

Package ‘MHCtools’

February 14, 2026

Type Package

Title Analysis of MHC Data in Non-Model Species

Version 1.6.0

Description Sixteen tools for bioinformatics processing and analysis of major histocompatibility complex (MHC) data. The functions are tailored for amplicon data sets that have been filtered using the dada2 method (for more information on dada2, visit <<https://benjjneb.github.io/dada2/>>), but even other types of data sets can be analyzed.

The ReplMatch() function matches replicates in data sets in order to evaluate genotyping success.

The GetReplTable() and GetReplStats() functions perform such an evaluation.

The CreateFas() function creates a fasta file with all the sequences in the data set.

The CreateSamplesFas() function creates individual fasta files for each sample in the data set.

The DistCalc() function calculates Grantham, Sandberg, or p-distances from pairwise comparisons of all sequences in a data set, and mean distances of all pairwise comparisons within each sample in a data set. The function additionally outputs five tables with physico-chemical z-descriptor values (based on Sandberg et al. 1998) for each amino acid position in all sequences in the data set. These tables may be useful for further downstream analyses, such as estimation of MHC supertypes.

The BootKmeans() function is a wrapper for the kmeans() function of the 'stats' package, which allows for bootstrapping. Bootstrapping k-estimates may be desirable in data sets, where e.g. BIC- vs. k-values do not produce clear inflection points ("elbows"). BootKmeans() performs multiple runs of kmeans() and estimates optimal k-values based on a user-defined threshold of BIC reduction. The method is an automated and bootstrapped version of visually inspecting elbow plots of BIC- vs. k-values.

The ClusterMatch() function is a tool for evaluating whether different k-means() clustering models identify similar clusters, and summarize bootstrap model stats as means for different estimated values of k. It is designed to take files produced by the BootKmeans() function as input, but other data can be analyzed if the descriptions of the required data formats are observed carefully.

The SynDist() function analyses of synonymous variation among aligned protein-coding DNA sequences, that is, nucleotide substitutions that do not translate to changes in

the amino acid sequences due to degeneracy of the genetic code. The SynDist() function calculates synonymous nucleotide changes per base and per codon in pairwise sequence comparisons, as well as mean synonymous variation among all pairwise comparisons of the sequences within each sample in a data set.

The PapaDiv() function compares parent pairs in the data set and calculate their joint MHC diversity, taking into account sequence variants that occur in both parents.

The HpltFind() function infers putative haplotypes from families in the data set.

The GetHpltTable() and GetHpltStats() functions evaluate the accuracy of the haplotype inference.

The CreateHpltOccTable() function creates a binary (logical) haplotype-sequence occurrence matrix from the output of HpltFind(), for easy overview of which sequences are present in which haplotypes.

The HpltMatch() function compares haplotypes to help identify overlapping and potentially identical types.

The NestTablesXL() function translates the output from HpltFind() to an Excel workbook, that provides a convenient overview for evaluation and curating of the inferred putative haplotypes.

License MIT + file LICENSE

Encoding UTF-8

LazyData true

Imports stats, utils, mgcv, grDevices, graphics, openxlsx

RoxygenNote 7.3.3

NeedsCompilation no

Author Jacob Roved [aut, cre]

Maintainer Jacob Roved <jacob.roved@biol.lu.se>

Depends R (>= 3.5.0)

Repository CRAN

Date/Publication 2026-02-14 17:20:02 UTC

Contents

BootKmeans	3
ClusterMatch	5
CreateFas	7
CreateHpltOccTable	8
CreateSamplesFas	9
DistCalc	10
GetHpltStats	12
GetHpltTable	13
GetReplStats	14
GetReplTable	15
HpltFind	16

HpltMatch 17
 hplt_occurrence_matrix 18
 k_summary_table 19
 NestTablesXL 19
 nest_table 21
 PapaDiv 21
 parents_table 22
 replicates_table 23
 ReplMatch 24
 sequence_table 25
 sequence_table_fas 25
 sequence_table_hplt 26
 sequence_table_repl 26
 sequence_table_SynDist 27
 SynDist 27
 z1_matrix 29
 z2_matrix 30
 z3_matrix 30
 z4_matrix 31
 z5_matrix 31

Index **33**

BootKmeans *BootKmeans() function*

Description

[BootKmeans](#) is a wrapper for the `kmeans()` function of the 'stats' package, which allows for bootstrapping. Bootstrapping k-estimates may be desirable in data sets, where the BIC- vs. k-values do not produce clear inflection points ("elbows").

Usage

```

BootKmeans(
  z1_matrix,
  z2_matrix,
  z3_matrix,
  z4_matrix,
  z5_matrix,
  threshold = 0.01,
  no_scans = 1000,
  max_k = 40,
  iter.max = 1e+06,
  nstart = 200,
  algorithm = "Hartigan-Wong",
  path_out = path_out
)
    
```

Arguments

<code>z1_matrix</code>	a matrix with numerical values of the first z-descriptor for each amino acid position in all sequences in the data set.
<code>z2_matrix</code>	a matrix with numerical values of the second z-descriptor for each amino acid position in all sequences in the data set.
<code>z3_matrix</code>	a matrix with numerical values of the third z-descriptor for each amino acid position in all sequences in the data set.
<code>z4_matrix</code>	a matrix with numerical values of the fourth z-descriptor for each amino acid position in all sequences in the data set.
<code>z5_matrix</code>	a matrix with numerical values of the fifth z-descriptor for each amino acid position in all sequences in the data set.
<code>threshold</code>	a numerical value between 0 and 1 specifying the threshold of reduction in BIC for selecting a k estimate for each kmeans clustering model. The value specifies a proportion of the max observed reduction in BIC when increasing k by 1 (default 0.01).
<code>no_scans</code>	an integer specifying the number of k estimation scans to run (default 1,000).
<code>max_k</code>	an integer specifying the hypothetical maximum number of clusters to detect (default 40). In each k estimation scan, the algorithm runs a kmeans() clustering model for each value of k between 1 and max_k.
<code>iter.max</code>	an integer specifying the maximum number of iterations allowed in each kmeans() clustering model (default 1,000,000).
<code>nstart</code>	an integer specifying the number of rows in the set of input matrices that will be chosen as initial centers in the kmeans() clustering models (default 200).
<code>algorithm</code>	character vector, specifying the method for the kmeans() clustering function, one of c("Hartigan-Wong", "Lloyd", "Forgy", "MacQueen"), default is "Hartigan-Wong".
<code>path_out</code>	a user defined path to the folder where the output files will be saved.

