = NULL,
  mutationDefinition = NULL,
  ambiguousMode = c("eitherOr", "and"),
  frequency = FALSE,
  combine = FALSE,
  nproc = 1
)
```
observedMutations

Arguments

- **db**: data.frame containing sequence data.
- **sequenceColumn**: character name of the column containing input sequences. IUPAC ambiguous characters for DNA are supported.
- **germlineColumn**: character name of the column containing the germline or reference sequence. IUPAC ambiguous characters for DNA are supported.
- **regionDefinition**: RegionDefinition object defining the regions and boundaries of the Ig sequences. If NULL, mutations are counted for entire sequence.
- **mutationDefinition**: MutationDefinition object defining replacement and silent mutation criteria. If NULL then replacement and silent are determined by exact amino acid identity.
- **ambiguousMode**: whether to consider ambiguous characters as "either or" or "and" when determining and counting the type(s) of mutations. Applicable only if sequenceColumn and/or germlineColumn contain(s) ambiguous characters. One of c("eitherOr", "and"). Default is "eitherOr".
- **frequency**: logical indicating whether or not to calculate mutation frequencies. Default is FALSE.
- **combine**: logical indicating whether for each sequence should the mutation counts for the different regions (CDR, FWR) and mutation types be combined and return one value of count/frequency per sequence instead of multiple values. Default is FALSE.
- **nproc**: number of cores to distribute the operation over. If the cluster has already been set the call function with nproc = 0 to not reset or reinitialize. Default is nproc = 1.

Details

Mutation counts are determined by comparing the input sequences (in the column specified by sequenceColumn) to the germline sequence (in the column specified by germlineColumn). See calcObservedMutations for more technical details, including criteria for which sequence differences are included in the mutation counts and which are not.

The mutations are binned as either replacement (R) or silent (S) across the different regions of the sequences as defined by regionDefinition. Typically, this would be the framework (FWR) and complementarity determining (CDR) regions of IMGT-gapped nucleotide sequences. Mutation counts are appended to the input db as additional columns.

Value

A modified db data.frame with observed mutation counts for each sequence listed. The columns names are dynamically created based on the regions in the regionDefinition. For example, when using the IMGT_V definition, which defines positions for CDR and FWR, the following columns are added:

- **MU_COUNT_CDR_R**: number of replacement mutations in CDR1 and CDR2 of the V-segment.
- **MU_COUNT_CDR_S**: number of silent mutations in CDR1 and CDR2 of the V-segment.
• **MU_COUNT_FWR_R**: number of replacement mutations in FWR1, FWR2 and FWR3 of the V-segment.
• **MU_COUNT_FWR_S**: number of silent mutations in FWR1, FWR2 and FWR3 of the V-segment.

If `frequency=TRUE`, R and S mutation frequencies are calculated over the number of non-N positions in the specified regions.

• **MU_FREQ_CDR_R**: frequency of replacement mutations in CDR1 and CDR2 of the V-segment.
• **MU_FREQ_CDR_S**: frequency of silent mutations in CDR1 and CDR2 of the V-segment.
• **MU_FREQ_FWR_R**: frequency of replacement mutations in FWR1, FWR2 and FWR3 of the V-segment.
• **MU_FREQ_FWR_S**: frequency of silent mutations in FWR1, FWR2 and FWR3 of the V-segment.

If `frequency=TRUE` and `combine=TRUE`, the mutations and non-N positions are aggregated and a single **MU_FREQ** value is returned

• **MU_FREQ**: frequency of replacement and silent mutations in the specified region

**See Also**

calcObservedMutations is called by this function to get the number of mutations in each sequence grouped by the `RegionDefinition`. See IMGT_SCHEMES for a set of predefined `RegionDefinition` objects. See expectedMutations for calculating expected mutation frequencies.

**Examples**

# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgG" & SAMPLE == "+7d")

# Calculate mutation frequency over the entire sequence
db_obs <- observedMutations(db, sequenceColumn="SEQUENCE_IMGT", germlineColumn="GERMLINE_IMGT_D_MASK", frequency=TRUE, nproc=1)

# Count of V-region mutations split by FWR and CDR
# With mutations only considered replacement if charge changes
db_obs <- observedMutations(db, sequenceColumn="SEQUENCE_IMGT", germlineColumn="GERMLINE_IMGT_D_MASK", regionDefinition=IMGT_V, mutationDefinition=CHARGE_MUTATIONS, nproc=1)
Description

plotBaselineDensity plots the probability density functions resulting from selection analysis using the BASELINe method.

Usage

plotBaselineDensity(
  baseline,
  idColumn,
  groupColumn = NULL,
  colorElement = c("id", "group"),
  colorValues = NULL,
  title = NULL,
  subsetRegions = NULL,
  sigmaLimits = c(-5, 5),
  facetBy = c("region", "group"),
  style = c("density"),
  sizeElement = c("none", "id", "group"),
  size = 1,
  silent = FALSE,
  ...
)

Arguments

baseline Baseline object containing selection probability density functions.
idColumn name of the column in the db slot of baseline containing primary identifiers.
groupColumn name of the column in the db slot of baseline containing secondary grouping identifiers. If NULL, organize the plot only on values in idColumn.
colorElement one of c("id","group") specifying whether the idColumn or groupColumn will be used for color coding. The other entry, if present, will be coded by line style.
colorValues named vector of colors for entries in colorElement, with names defining unique values in the colorElement column and values being colors. Also controls the order in which values appear on the plot. If NULL alphabetical ordering and a default color palette will be used.
title string defining the plot title.
subsetRegions character vector defining a subset of regions to plot, corresponding to the regions for which the baseline data was calculated. If NULL all regions in baseline are plotted.
plotBaselineDensity

 sigmaLimits numeric vector containing two values defining the \( \text{c(lower,upper)} \) bounds of the selection scores to plot.

 facetBy one of \( \text{c("region","group")} \) specifying which category to facet the plot by, either values in \text{groupColumn \("group"\)} or regions defined in the \text{regions slot} of the baseline object \("region"\). If this is set to "group", then the region will behave as the \text{groupColumn} for purposes of the \text{colorElement} argument.

 style type of plot to draw. One of:
  
  - "density": plots a set of curves for each probability density function in \text{baseline}, with colors determined by values in the \text{colorElement} column. Faceting is determined by the \text{facetBy} argument.

 sizeElement one of \( \text{c("none","id","group")} \) specifying whether the lines in the plot should be all of the same size \( \text{(none)} \) or have their sizes depend on the values in \text{id} or \text{code}.

 size numeric scaling factor for lines, points and text in the plot.

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

 ... additional arguments to pass to \text{ggplot2::theme}.

 Value

 A \text{ggplot} object defining the plot.

 See Also

 Takes as input a \text{Baseline} object returned from \text{groupBaseline}.

 Examples

 # Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE %in% c("IgM", "IgG"))

 # Collapse clones
db <- collapseClones(db, sequenceColumn="SEQUENCE_IMGT",
 germlineColumn="GERMLINE_IMGT_D_MASK",
 method="thresholdedFreq", minimumFrequency=0.6,
 includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

 # Calculate BASELINE
baseline <- calcBaseline(db,
 sequenceColumn="CLONAL_SEQUENCE",
 germlineColumn="CLONAL_GERMLINE",
 testStatistic="focused",
 regionDefinition=IMGT_V,
 targetingModel=HH_S5F,
 nproc=1)

 # Grouping the PDFs by the sample and isotype annotations
grouped <- groupBaseline(baseline, groupBy=c("SAMPLE", "ISOTYPE"))

# Plot density faceted by region with custom isotype colors
isotype_colors <- c("IgM"="darkorchid", "IgD"="firebrick",
"IgG"="seagreen", "IgA"="steelblue")
plotBaselineDensity(grouped, "SAMPLE", "ISOTYPE", colorValues=isotype_colors,
                colorElement="group", sigmaLimits=c(-1, 1))

# Facet by isotype instead of region
sample_colors <- c("-1h"="steelblue", "+7d"="firebrick")
plotBaselineDensity(grouped, "SAMPLE", "ISOTYPE", facetBy="group",
                colorValues=sample_colors, sigmaLimits=c(-1, 1))

---

**plotBaselineSummary**  
Plots BASELINe summary statistics

**Description**
plotBaselineSummary plots a summary of the results of selection analysis using the BASELINe method.

**Usage**