Details

BootKmeans() performs multiple runs of kmeans() scanning k-values from 1 to a maximum value defined by the user. In each scan, an optimal k-value is estimated using a user-defined threshold of BIC reduction. The method is an automated version of visually inspecting elbow plots of BIC- vs. k-values. The number of scans to be performed is defined by the user.

For each k-estimate scan, the algorithm produces a summary of the stats incl. total within SS, AIC, and BIC, an elbow plot (BIC vs. k), and a set of cluster files corresponding to the estimated optimal k-value. It also produces a table summarizing the stats of the final selected kmeans() models corresponding to the estimated optimal k-values of each scan.

After running BootKmeans() on a data set, it is recommended to subsequently evaluate the repeatability of the bootstrapped k-estimation scans with the ClusterMatch() function also included in MHCtools.

Input data format: A set of five z-matrices containing numerical values of the z-descriptors (z1-z5) for each amino acid position in a sequence alignment. Each column should represent an amino acid position and each row one sequence in the alignment.

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

The function produces three folders in `path_out`, which contain for each scan the estimated k-clusters saved as `.RData` files, an elbow plot saved as `.pdf`, and a stats summary table saved as a `.csv` file. In `path_out` a summary of all scans performed in the bootstrap run is also saved as `.csv`. This table is also shown in the console. Should alternative elbow plots be desired, they may be produced manually with the stats presented in the summary tables for each scan.

Note

If z-matrices were generated with the `DistCalc()` function, please make sure to load the z-matrices from the `.csv` files exported by `DistCalc()`. Calling e.g. `'z1_matrix'` without loading the exported tables will engage the default test data set in MHCtools.

Setting `max_k` too high can cause kmeans to fail with the error "more cluster centers than distinct data points" - this problem can be solved by reducing `max_k`.

AIC and BIC are calculated from the kmeans model objects by the following formulae: - $AIC = D + 2 * m * k$ - $BIC = D + \log(n) * m * k$ in which: - $m = \text{ncol}(\text{fit}\$centers)$ - $n = \text{length}(\text{fit}\$cluster)$ - $k = \text{nrow}(\text{fit}\$centers)$ - $D = \text{fit}\$tot.withinss$

See Also

[ClusterMatch](#); [DistCalc](#)

Examples

```
z1_matrix <- z1_matrix
z2_matrix <- z2_matrix
z3_matrix <- z3_matrix
z4_matrix <- z4_matrix
z5_matrix <- z5_matrix
path_out <- tempdir()
BootKmeans(z1_matrix, z2_matrix, z3_matrix, z4_matrix, z5_matrix, threshold=0.01,
no_scans=10, max_k=20, iter.max=10, nstart=10, algorithm="Hartigan-Wong",
path_out=path_out)
```

Description

`ClusterMatch` is a tool for evaluating whether `k-means()` clustering models with similar estimated values of `k` identify similar clusters. `ClusterMatch()` also summarizes model stats as means for different estimated values of `k`. It is designed to take files produced by the `BootKmeans()` function as input, but other data can be analyzed if the descriptions of the data formats given below are observed carefully.

Usage

```
ClusterMatch(filepath, path_out, k_summary_table)
```

Arguments

<code>filepath</code>	a user defined path to a folder that contains the set of K-cluster files to be matched against each other. The algorithm will attempt to load all files in the folder, so it should contain only the relevant K-cluster files. If the clusters were generated using the <code>BootKmeans()</code> function, such a folder (named Clusters) was created by the algorithm in the output path given by the user. Each K-cluster file should correspond to the <code>model\$cluster</code> object in <code>kmeans()</code> saved as a <code>.RData</code> file. Such files are generated as part of the output from <code>BootKmeans()</code> . <code>ClusterMatch()</code> assumes that the file names contain the string "model_" followed by a model number, which must match the corresponding row numbers in <code>k_summary_table</code> . If the data used was generated with the <code>BootKmeans()</code> function, the formats and numbers will match by default.
<code>path_out</code>	a user defined path to the folder where the output files will be saved.
<code>k_summary_table</code>	a data frame summarizing the stats of the <code>kmeans()</code> models that produced the clusters in the K-cluster files. If the data used was generated with the <code>BootKmeans()</code> function, a compatible <code>k_summary_table</code> was produced in the output path with the file name "k_means_bootstrap_summary_stats_<date>.csv". If other data is analyzed, please observe these formatting requirements: The <code>k_summary_table</code> must contain the data for each <code>kmeans()</code> model in rows and as minimum the following columns: - <code>k</code> -value (colname: <code>k.est</code>) - residual total within sums-of-squares (colname: <code>Tot.withinss.resid</code>) - residual AIC (colname: <code>AIC.resid</code>) - residual BIC (colname: <code>BIC.resid</code>) - delta BIC/max BIC (colname: <code>prop.delta.BIC</code>) - delta BIC/ <code>k.est</code> (colname: <code>delta.BIC.over.k</code>) It is crucial that the models have the same numbers in the K-cluster file names and in the <code>k_summary_table</code> , and that the rows of the table are ordered by the model number.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), *HLA Typing: Methods and Protocols* (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

The function returns a summary table, which for each estimated number of clusters (i.e. the k -values of the models) lists: - number of models that found i clusters - mean residual total within sums-of-squares - mean residual AIC - mean residual BIC - mean delta BIC/max BIC - mean delta BIC/ k - mean number of allele assignments that fall outside of the i most abundant clusters across all pairwise comparisons between the models that found i clusters - mean proportion of allele assignments that fall outside of the i most abundant clusters across all pairwise comparisons between the models that found i clusters The summary table is also saved as a .csv file in the output path.

See Also

[BootKmeans](#)

Examples

```
filepath <- system.file("extdata/ClusterMatch", package="MHCtools")
path_out <- tempdir()
k_summary_table <- k_summary_table
ClusterMatch(filepath, path_out, k_summary_table)
```

CreateFas

CreateFas() function

Description

[CreateFas](#) creates a FASTA file with all the sequences in a 'dada2' sequence table.

Usage

```
CreateFas(seq_table, path_out)
```

Arguments

seq_table	seq_table is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns.
path_out	is a user defined path to the folder where the output files will be saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A FASTA file with all the sequences in a 'dada2' sequence table. The sequences are named in the FASTA file by an index number corresponding to their column number in the sequence table.

See Also

[CreateSamplesFas](https://benjjneb.github.io/dada2/); for more information about 'dada2' visit <<https://benjjneb.github.io/dada2/>>

Examples

```
seq_table <- sequence_table_fas
path_out <- tempdir()
CreateFas(seq_table, path_out)
```

CreateHpltOccTable *CreateHpltOccTable() function*

Description

[CreateHpltOccTable](#) is designed to create a haplotype-sequence occurrence matrix from the set of R lists with putative haplotypes output by the `HpltFind()` function. `CreateHpltOccTable()` assumes that data originated from a diploid species.

Usage

```
CreateHpltOccTable(seq_table, filepath, path_out)
```

Arguments

<code>seq_table</code>	<code>seq_table</code> is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns.
<code>filepath</code>	is a user defined path to the folder where the output files from the <code>HpltFind()</code> function have been saved.
<code>path_out</code>	is a user defined path to the folder where the output files will be saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A binary (logical) occurrence matrix with the data set sequences (inherited from `seq_table`) in columns and the putative haplotypes inferred by the `HpltFind()` function in rows.

See Also

[HpltFind](https://benjjneb.github.io/dada2/); for more information about 'dada2' visit <<https://benjjneb.github.io/dada2/>>

Examples

```
seq_table <- sequence_table
filepath <- system.file("extdata/HpltFindOut/", package="MHCtools")
path_out <- tempdir()
CreateHpltOccTable(seq_table, filepath, path_out)
```

CreateSamplesFas *CreateSamplesFas()* function

Description

[CreateSamplesFas](#) creates a set of FASTA files with the sequences present in each sample in a 'dada2' sequence table.

Usage

```
CreateSamplesFas(seq_table, path_out)
```

Arguments

seq_table	seq_table is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns.
path_out	is a user defined path to the folder where the output files will be saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A set of FASTA files with the sequences present in each sample in the sequence table. The sequences are named in the FASTA files by an index number corresponding to their column number in the sequence table, thus identical sequences will have identical sample names in all the FASTA files.

See Also

[CreateFas](#); for more information about 'dada2' visit <<https://benjjneb.github.io/dada2/>>

Examples

```
seq_table <- sequence_table_fas
path_out <- tempdir()
CreateSamplesFas(seq_table, path_out)
```

DistCalc

DistCalc() function

Description

[DistCalc](#) calculates Grantham distances, Sandberg distances, or p-distances from pairwise comparisons of aligned sequences.

Usage

```
DistCalc(
  seq_file,
  path_out,
  input_fasta = NULL,
  input_seq = "aa",
  aa_dist = NULL,
  codon_pos = NULL,
  dist_type = "G"
)
```

Arguments

seq_file	is a sequence occurrence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns. Optionally, a fasta file can be supplied as input in the format rendered by read.fasta() from the package 'seqinr'.
path_out	is a user defined path to the folder where the output files will be saved.
input_fasta	optional, a logical (TRUE/FALSE) that indicates whether the input file is a fasta file (TRUE) or a 'dada2'-style sequence table (NULL/FALSE). The default is NULL/FALSE.
input_seq	defines the type of sequences in seq_file. It may take the values 'nucl' or 'aa'.
aa_dist	is optional, a logical (TRUE/FALSE) that determines whether nucleotide sequences should be translated to amino acid sequences before distance calculation, default is NULL/FALSE. Note that aa_dist must be set to TRUE, if Grantham or Sandberg distances are calculated from an alignment of nucleotide sequences.
codon_pos	is optional, a vector of comma separated integers specifying which codons to include in distance calculations. If omitted, distance calculations are made using all codons. Note: When calculating nucleotide P-distances, codon_pos should be specified as a vector of nucleotide positions.

`dist_type` is used to specify which kind of distances that are calculated. It takes the values 'G' for Grantham distances, 'S' for Sandberg distances, or 'P' for p-distances. The argument is optional with 'G' as default setting.

Details

The `DistCalc()` function takes a fasta file or a 'dada2'-style sequence occurrence table (with aligned sequences as column names and samples in rows) as input and produces a matrix with pairwise distances for all sequences in the data set. If calculation of Sandberg distances is specified, the function additionally outputs five tables with physico-chemical z-descriptor values (based on Sandberg et al. 1998) for each amino acid position in all sequences in the data set. These tables may be useful for further downstream analyses, such as estimation of MHC supertypes. If a sequence occurrence table is provided as input, the `DistCalc()` function furthermore produces a table with the mean distances from all pairwise comparisons of the sequences in each sample in the data set. (Note: The mean distance will be NA for samples that have 0 or 1 sequence(s).)

Grantham distances and Sandberg distances are calculated as described in Pierini & Lenz 2018. The Grantham distances produced by `DistCalc()` are simply the mean Grantham distances (Grantham 1974) between all amino acid codons in sequence pairs. When calculating Sandberg distances, `DistCalc()` first computes Euclidian distances between all amino acid pairs based on the five physico-chemical z-descriptors defined in Sandberg et al. 1998. Sandberg distances are then calculated as the mean Euclidian distances between all amino acid codons in sequence pairs. P-distances calculated by `DistCalc()` are simply the proportion of varying codons between pairs of sequences.