```r
plotBaselineSummary(
  baseline,
  idColumn,
  groupColumn = NULL,
  groupColors = NULL,
  subsetRegions = NULL,
  facetBy = c("region", "group"),
  title = NULL,
  style = c("summary"),
  size = 1,
  silent = FALSE,
  ...
)
```

**Arguments**

- `baseline`: either a data.frame returned from `summarizeBaseline` or a Baseline object returned from `groupBaseline` containing selection probability density functions and summary statistics.
- `idColumn`: name of the column in baseline containing primary identifiers. If the input is a Baseline object, then this will be a column in the stats slot of baseline.
groupColumn

name of the column in baseline containing secondary grouping identifiers. If the input is a Baseline object, then this will be a column in the stats slot of baseline.

groupColors
	named vector of colors for entries in groupColumn, with names defining unique values in the groupColumn and values being colors. Also controls the order in which groups appear on the plot. If NULL alphabetical ordering and a default color palette will be used. Has no effect if facetBy="group".

subsetRegions

character vector defining a subset of regions to plot, corresponding to the regions for which the baseline data was calculated. If NULL all regions in baseline are plotted.

facetBy

one of c("group", "region") specifying which category to facet the plot by, either values in groupColumn ("group") or regions defined in baseline ("region"). The data that is not used for faceting will be color coded.

title

string defining the plot title.

style

type of plot to draw. One of:

- "summary": plots the mean and confidence interval for the selection scores of each value in idColumn. Faceting and coloring are determined by values in groupColumn and regions defined in baseline, depending upon the facetBy argument.

size

numeric scaling factor for lines, points and text 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

Takes as input either a Baseline object returned by groupBaseline or a data.frame returned from summarizeBaseline.

Examples

# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE %in% c("IgM", "IgG"))

# Collapse clones
db <- collapseClones(db, sequenceColumn="SEQUENCE_IMGT",
                      germlineColumn="GERMLINE_IMGT_D_MASK",
                      method="thresholdedFreq", minimumFrequency=0.6,
                      includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

# Calculate BASELINE
baseline <- calcBaseline(db,
    sequenceColumn="CLONAL_SEQUENCE",
    germlineColumn="CLONAL_GERMLINE",
    testStatistic="focused",
    regionDefinition=IMGT_V,
    targetingModel=HH_S5F,
    nproc=1)

# Grouping the PDFs by sample and isotype annotations
grouped <- groupBaseline(baseline, groupBy=c("SAMPLE", "ISOTYPE"))

# Plot mean and confidence interval by region with custom group colors
isotype_colors <- c("IgM"="darkorchid", "IgD"="firebrick",
    "IgG"="seagreen", "IgA"="steelblue")
plotBaselineSummary(grouped, "SAMPLE", "ISOTYPE",
    groupColors=isotype_colors)

# Facet by group instead of region
plotBaselineSummary(grouped, "SAMPLE", "ISOTYPE", facetBy="group")

---

plotDensityThreshold  Plot findThreshold results for the density method

Description

plotDensityThreshold plots the results from "density" method of findThreshold, including the smoothed density estimate, input nearest neighbor distance histogram, and threshold selected.

Usage

plotDensityThreshold(
    data,
    cross = NULL,
    xmin = NULL,
    xmax = NULL,
    breaks = NULL,
    binwidth = NULL,
    title = NULL,
    size = 1,
    silent = FALSE,
    ...
)

Arguments

data  DensityThreshold object output by the “density” method of findThreshold.
cross numeric vector of distances from `distToNearest` to draw as a histogram below the data histogram for comparison purposes.

`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 to the bandwidth parameter determined by `findThreshold`.

`title` string defining the plot title.

`size` numeric value for the plot line sizes.

`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 `DensityThreshold` for the the input object definition and `findThreshold` for generating the input object. See `distToNearest` calculating nearest neighbor distances.

Examples