The `DistCalc()` function includes an option for the user to specify which codons to compare, which is useful e.g. if conducting the analysis only on codons involved in specific functions, such as peptide binding of an MHC molecule. Note: When calculating nucleotide P-distances, `codon_pos` is applied directly on the nucleotide sequences. This allows the user to calculate divergence in e.g. first, second, or third codon positions. Hence, `codon_pos` should be specified as a vector of nucleotide positions when calculating nucleotide P-distances.

`DistCalc()` also accepts calculating amino acid distances directly from protein-coding DNA sequences using the standard genetic code.

The `DistCalc()` function accepts the following characters in the sequences: Nucleotide sequences: A,T,G,C Amino acid sequences: A,R,N,D,C,Q,E,G,H,I,L,K,M,F,P,S,T,W,Y,V

It accepts gaps defined by '-'. Nucleotide triplets containing gaps are translated to 'X', if amino acid distances are calculated directly from DNA nucleotide sequences. Please note that '-' or 'X' are treated as unique characters in p-distance calculations. The function will not accept 'X' or gaps in Grantham or Sandberg distance calculations. If you wish to exclude codons with 'X' or gaps from distance calculations, please use the `codon_pos` option to specify which codons to compare.

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

If you calculated Grantham or Sandberg distances, please additionally cite either of the following references: Grantham R. (1974). Amino acid difference formula to help explain protein evolution. *Science* 185:862-864. Sandberg M, Eriksson L, Jonsson J, Sjostrom M, Wold S. (1998). New chemical descriptors relevant for the design of biologically active peptides. A multivariate characterization of 87 amino acids. *JMed Chem.* 41(14):2481-2491.

Value

The function returns a matrix with distances from all pairwise sequence comparisons, where n is the number of sequences. If a sequence occurrence table is given as input file, the function additionally returns a table with the mean distance for each sample in the data set. If a sequence occurrence table is given as input file, the sequences are named in the output matrix by an index number that corresponds to their column number in the input file. If calculation of Sandberg distances is specified, the function additionally outputs five tables with physico-chemical z-descriptor values for each amino acid position in all sequences in the data set. All output tables are saved as .csv files in the output path.

See Also

For more information about 'dada2', visit <<https://benjjneb.github.io/dada2/>>

Examples

```
seq_file <- sequence_table_fas
path_out <- tempdir()
DistCalc(seq_file, path_out, input_fasta=NULL, input_seq="nucl", aa_dist=NULL,
codon_pos=c(1,2,3,4,5,6,7,8), dist_type="P")
```

 GetHpltStats

GetHpltStats() function

Description

[GetHpltStats](#) uses the output files produced by the HpltFind() function to calculate the mean of the mean proportion of incongruent sequences across all nests in the data set.

Usage

```
GetHpltStats(filepath)
```

Arguments

filepath is a user defined path to the folder where the output files from the HpltFind() function have been saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A mean of the mean proportion of incongruent sequences for each nest.

See Also

[HpltFind](#); [GetHpltTable](#)

Examples

```
filepath <- system.file("extdata/HpltFindOut/", package="MHCtools")
GetHpltStats(filepath)
```

GetHpltTable	<i>GetHpltTable() function</i>
--------------	--------------------------------

Description

[GetHpltTable](#) uses the output files produced by the [HpltFind\(\)](#) function to produce a table with the mean proportion of incongruent sequences for each nest. If the mean proportion of incongruent sequences is generally low, but certain nests have many incongruent sequences, biological reasons may be causing the mismatches, e.g. extra-pair fertilizations or recombination events.

Usage

```
GetHpltTable(filepath)
```

Arguments

`filepath` is a user defined path to the folder where the output files from the [HpltFind\(\)](#) function have been saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A table with the mean proportion of incongruent sequences for each nest.

See Also

[HpltFind](#); [GetHpltStats](#)

Examples

```
filepath <- system.file("extdata/HpltFindOut/", package="MHCtools")
GetHpltTable(filepath)
```

GetReplStats

GetReplStats function

Description

[GetReplStats](#) uses the output files produced by the `ReplMatch()` function to calculate statistics on the agreement between replicated samples in the sequencing experiment.

Usage

```
GetReplStats(filepath)
```

Arguments

`filepath` is a user defined path to the folder where the output files from the `ReplMatch()` function have been saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A list containing the number of replicate sets with zero incongruent sequences, the proportion of replicate sets with zero incongruent sequences, the mean of the mean proportion of incongruent sequences across all replicate sets, and the repeatability of the sequencing experiment.

See Also

[ReplMatch](#); [GetReplTable](#)

Examples

```
filepath <- system.file("extdata/ReplMatchOut/", package="MHCtools")
GetReplStats(filepath)
```

GetReplTable	<i>GetReplTable function</i>
--------------	------------------------------

Description

[GetReplTable](#) uses the output files produced by the `ReplMatch()` function to produce a table with the replicate sets and their respective mean proportion of incongruent sequences.

Usage

```
GetReplTable(filepath)
```

Arguments

`filepath` is a user defined path to the folder where the output files from the `ReplMatch()` function have been saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A table with the mean proportion of incongruent sequences for each replicate set.

See Also

[ReplMatch](#); [GetReplStats](#)

Examples

```
filepath <- system.file("extdata/ReplMatchOut/", package="MHCtools")
GetReplTable(filepath)
```

HpltFind

*HpltFind() function***Description**

HpltFind is designed to automatically infer major histocompatibility complex (MHC) haplotypes from the genotypes of parents and offspring in families (defined as nests) in non-model species, where MHC sequence variants cannot be identified as belonging to individual loci. **HpltFind()** assumes that data originated from a diploid species. The functions **GetHpltTable()**, **GetHpltStats()**, and **NestTablesXL()** are designed to evaluate the output files.

Usage

```
HpltFind(nest_table, seq_table, alpha = 0.8, path_out)
```

Arguments

nest_table	is a table containing the sample names of parents and offspring in each nest. This table should be organized so that the individual names are in the first column (Sample_ID), and the nest number is in the second column (Nest). For each nest, the first two rows should be the parents, followed immediately by the offspring in the subsequent rows, and then followed by the next nest, and so on. It is assumed that nests are numbered consecutively beginning at 1.
seq_table	seq_table is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns.
alpha	a numerical value between 0 and 1 (default 0.8) specifying a threshold by which a set of sequences overlapping between a chick and a parent will be assigned to the putative parental A haplotype or passed to the B haplotype. Typical values are in the range 0.6-0.9. In data sets with many different MHC alleles per individual (i.e. many MHC gene copies), alpha may be set high. In data sets with fewer MHC alleles per individual, it should be set lower. A range of alpha values may be tested to find the optimal setting for a given data set, e.g. by evaluating the mean proportion of incongruent sequences across the data set using GetHpltStats() .
path_out	is a user defined path to the folder where the output files will be saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A set of R lists containing for each nest the putative haplotypes, the names of sequences that could not be resolved with certainty in each parent, the names of the sequences that were incongruent in the genotypes of the nest, and the mean proportion of incongruent sequences (which is a measure of the haplotype inference success and largely influenced by the exactness of the genotyping experiment). The sequences are named in the output by an index number corresponding to their column number in the sequence table, thus identical sequences will have identical sample names in all the output files. These files can be reopened in R e.g. using the `readRDS()` function in the base package. Note: `HpltFind()` will overwrite any existing files with the same output file names in `path_out`.

See Also

[GetHpltTable](#); [GetHpltStats](#); [NestTablesXL](#); [CreateHpltOccTable](#); for more information about 'dada2' visit <https://benjjneb.github.io/dada2/>

Examples

```
nest_table <- nest_table
seq_table <- sequence_table
path_out <- tempdir()
HpltFind(nest_table, seq_table, alpha=0.8, path_out)
```

HpltMatch

HpltMatch() function

Description

Putative haplotypes may be identical to each other, or they may differ only by incongruent or unresolved sequences. It is therefore useful to curate putative haplotypes by comparing them to identify potentially overlapping types as candidates for further investigation. `HpltMatch` calculates the proportion of matching sequences between pairs of haplotypes and produces a .csv table with values in a lower left matrix. If a threshold value is specified, a list of haplotype matches where the proportion of matching sequences exceeds the threshold will be produced.

Usage

```
HpltMatch(hplt_occ_matrix, path_out, threshold = NULL)
```

Arguments

<code>hplt_occ_matrix</code>	A binary (logical) occurrence matrix with the data set sequences in columns and the putative haplotypes in rows, as produced by the <code>CreateHpltOccTable()</code> function.
<code>path_out</code>	a user defined path to the folder where the output file(s) will be saved.
<code>threshold</code>	a numerical value between 0 and 1 (default NULL) specifying a threshold for the proportion of matching sequences between haplotypes.

Details

Note: The `NestTablesXL()` function provides a useful format for further investigation of potentially overlapping haplotypes.

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. *Cran*. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), *HLA Typing: Methods and Protocols* (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A table specifying the proportions of matching sequences between pairs of haplotypes (in a lower left matrix). If a threshold value is specified, a list of haplotype matches where the proportion of matching sequences exceeds the threshold will be printed to the console. The list will also be saved in the output path, and can be reopened in R e.g. using the `readRDS()` function in the base package. Note: `HpltMatch()` will overwrite any existing files with the same output file names in `path_out`.

See Also

[HpltFind](#); [CreateHpltOccTable](#); [NestTablesXL](#)

Examples

```
hplt_occ_matrix <- hplt_occurrence_matrix
path_out <- tempdir()
HpltMatch(hplt_occ_matrix, path_out, threshold=NULL)
```

`hplt_occurrence_matrix`

Data `hplt_occurrence_matrix`

Description

`hplt_occurrence_matrix` is an example of a binary occurrence matrix derived from a randomized real major histocompatibility complex (MHC) data set. The matrix was generated using the `CreateHpltOccTable()` function.

Usage

```
hplt_occurrence_matrix
```

Format

`hplt_occurrence_matrix` is a data frame with 136 putative haplotypes in rows and 329 MHC sequence variants in columns.

Source

original data.

<code>k_summary_table</code>	<i>k_summary_table.rda</i>
------------------------------	----------------------------

Description

`k_summary_table` contains the results from a bootstrapped kmeans clustering analysis performed on the test data in the tables `z1_matrix_test_data`, `z2_matrix_test_data`, `z3_matrix_test_data`, `z4_matrix_test_data`, and `z5_matrix_test_data` using `BootKmeans()`.

Usage

`k_summary_table`

Format

`k_summary_table` is a data frame with observations from 10 k-estimation scans in rows and their respective stats in 11 columns.

Source

original data.

<code>NestTablesXL</code>	<i>NestTablesXL() function</i>
---------------------------	--------------------------------

Description

[NestTablesXL](#) reads the R lists output by the `HpltFind()` function and translates them to an Excel workbook for more convenient evaluation of the inferred haplotypes and curation of unresolved and incongruent sequences. The workbook contains separate tabs for each nest in the data set and provides an overview of the genotypes of the samples in each nest and the inferred haplotypes.

Usage

`NestTablesXL(nest_table, seq_table, filepath, path_out)`

Arguments

<code>nest_table</code>	is a table containing the sample names of parents and offspring in each nest. This table should be organized so that the individual names are in the first column (<code>Sample_ID</code>), and the nest number is in the second column (<code>Nest</code>). For each nest, the first two rows should be the parents, followed immediately by the offspring in the subsequent rows, and then followed by the next nest, and so on. It is assumed that nests are numbered consecutively beginning at 1. Please use the same table as was used to generate the haplotypes using <code>HpltFind()</code> .
<code>seq_table</code>	<code>seq_table</code> is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns. Please use the same table as was used to generate the haplotypes using <code>HpltFind()</code> .
<code>filepath</code>	is a user defined path to the folder where the output files from the <code>HpltFind()</code> function have been saved.
<code>path_out</code>	is a user defined path to the folder where the output file will be saved.