```r
# Subset example data to one sample as a demo
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, SAMPLE == "-1h")

# Use nucleotide Hamming distance and normalize by junction length
db <- distToNearest(db, model="ham", normalize="len", nproc=1)

# To find the threshold cut, call findThreshold function for "gmm" method.
output <- findThreshold(db$DIST_NEAREST, method="density")
print(output)

# Plot
plotDensityThreshold(output)
```
plotGmmThreshold

Description

plotGmmThreshold plots the results from "gmm" method of findThreshold, including the Gaussian distributions, input nearest neighbor distance histogram, and threshold selected.

Usage

plotGmmThreshold(
  data,
  cross = NULL,
  xmin = NULL,
  xmax = NULL,
  breaks = NULL,
  binwidth = NULL,
  title = NULL,
  size = 1,
  silent = FALSE,
  ...
)

Arguments

data [GmmThreshold object output by the "gmm" method of findThreshold.]
cross numeric vector of distances from distToNearest to draw as a histogram below
          the data histogram for comparison purposes.
xmin minimal 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.
plotMutability

Plot mutability probabilities

Description

plotMutability plots the mutability rates of a TargetingModel.

Usage

plotMutability(
  model,
  nucleotides = c("A", "C", "G", "T"),
  mark = NULL,
  style = c("hedgehog", "bar"),
  size = 1,
  silent = FALSE,
  ...
)

Arguments

model TargetingModel object or vector containing normalized mutability rates.
nucleotides vector of center nucleotide characters to plot.
mark vector of 5-mer motifs to highlight in the plot. If NULL only highlight classical hot and cold spot motifs.
**style**

type of plot to draw. One of:

- "hedgehog": circular plot showing higher mutability scores further from the circle. The 5-mer is denoted by the values of the inner circle. The 5-mer is read from the most interior position of the 5-mer (5') to most exterior position (3'), with the center nucleotide in the center ring. Note, the order in which the 5-mers are plotted is different for nucleotides c("A", "C") and c("G", "T").
- "bar": bar plot of mutability similar to the hedgehog style with the most 5' positions of each 5-mer at the base of the plot.

**size**

numeric scaling factor for lines and text in the plot.

**silent**

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

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

**Value**

A named list of ggplot objects defining the plots, with names defined by the center nucleotide for the plot object.

**See Also**

Takes as input a TargetingModel object. See createTargetingModel for model building.

**Examples**

```r
# Plot one nucleotide in circular style
plotMutability(HH_S5F, "C")

# Plot two nucleotides in barchart style
plotMutability(HH_S5F, c("G", "T"), style="bar")
```

---

**plotTune**

*Visualize parameter tuning for minNumMutations and minNumSeqMutations*

**Description**

Visualize results from minNumMutationsTune and minNumSeqMutationsTune

**Usage**

```r
plotTune(
  tuneMtx,
  thresh,
  criterion = c("5mer", "3mer", "1mer", "3mer+1mer", "measured", "inferred"),
  pchs = 1,
)```
Arguments

- `tuneMtx`: a matrix or a list of matrices produced by either `minNumMutationsTune` or `minNumSeqMutationsTune`. In the case of a list, it is assumed that each matrix corresponds to a sample and that all matrices in the list were produced using the same set of trial values of `minNumMutations` or `minNumSeqMutations`.
- `thresh`: a number or a vector of indicating the value or the range of values of `minNumMutations` or `minNumSeqMutations` to plot. Should correspond to the columns of `tuneMtx`.
- `criterion`: one of "5mer", "3mer", "1mer", or "3mer+1mer" (for `tuneMtx` produced by `minNumMutationsTune`), or either "measured" or "inferred" (for `tuneMtx` produced by `minNumSeqMutationsTune`).
- `pchs`: point types to pass on to `plot`.
- `ltys`: line types to pass on to `plot`.
- `cols`: colors to pass on to `plot`.
- `plotLegend`: whether to plot legend. Default is `TRUE`. Only applicable if `tuneMtx` is a named list with names of the matrices corresponding to the names of the samples.
- `legendPos`: position of legend to pass on to `legend`. Can be either a numeric vector specifying x-y coordinates, or one of "topright", "center", etc. Default is "topright".
- `legendHoriz`: whether to make legend horizontal. Default is `FALSE`.
- `legendCex`: numeric values by which legend should be magnified relative to 1.