Details

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

An Excel workbook with individual tabs for each nest in `nest_table`. Each tab contains a binary (logical) occurrence matrix with the samples from each nest in columns and sequences (inherited from `seq_table`) in rows. The order of the samples is derived from `nest_table`, with parents in the two leftmost columns. Each tab also lists the putative haplotypes inferred by the `HpltFind()` function and provides lists of unresolved sequences in haplotypes, sequences with unidentified decent (i.e., present in parents but not in offspring), sequences not assigned to haplotypes, and sequences with unidentified origin (i.e., present in offspring but not in parents). Note: `NestTablesXL()` will overwrite any existing file with the output file name in `path_out`.

See Also

[HpltFind](https://benjjneb.github.io/dada2/); for more information about 'dada2' visit <<https://benjjneb.github.io/dada2/>>

Examples

```
nest_table <- nest_table
seq_table <- sequence_table_hplt
filepath <- system.file("extdata/HpltFindOut/", package="MHCtools")
path_out <- tempdir()
NestTablesXL(nest_table, seq_table, filepath, path_out)
```

nest_table	<i>Data nest_table</i>
------------	------------------------

Description

nest_table, parents_table, and sequence_table comprise a randomized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment with data from parents and offspring. The nucleotide sequences have been replaced with randomly generated sequences, and sample names have been anonymized.

Usage

```
nest_table
```

Format

nest_table is a data frame with 213 samples in rows and 2 columns:

Sample_ID Sample ID

Nest Nest index number

Source

original data.

PapaDiv	<i>PapaDiv() function</i>
---------	---------------------------

Description

[PapaDiv](#) calculates the joint major histocompatibility complex (MHC) diversity in parent pairs, taking into account alleles that are shared between the parents. The joint diversity in parent pairs is often of interest in studies of mate choice, fitness, and heritability.

Usage

```
PapaDiv(parents_table, seq_table, path_out)
```

Arguments

parents_table is a table containing the sample names of the parents in each nest. This table should be organized so that each row represents one nest, with the individual names of the mothers in the first column (Mother), and the individual names of the fathers in the second column (Father).

seq_table seq_table is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns.

path_out is a user defined path to the folder where the output files will be saved.

Details

The `PapaDiv()` function outputs a set of R lists containing for the joint diversity of each parent pair, the proportion of sequences that are shared between the parents, the diversity of each of the parents, the observed sequence variants in each parent, the matched sequence variants, and the incongruent sequence variants in each parent.

In addition, `PapaDiv()` produces a summary table with the names of the parents in a pair, their respective MHC diversities, and the joint parent pair diversity.

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

a set of R lists containing for the joint diversity of each parent pair, the proportion of sequences that are shared between the parents, the diversity of each of the parents, the observed sequence variants in each parent, the matched sequence variants, and the incongruent sequence variants in each parent. The sequences are named in the output by an index number corresponding to their column number in the sequence table, thus identical sequences will have identical sample names in all the output files. These files are saved in a sub folder in the output path called `Parent_pairs` (created by `PapaDiv()`) and can be reopened in R e.g. using the `readRDS()` function in the base package. For downstream data analysis, the `PapaDiv()` function also produces a summary table with the names of the parents in a pair, their respective MHC diversities, and the joint parent pair diversity. This table is saved as a `.csv` file in the output path.

See Also

For more information about 'dada2' visit <https://benjjneb.github.io/dada2/>

Examples

```
parents_table <- parents_table
seq_table <- sequence_table
path_out <- tempdir()
PapaDiv(parents_table, seq_table, path_out)
```

parents_table

Data parents_table

Description

`nest_table`, `parents_table`, and `sequence_table` comprise a randomized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment with data from parents and offspring. The nucleotide sequences have been replaced with randomly generated sequences, and sample names have been anonymized.

Usage

parents_table

Format

parents_table is a data frame with 57 parent pairs in rows and 2 columns:

Mother Mother ID

Father Father ID

Source

original data.

replicates_table	<i>Data replicates_table</i>
------------------	------------------------------

Description

replicates_table and sequence_table_repl comprise a randomized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment with technical replicates. The nucleotide sequences have been replaced with randomly generated sequences, and sample names have been anonymized.

Usage

replicates_table

Format

replicates_table is a data frame with 111 technical replicate samples in rows and 2 columns:

Sample_ID Technical replicate sample ID

Replic_set Index number of replicate set

Source

original data.

ReplMatch	<i>ReplMatch()</i> function
-----------	-----------------------------

Description

In amplicon filtering it is sometimes valuable to compare technical replicates in order to estimate the accuracy of a genotyping experiment. This may be done both to optimize filtering settings and to estimate repeatability to report in a publication. `ReplMatch` is designed to automatically compare technical replicates in an amplicon filtering data set and report the proportion of mismatches. The functions `GetReplTable()` and `GetReplStats()` are designed to evaluate the output files.

Usage

```
ReplMatch(repl_table, seq_table, path_out)
```

Arguments

<code>repl_table</code>	is a table containing the sample names of technical replicates in the data set. This table should be organized so that the individual names are in the first column (<code>Sample_ID</code>), and the index number of the replicate set is in the second column (<code>Replic_set</code>). Replicate sets may contain more than two replicates, but sets must be numbered consecutively beginning at 1.
<code>seq_table</code>	<code>seq_table</code> is a sequence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequence variants in columns.
<code>path_out</code>	is a user defined path to the folder where the output files will be saved.

Details

Note: `ReplMatch()` will throw a warning if all samples in a replicate set have 0 sequences. In that case, the `mean_props` for that replicate set and the repeatability for the data set will be `NaN`, and `ReplMatch()` will report which replicate set is problematic and suggest to remove it from the `repl_table`. If removing replicate sets, beware that the replicate sets in `repl_table` must be numbered consecutively beginning at 1.