Details

For `tuneMtx` produced by `minNumMutationsTune`, for each sample, depending on `criterion`, the numbers of 5-mers for which substitution rates are directly computed ("5mer"), inferred based on inner 3-mers ("3mer"), inferred based on central 1-mers ("1mer"), or inferred based on inner 3-mers and central 1-mers ("3mer+1mer") are plotted on the y-axis against values of `minNumMutations` on the x-axis.

For `tuneMtx` produced by `minNumSeqMutationsTune`, for each sample, depending on `criterion`, the numbers of 5-mers for which mutability rates are directly measured ("measured") or inferred ("inferred") are plotted on the y-axis against values of `minNumSeqMutations` on the x-axis.

Note that legends will be plotted only if `tuneMtx` is a supplied as a named list of matrices, ideally with names of each matrix corresponding to those of the samples based on which the matrices were produced, even if `plotLegend=TRUE`.

See Also

See `minNumMutationsTune` and `minNumSeqMutationsTune` for generating `tuneMtx`.
Examples

```r
# Subset example data to one isotype and sample as demos
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgA")

tuneMtx = list()
for (i in 1:length(unique(db$SAMPLE))) {
  # Get data corresponding to current sample
  curDb = db[db$SAMPLE == unique(db$SAMPLE)[i], ]

  # Count the number of mutations per 5-mer
  subCount = createSubstitutionMatrix(db=curDb, model="S", multipleMutation="independent",
                                       returnModel="5mer", numMutationsOnly=TRUE)

  # Tune over minNumMutations = 5..50
  subTune = minNumMutationsTune(subCount, seq(from=5, to=50, by=5))

  tuneMtx = c(tuneMtx, list(subTune))
}

# Name tuneMtx after sample names
names(tuneMtx) = unique(db$SAMPLE)

# plot with legend for both samples for a subset of minNumMutations values
plotTune(tuneMtx, thresh=c(5, 15, 25, 40), criterion="3mer",
         pchs=16:17, lty=1:2, cols=2:3,
         plotLegend=TRUE, legendPos=c(5, 100))

# plot for only 1 sample for all the minNumMutations values (no legend)
plotTune(tuneMtx[[1]], thresh=seq(from=5, to=50, by=5), criterion="3mer")
```

RegionDefinition-class

*S4 class defining a region definition*

**Description**

RegionDefinition defines a common data structure for defining the region boundaries of an Ig sequence.

**Slots**

- `name` name of the RegionDefinition.
- `description` description of the model and its source.
- `boundaries` factor defining the region boundaries of the sequence. The levels and values of boundaries determine the number of regions.
seqLength  length of the sequence.
regions  levels of the boundaries; e.g. c("CDR","FWR").
labels  labels for the boundary and mutations combinations; e.g., c("CDR_R","CDR_S","FWR_R","FWR_S").
citation  publication source.

See Also

See IMGT_SCHEMES for a set of predefined RegionDefinition objects.

---

**Description**

Dramatic improvements in high-throughput sequencing technologies now enable large-scale characterization of Ig repertoires, defined as the collection of transmembrane antigen-receptor proteins located on the surface of T and B lymphocytes. The shazam package provides tools for advanced analysis of somatic hypermutation (SHM) in immunoglobulin (Ig) sequences. The key functions in shazam, broken down topic, are described below.

**Mutational profiling**

shazam provides tools to quantify the extent and nature of SHM within full length V(D)J sequences as well as sub-regions (eg, FWR and CDR). Quantification of expected mutational loaded, under specific SHM targeting models, can also be performed along with model driven simulations of SHM.

- collapseClones: Build clonal consensus sequences.
- consensusSequence: Build a single consensus sequence.
- observedMutations: Compute observed mutation counts and frequencies.
- expectedMutations: Compute expected mutation frequencies.
- shmulateSeq: Simulate mutations in a single sequence.
- shmulateTree: Simulate mutations over a lineage tree.

**SHM targeting models**

Computational models and analyses of SHM have separated the process into two independent components:

1. A mutability model that defines where mutations occur.
2. A nucleotide substitution model that defines the resulting mutation.

Collectively these are what form the targeting model of SHM. shazam provides empirically derived targeting models for both humans and mice, along with tools to build these mutability and substitution models from data.
• createTargetingModel: Build a 5-mer targeting model.
• plotMutability: Plot 5-mer mutability rates.
• HH_S5F: Human 5-mer SHM targeting model.
• MK_RS5NF: Mouse 5-mer SHM targeting model.