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

A set of R lists containing for each replicate set the observed sequence variants, the names of the sequences that were incongruent in the replicates, and the mean proportion of incongruent sequences (if 100 matches are expected between the replicates, this is equivalent of an error rate in the sequencing process). The sequences are named in the output by an index number corresponding to their column number in the sequence table, thus identical sequences will have identical sample names in all the output files. These files can be reopened in R e.g. using the `readRDS()` function in the base package.

See Also

[GetReplTable](#); [GetReplStats](#); for more information about 'dada2' visit <<https://benjjneb.github.io/dada2/>>

Examples

```
repl_table <- replicates_table
seq_table <- sequence_table_repl
path_out <- tempdir()
ReplMatch(repl_table, seq_table, path_out)
```

sequence_table	<i>Data sequence_table</i>
----------------	----------------------------

Description

nest_table, parents_table, and sequence_table comprise a randomized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment with data from parents and offspring. The nucleotide sequences have been replaced with randomly generated sequences, and sample names have been anonymized.

Usage

```
sequence_table
```

Format

sequence_table is a data frame with 334 samples in rows and 329 DNA sequence variants in columns.

Source

original data.

sequence_table_fas	<i>Data sequence_table_fas</i>
--------------------	--------------------------------

Description

sequence_table_fas is a randomized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment. The nucleotide sequences have been replaced with randomly generated sequences, and sample names have been anonymized.

Usage

```
sequence_table_fas
```

Format

sequence_table_fas is a data frame with 100 samples in rows and 166 DNA sequence variants in columns.

Source

original data.

sequence_table_hplt *Data sequence_table_hplt*

Description

sequence_table_hplt is a randomized test data set derived from a real major histocompatibility complex (MHC) genotyping experiment. This table differs from the sequence_table by having the nucleotide sequences replaced with sequence names. Sample names have been anonymized from the real data set.

Usage

sequence_table_hplt

Format

sequence_table_hplt is a data frame with 334 samples in rows and 329 sequence variants in columns.

Source

original data.

sequence_table_repl *Data sequence_table_repl*

Description

replicates_table and sequence_table_repl comprise a randomized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment with technical replicates. The nucleotide sequences have been replaced with randomly generated sequences, and sample names have been anonymized.

Usage

sequence_table_repl

Format

sequence_table_repl is a data frame with 412 samples in rows and 511 DNA sequence variants in columns.

Source

original data.

sequence_table_SynDist

Data sequence_table_SynDist

Description

Sequence_table_SynDist comprises an anonymized dataset derived from a real major histocompatibility complex (MHC) genotyping experiment.

Usage

sequence_table_SynDist

Format

sequence_table_SynDist is a data frame with 20 samples in rows and 20 DNA sequence variants in columns.

Source

original data.

SynDist

SynDist() function

Description

[SynDist](#) identifies and quantifies synonymous variation among aligned protein-coding DNA sequences, that is, nucleotide substitutions that do not translate to changes in the amino acid sequences, due to degeneracy of the genetic code.

Usage

```
SynDist(
  seq_file,
  path_out,
  input_fasta = NULL,
  codon_pos = NULL,
  analysis = "dist"
)
```

Arguments

seq_file	is a sequence occurrence table as output by the 'dada2' pipeline, which has samples in rows and nucleotide sequences in columns. Optionally, a fasta file can be supplied as input in the format rendered by read.fasta() from the package 'seqinr'.
path_out	is a user defined path to the folder where the output files will be saved.
input_fasta	optional, a logical (TRUE/FALSE) that indicates whether the input file is a fasta file (TRUE) or a 'dada2'-style sequence table (NULL/FALSE). The default is NULL/FALSE.
codon_pos	is optional, a vector of comma separated integers specifying which codons to include in analyses. If omitted, analyses are made using all codons. Note: With SynDist(), codon_pos should always be specified as codons, i.e. numbered nucleotide triplets in open reading frame.
analysis	is used to specify the desired kind of analysis. It takes the values 'dist' for quantification of pairwise synonymous variation between sequences, or 'codon' for quantification of synonymous substitutions per nucleotide or codon position. The argument is optional with 'dist' as default.

Details

The SynDist() function takes a fasta file or a 'dada2'-style sequence occurrence table (with aligned sequences as column names and samples in rows) as input and identifies synonymous variation by pairwise sequence comparisons.

SynDist() can do qualitative or quantitative analysis of synonymous variation. If analysis="codon" is specified, the function identifies synonymous nucleotide variation and outputs tables with the number of observations of synonymous nucleotide changes per base and per codon among all pairwise sequence comparisons in the data set. These tables also specify, for each base or codon position, the proportions of the total pairwise comparisons that harbor synonymous substitutions.

If analysis="dist", the function produces a distance matrix specifying the number and proportion (p-distance) of synonymous nucleotide changes in each pairwise sequence comparison in the data set. In the distance matrix, synonymous p-distance is calculated as the number of synonymous nucleotide changes observed in each pairwise sequence comparison divided by the sequence length (number of bases). If a 'dada2'-style sequence occurrence table is provided as input, the SynDist() function furthermore produces two tables with the mean number of synonymous variations and mean synonymous p-distances among all pairwise comparisons of the sequences in each sample in the data set. (Note: The means will be NA for samples that have 0 or 1 sequence(s).)

The SynDist() function includes an option for the user to specify which codons to compare. This is useful e.g. if the sequences contain gaps in some codons, which should be excluded from quantitative analysis.

SynDist() translates the supplied DNA sequences to amino acid sequences using the standard genetic code and sequences must be aligned in open reading frame. The function only accepts the following characters in the sequences: -,a,t,g,c,A,T,G,C

Nucleotide triplets containing gaps (-) are translated to 'X', similar to stop codons. Please note that '-' are treated as unique characters in p-distance calculations. The function will give warnings if gaps or stop codons are detected. If you wish to exclude stop codons or gaps from distance calculations, please use the codon_pos option to specify which codons to compare.