Quantification of selection pressure

Bayesian Estimation of Antigen-driven Selection in Ig Sequences is a novel method for quantifying antigen-driven selection in high-throughput Ig sequence data. Targeting models created using shazam can be used to estimate the null distribution of expected mutation frequencies used by BASELINe, providing measures of selection pressure informed by known AID targeting biases.

• calcBaseline: Calculate the BASELINe probability density functions (PDFs).
• groupBaseline: Combine PDFs from sequences grouped by biological or experimental relevance.
• summarizeBaseline: Compute summary statistics from BASELINe PDFs.
• testBaseline: Perform significance testing for the difference between BASELINe PDFs.
• plotBaselineDensity: Plot the probability density functions resulting from selection analysis.
• plotBaselineSummary: Plot summary statistics resulting from selection analysis.

Mutational distance calculation

shazam provides tools to compute evolutionary distances between sequences or groups of sequences, which can leverage SHM targeting models. This information is particularly useful in understanding and defining clonal relationships.

• findThreshold: Identify clonal assignment threshold based on distances to nearest neighbors.
• distToNearest: Tune clonal assignment thresholds by calculating distances to nearest neighbors.
• calcTargetingDistance: Construct a nucleotide distance matrix from a 5-mer targeting model.

References

shmulateSeq  

Simulate mutations in a single sequence

Description

Generates random mutations in a sequence iteratively using a targeting model. Targeting probabilities at each position are updated after each iteration.

Usage

shmulateSeq(
  sequence, 
  numMutations, 
  targetingModel = HH_S5F, 
  start = 1, 
  end = nchar(sequence) 
)

Arguments

sequence  
sequence string in which mutations are to be introduced. Accepted alphabet: {A,T,G,C,N,.}. Note that - is not accepted.

numMutations  
number of mutations to be introduced into sequence.

targetingModel  
5-mer TargetingModel object to be used for computing probabilities of mutations at each position. Defaults to HH_S5F.

start  
Initial position in sequence where mutations can be introduced. Default: 1

end  
Last position in sequence where mutations can be introduced. Default: last position (sequence length).

Details

If the input sequence has a non-triplet overhang at the end, it will be trimmed to the last codon. For example, ATGCATGC will be trimmed to ATGCAT.

Mutations are not introduced to positions in the input sequence that contain . or N.

Value

A string defining the mutated sequence.

See Also

See shmulateTree for imposing mutations on a lineage tree. See HH_S5F and MK_RS5NF for predefined TargetingModel objects.
Examples

```r
# Define example input sequence
sequence <- "NGATCTGACGACGCGGCTATTACTGTGCGAGATA.TTTA"

# Simulate using the default human 5-mer targeting model
shmulateSeq(sequence, numMutations=6)
```

Description

`shmulateTree` returns a set of simulated sequences generated from an input sequence and a lineage tree. The input sequence is used to replace the most recent common ancestor (MRCA) node of the igraph object defining the lineage tree. Sequences are then simulated with mutations corresponding to edge weights in the tree. Sequences will not be generated for groups of nodes that are specified to be excluded.

Usage

```r
shmulateTree(
  sequence,
  graph,
  targetingModel = HH_S5F,
  field = NULL,
  exclude = NULL,
  junctionWeight = NULL,
  start = 1,
  end = nchar(sequence)
)
```

Arguments

- `sequence`: string defining the MRCA sequence to seed mutations from.
- `graph`: igraph object defining the seed lineage tree, with vertex annotations, whose edges are to be recreated.
- `targetingModel`: 5-mer `TargetingModel` object to be used for computing probabilities of mutations at each position. Defaults to `HH_S5F`.
- `field`: annotation to use for both unweighted path length exclusion and consideration as the MRCA node. If `NULL` do not exclude any nodes.
- `exclude`: vector of annotation values in `field` to exclude from potential MRCA set. If `NULL` do not exclude any nodes. Has no effect if `field=NULL`.
- `junctionWeight`: fraction of the nucleotide sequence that is within the junction region. When specified this adds a proportional number of mutations to the immediate offspring nodes of the MRCA. Requires a value between 0 and 1. If `NULL` then edge weights are unmodified from the input graph.
slideWindowDb

Description

slideWindowDb determines whether each input sequence in a data.frame contains equal to or more than a given number of mutations in a given length of consecutive nucleotides (a "window") when compared to their respective germline sequence.
Usage

```r
slideWindowDb(
  db,
  sequenceColumn = "SEQUENCE_IMGT",
  germlineColumn = "GERMLINE_IMGT_D_MASK",
  mutThresh, 
  windowSize
)
```

Arguments

- `db` data.frame containing sequence data.
- `sequenceColumn` name of the column containing IMGT-gapped sample sequences.
- `germlineColumn` name of the column containing IMGT-gapped germline sequences.
- `mutThresh` threshold on the number of mutations in `windowSize` consecutive nucleotides. Must be between 1 and `windowSize` inclusive.
- `windowSize` length of consecutive nucleotides. Must be at least 2.

Value

a logical vector. The length of the vector matches the number of input sequences in `db`. Each entry in the vector indicates whether the corresponding input sequence should be filtered based on the given parameters.

See Also

See `slideWindowSeq` for applying the sliding window approach on a single sequence. See `slideWindowTune` for parameter tuning for `mutThresh` and `windowSize`.

Examples

```r
# Use an entry in the example data for input and germline sequence
data(ExampleDb, package="alakazam")

# Apply the sliding window approach on a subset of ExampleDb
slideWindowDb(db = ExampleDb[1:10,], mutThresh=6, windowSize=10)
```

---

**slideWindowSeq**  
 **Sliding window approach towards filtering a single sequence**

Description

`slideWindowSeq` determines whether an input sequence contains equal to or more than a given number of mutations in a given length of consecutive nucleotides (a "window") when compared to a germline sequence.
Usage

`slideWindowSeq(inputSeq, germlineSeq, mutThresh, windowSize)`

Arguments

- `inputSeq`: input sequence.
- `germlineSeq`: germline sequence.
- `mutThresh`: threshold on the number of mutations in windowSize consecutive nucleotides. Must be between 1 and windowSize inclusive.
- `windowSize`: length of consecutive nucleotides. Must be at least 2.

Value

TRUE if there are equal to or more than mutThresh number of mutations in any window of windowSize consecutive nucleotides (i.e. the sequence should be filtered); FALSE if otherwise.

See Also

- `calcObservedMutations` is called by `slideWindowSeq` to identify observed mutations. See `slideWindowDb` for applying the sliding window approach on a `data.frame`. See `slideWindowTune` for parameter tuning for `mutThresh` and `windowSize`.

Examples

```r
# Use an entry in the example data for input and germline sequence
data(ExampleDb, package="alakazam")
in_seq <- ExampleDb[100, "SEQUENCE.IMGT"]
germ_seq <- ExampleDb[100, "GERMLINE.IMGT_D_MASK"]

# Determine if in_seq has 6 or more mutations in 10 consecutive nucleotides
slideWindowSeq(inputSeq=in_seq, germlineSeq=germ_seq, mutThresh=6, windowSize=10)
```

Description

Apply `slideWindowDb` over a search grid made of combinations of `mutThresh` and `windowSize` to help with picking a pair of values for these parameters. Parameter tuning can be performed by choosing a combination that gives a reasonable number of filtered/remaining sequences.
Usage

slideWindowTune(
  db,
  sequenceColumn = "SEQUENCE_IMGT",
  germlineColumn = "GERMLINE_IMGT_D_MASK",
  dbMutList = NULL,
  mutThreshRange,
  windowSizeRange,
  verbose = TRUE
)

Arguments

db data.frame containing sequence data.
sequenceColumn name of the column containing IMGT-gapped sample sequences.
germlineColumn name of the column containing IMGT-gapped germline sequences.
dbMutList if supplied, this should be a list consisting of data.frames returned as $pos in the nested list produced by calcObservedMutations with returnRaw=TRUE; otherwise, calcObservedMutations is called on columns sequenceColumn and germlineColumn of db. Default is NULL.
mutThreshRange range of threshold on the number of mutations in windowSize consecutive nucleotides to try. Must be between 1 and maximum windowSizeRange inclusive.
windowSizeRange range of length of consecutive nucleotides to try. The lower end must be at least 2.
verbose whether to print out messages indicating current progress. Default is TRUE.

Details

If, in a given combination of mutThresh and windowSize, mutThresh is greater than windowSize, NAs will be returned for that particular combination. A message indicating that the combination has been "skipped" will be printed if verbose=TRUE.

If calcObservedMutations was previously run on db and saved, supplying $pos from the saved result as dbMutList could save time by skipping a second call of calcObservedMutations. This could be helpful especially when db is large.

Value

a list of logical matrices. Each matrix corresponds to a windowSize in windowSizeRange. Each column in a matrix corresponds to a mutThresh in mutThreshRange.

See Also

slideWindowDb is called on db for tuning. See slideWindowTunePlot for visualization. See calcObservedMutations for generating dbMutList.
Examples

```r
# Load and subset example data
data(ExampleDb, package="alakazam")
db <- ExampleDb[1:5, ]

# Try out thresholds of 2-4 mutations in window sizes of 7-9 nucleotides.
# In this case, all combinations are legal.
slideWindowTune(db, mutThreshRange=2:4, windowSizeRange=7:9)

# Illegal combinations are skipped, returning NAs.
slideWindowTune(db, mutThreshRange=2:4, windowSizeRange=2:4,
                verbose=FALSE)

# Run calcObservedMutations separately
exDbMutList <- sapply(1:5, function(i) {
    calcObservedMutations(inputSeq=db[i, "SEQUENCE_IMGT"],
                           germlineSeq=db[i, "GERMLINE_IMGT_D_MASK"],
                           returnRaw=TRUE)$pos
})
slideWindowTune(db, dbMutList=exDbMutList,
                mutThreshRange=2:4, windowSizeRange=2:4)
```

slideWindowTunePlot

Visualize parameter tuning for sliding window approach

Description

Visualize results from slideWindowTune

Usage

```r
slideWindowTunePlot(
    tuneList,
    plotFiltered = TRUE,
    percentage = FALSE,
    jitter.x = FALSE,
    jitter.x.amt = 0.1,
    jitter.y = FALSE,
    jitter.y.amt = 0.1,
    pchs = 1,
    ltys = 2,
    cols = 1,
    plotLegend = TRUE,
    legendPos = "topright",
    legendHoriz = FALSE,
    legendCex = 1,
    title = NULL
)
```
**Arguments**