If you publish data or results produced with MHCtools, please cite both of the following references: Roved, J. (2022). MHCtools: Analysis of MHC data in non-model species. Cran. Roved, J. (2024). MHCtools 1.5: Analysis of MHC sequencing data in R. In S. Boegel (Ed.), HLA Typing: Methods and Protocols (2nd ed., pp. 275–295). Humana Press. https://doi.org/10.1007/978-1-0716-3874-3_18

Value

When `analysis="dist"`, the function produces a .csv distance matrix with the number of synonymous substitutions in each pairwise sequence comparison in the upper right matrix and the synonymous p-distance in each pairwise sequence comparison in the lower left matrix. If a sequence occurrence table is given as input file, the function additionally produces two tables with the mean number of synonymous substitutions and the mean synonymous p-distance across all pairwise sequence comparisons for each sample in the data set. If a sequence occurrence table is given as input file, the sequences are named in the output matrix by an index number that corresponds to their column number in the input file. If `analysis="codon"`, the function produces two .csv summary tables, one with the total number of synonymous substitutions per nucleotide position across all pairwise sequence comparisons and one with the number of synonymous codon variations per codon across all pairwise sequence comparisons. Note that in the codon summary table, the synonymous codon variation does not quantify the number of nucleotide variations between the synonymous codons, since that can be derived from the nucleotide summary table. Each summary table also contains a column that specifies the proportion of the observed number of synonymous variations (per nucleotide position or codon) out of the number of pairwise sequence comparisons. E.g., if three sequences are compared and a synonymous substitution is observed for a given codon once (i.e., between two of the three sequences), that gives a proportion of synonymous observations of one out of three pairwise sequence comparisons for that codon.

See Also

For more information about 'dada2', visit <https://benjjneb.github.io/dada2/>

Examples

```
seq_file <- sequence_table_SynDist
path_out <- tempdir()
SynDist(seq_file, path_out, input_fasta=NULL, codon_pos=c(1,2,3,4,5,6,7,8),
analysis="dist")
```

`z1_matrix`

`z1_matrix.rda`

Description

`z1_matrix` comprise a randomized dataset derived from 70 nucleotide sequences of a real major histocompatibility complex (MHC) genotyping experiment. z1-descriptor values have been extracted from a subset of 8 amino acid codons and sequence names have been anonymized.

Usage

z1_matrix

Format

z1_matrix is a data frame with 70 sequences in rows and z1-descriptor variables for 8 sequence codons in columns.

Source

original data.

z2_matrix	<i>z2_matrix.rda</i>
-----------	----------------------

Description

z2_matrix comprise a randomized dataset derived from 70 nucleotide sequences of a real major histocompatibility complex (MHC) genotyping experiment. z2-descriptor values have been extracted from a subset of 8 amino acid codons and sequence names have been anonymized.

Usage

z2_matrix

Format

z2_matrix is a data frame with 70 sequences in rows and z2-descriptor variables for 8 sequence codons in columns.

Source

original data.

z3_matrix	<i>z3_matrix.rda</i>
-----------	----------------------

Description

z3_matrix comprise a randomized dataset derived from 70 nucleotide sequences of a real major histocompatibility complex (MHC) genotyping experiment. z3-descriptor values have been extracted from a subset of 8 amino acid codons and sequence names have been anonymized.

Usage

z3_matrix

Format

z3_matrix is a data frame with 70 sequences in rows and z3-descriptor variables for 8 sequence codons in columns.

Source

original data.

z4_matrix	<i>z4_matrix.rda</i>
-----------	----------------------

Description

z4_matrix comprise a randomized dataset derived from 70 nucleotide sequences of a real major histocompatibility complex (MHC) genotyping experiment. z4-descriptor values have been extracted from a subset of 8 amino acid codons and sequence names have been anonymized.

Usage

```
z4_matrix
```

Format

z4_matrix is a data frame with 70 sequences in rows and z4-descriptor variables for 8 sequence codons in columns.

Source

original data.

z5_matrix	<i>z5_matrix.rda</i>
-----------	----------------------

Description

z5_matrix comprise a randomized dataset derived from 70 nucleotide sequences of a real major histocompatibility complex (MHC) genotyping experiment. z5-descriptor values have been extracted from a subset of 8 amino acid codons and sequence names have been anonymized.

Usage

```
z5_matrix
```

Format

z5_matrix is a data frame with 70 sequences in rows and z5-descriptor variables for 8 sequence codons in columns.

Source

original data.

Index

* datasets

hplt_occurrence_matrix, 18
k_summary_table, 19
nest_table, 21
parents_table, 22
replicates_table, 23
sequence_table, 25
sequence_table_fas, 25
sequence_table_hplt, 26
sequence_table_repl, 26
sequence_table_SynDist, 27
z1_matrix, 29
z2_matrix, 30
z3_matrix, 30
z4_matrix, 31
z5_matrix, 31

BootKmeans, 3, 3, 7

ClusterMatch, 5, 5, 6
CreateFas, 7, 7, 9
CreateHpltOccTable, 8, 8, 17, 18
CreateSamplesFas, 8, 9, 9

DistCalc, 5, 10, 10

GetHpltStats, 12, 12, 13, 17
GetHpltTable, 13, 13, 17
GetReplStats, 14, 14, 15, 25
GetReplTable, 14, 15, 15, 25

hplt_occurrence_matrix, 18
HpltFind, 9, 13, 16, 16, 18, 20
HpltMatch, 17, 17

k_summary_table, 19

nest_table, 21
NestTablesXL, 17–19, 19

PapaDiv, 21, 21

parents_table, 22

replicates_table, 23
ReplMatch, 14, 15, 24, 24

sequence_table, 25
sequence_table_fas, 25
sequence_table_hplt, 26
sequence_table_repl, 26
sequence_table_SynDist, 27
SynDist, 27, 27

z1_matrix, 29
z2_matrix, 30
z3_matrix, 30
z4_matrix, 31
z5_matrix, 31