- **tuneList**: a list of logical matrices returned by `slideWindowTune`.
- **plotFiltered**: whether to plot the number of filtered sequences (as opposed to the number of remaining sequences). Default is TRUE.
- **percentage**: whether to plot on the y-axis the percentage of filtered sequences (as opposed to the absolute number). Default is FALSE.
- **jitter.x**: whether to jitter x-axis values. Default is FALSE.
- **jitter.x.amt**: amount of jittering to be applied on x-axis values if `jitter.x=TRUE`. Default is 0.1.
- **jitter.y**: whether to jitter y-axis values. Default is FALSE.
- **jitter.y.amt**: amount of jittering to be applied on y-axis values if `jitter.y=TRUE`. Default is 0.1.
- **pchs**: point types to pass on to `plot`.
- **ltys**: line types to pass on to `plot`.
- **cols**: colors to pass on to `plot`.
- **plotLegend**: whether to plot legend. Default is TRUE.
- **legendPos**: position of legend to pass on to `legend`. Can be either a numeric vector specifying x-y coordinates, or one of "topright", "center", etc. Default is "topright".
- **legendHoriz**: whether to make legend horizontal. Default is FALSE.
- **legendCex**: numeric values by which legend should be magnified relative to 1.
- **title**: plot main title. Default is NULL (no title)

**Details**

For each `WindowSize`, the numbers of sequences filtered or remaining after applying the sliding window approach are plotted on the y-axis against thresholds on the number of mutations in a window on the x-axis.

When plotting, a user-defined amount of jittering can be applied on values plotted on either axis or both axes via adjusting `jitter.x`, `jitter.y`, `jitter.x.amt` and `jitter.y.amt`. This may be helpful with visually distinguishing lines for different window sizes in case they are very close or identical to each other. If plotting percentages (`percentage=TRUE`) and using jittering on the y-axis values (`jitter.y=TRUE`), it is strongly recommended that `jitter.y.amt` be set very small (e.g. 0.01).

NA for a combination of `mutThresh` and `WindowSize` where `mutThresh` is greater than `WindowSize` will not be plotted.

**See Also**

See `slideWindowTune` for how to get `tuneList`. See `jitter` for use of amount of jittering.
### Examples

```r
# Use an entry in the example data for input and germline sequence
data(ExampleDb, package="alakazam")

# Try out thresholds of 2-4 mutations in window sizes of 3-5 nucleotides
# on a subset of ExampleDb
tuneList <- slideWindowTune(db = ExampleDb[1:10, ],
                           mutThreshRange = 2:4, windowSizeRange = 3:5,
                           verbose = FALSE)

# Visualize
# Plot numbers of sequences filtered without jittering y-axis values
slideWindowTunePlot(tuneList, pchs=1:3, lty=1:3, col=1:3,
                    plotFiltered=TRUE, jitter.y=FALSE)

# Notice that some of the lines overlap
# Jittering could help
slideWindowTunePlot(tuneList, pchs=1:3, lty=1:3, col=1:3,
                    plotFiltered=TRUE, jitter.y=TRUE)

# Plot numbers of sequences remaining instead of filtered
slideWindowTunePlot(tuneList, pchs=1:3, lty=1:3, col=1:3,
                    plotFiltered=FALSE, jitter.y=TRUE,
                    legendPos="bottomright")

# Plot percentages of sequences filtered with a tiny amount of jittering
slideWindowTunePlot(tuneList, pchs=1:3, lty=1:3, col=1:3,
                    plotFiltered=TRUE, percentage=TRUE,
                    jitter.y=TRUE, jitter.y.amt=0.01)
```

### summarizeBaseline

`summarizeBaseline` calculates BASELINe summary statistics such as the mean selection strength (mean Sigma), the 95% confidence intervals and p-values for the presence of selection.

#### Description

`summarizeBaseline` calculates BASELINe statistics such as the mean selection strength (mean Sigma), the 95% confidence intervals and p-values for the presence of selection.

#### Usage

```r
summarizeBaseline(baseline, returnType = c("baseline", "df"), nproc = 1)
```

#### Arguments

- **baseline**: Baseline object returned by `calcBaseline` containing annotations and BASELINe posterior probability density functions (PDFs) for each sequence.
summarizeBaseline

**returnType**
One of c("baseline", "df") defining whether to return a Baseline object ("baseline") with an updated stats slot or a data.frame ("df") of summary statistics.

**nproc**
number of cores to distribute the operation over. If nproc = 0 then the cluster has already been set and will not be reset.

**Details**
The returned p-value can be either positive or negative. Its magnitude (without the sign) should be interpreted as per normal. Its sign indicates the direction of the selection detected. A positive p-value indicates positive selection, whereas a negative p-value indicates negative selection.

**Value**
Either a modified Baseline object or data.frame containing the mean BASELINe selection strength, its 95% confidence intervals, and a p-value for the presence of selection.

**References**

**See Also**
See calcBaseline for generating Baseline objects and groupBaseline for convolving groups of BASELINe PDFs.

**Examples**

```r
# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE == "IgG")

# Collapse clones
db <- collapseClones(db, sequenceColumn="SEQUENCE_IMGT",
                     germlineColumn="GERMLINE_IMGT_D_MASK",
                     method="thresholdedFreq", minimumFrequency=0.6,
                     includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

# Calculate BASELINe
baseline <- calcBaseline(db,
                         sequenceColumn="CLONAL_SEQUENCE",
                         germlineColumn="CLONAL_GERMLINE",
                         testStatistic="focused",
                         regionDefinition=IMGT_V,
                         targetingModel=HH_S5F,
                         nproc = 1)

# Grouping the PDFs by the sample annotation
grouped <- groupBaseline(baseline, groupBy="SAMPLE")
```
# Get a data.frame of the summary statistics
stats <- summarizeBaseline(grouped, returnType="df")

TargetingModel-class  
S4 class defining a targeting model

Description

TargetingModel defines a common data structure for mutability, substitution and targeting of immunoglobulin (Ig) sequencing data in a 5-mer microsequence context.

Usage

```r
## S4 method for signature 'TargetingModel,missing'
plot(x, y, ...)
```

Arguments

- `x` TargetingModel object.
- `y` ignored.
- `...` arguments to pass to `plotMutability`.

Slots

- `name` Name of the model.
- `description` Description of the model and its source data.
- `species` Genus and species of the source sequencing data.
- `date` Date the model was built.
- `citation` Publication source.
- `substitution` Normalized rates of the center nucleotide of a given 5-mer mutating to a different nucleotide. The substitution model is stored as a 5x3125 matrix of rates. Rows define the mutated nucleotide at the center of each 5-mer, one of c("A", "C", "G", "T", "N"), and columns define the complete 5-mer of the unmutated nucleotide sequence.
- `mutability` Normalized rates of a given 5-mer being mutated. The mutability model is stored as a numeric vector of length 3125 with mutability rates for each 5-mer. Note that "normalized" means that the mutability rates for the 1024 5-mers that contain no "N" at any position sums up to 1 (as opposed to the entire vector summing up to 1).
- `targeting` Rate matrix of a given mutation occurring, defined as `mutability * substitution`. The targeting model is stored as a 5x3125 matrix. Rows define the mutated nucleotide at the center of each 5-mer, one of c("A", "C", "G", "T", "N"), and columns define the complete 5-mer of the unmutated nucleotide sequence.

See Also

See `createTargetingModel` building models from sequencing data.
testBaseline  

Two-sided test of BASELINe PDFs

Description

TestBaseline performs a two-sample significance test of BASELINe posterior probability density functions (PDFs).

Usage

```r
testBaseline(baseline, groupBy)
```

Arguments

- `baseline`: Baseline object containing the db and grouped BASELINe PDFs returned by `groupBaseline`.
- `groupBy`: string defining the column in the db slot of the Baseline containing sequence or group identifiers.

Value

A data.frame with test results containing the following columns:

- `REGION`: sequence region, such as "CDR" and "FWR".
- `TEST`: string defining the groups be compared. The string is formatted as the conclusion associated with the p-value in the form GROUP1 != GROUP2. Meaning, the p-value for rejection of the null hypothesis that GROUP1 and GROUP2 have equivalent distributions.
- `PVALUE`: two-sided p-value for the comparison.
- `FDR`: FDR corrected PVALUE.

References


See Also

To generate the Baseline input object see `groupBaseline`.

Examples

```r
# Subset example data
data(ExampleDb, package="alakazam")
db <- subset(ExampleDb, ISOTYPE %in% c("IgM", "IgG", "IgA"))

# Collapse clones
```
U5N <- collapseClones(db, sequenceColumn="SEQUENCE_IMGT",
                          germlineColumn="GERMLINE_IMGT_D_MASK",
                          method="thresholdedFreq", minimumFrequency=0.6,
                          includeAmbiguous=FALSE, breakTiesStochastic=FALSE)

# Calculate Baseline
baseline <- calcBaseline(db,
                          sequenceColumn="CLONAL_SEQUENCE",
                          germlineColumn="CLONAL_GERMLINE",
                          testStatistic="focused",
                          regionDefinition=IMGT_V,
                          targetingModel=HH_S5F,
                          nproc=1)

# Group PDFs by the isotype
grouped <- groupBaseline(baseline, groupBy="ISOTYPE")

# Visualize isotype PDFs
plot(grouped, "ISOTYPE")

# Perform test on isotype PDFs
testBaseline(grouped, groupBy="ISOTYPE")

U5N

Uniform 5-mer null targeting model.

Description

A null 5-mer model of somatic hypermutation targeting where all substitution, mutability and targeting rates are uniformly distributed.

Usage

U5N

Format

A TargetingModel object.

See Also

See HH_S5F and HKL_S5F for the human 5-mer targeting models; and MK_RS5NF for the mouse 5-mer targeting model.
writeTargetingDistance

Write targeting model distances to a file

Description

writeTargetingDistance writes a 5-mer targeting distance matrix to a tab-delimited file.

Usage

writeTargetingDistance(model, file)

Arguments

model  
TargetingModel object with mutation likelihood information.

file  
name of file to write.

Details

The targeting distance write as a tab-delimited 5x3125 matrix. Rows define the mutated nucleotide at the center of each 5-mer, one of c("A","C","G","T","N"), and columns define the complete 5-mer of the unmutated nucleotide sequence. NA values in the distance matrix are replaced with distance 0.

See Also

Takes as input a TargetingModel object and calculates distances using calcTargetingDistance.

Examples

```r
# Not run:
# Write HS5F targeting model to working directory as hs5f.tab
writeTargetingDistance(HH_S5F, "hs5f.tab")

# End(Not run)
```
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