

Package ‘disclapmix2’

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Type Package

Title Mixtures of Discrete Laplace Distributions using Numerical Optimisation

Version 0.6.1

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Description

Fit a mixture of Discrete Laplace distributions using plain numerical optimisation. This package has similar applications as the 'disclapmix' package that uses an EM algorithm.

License GPL (>= 2)

Imports Rcpp (>= 1.0.3), cluster

LinkingTo Rcpp

RoxygenNote 7.2.1

Encoding UTF-8

Suggests testthat, disclapmix, readxl

NeedsCompilation yes

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`disclapmix2`*Discrete Laplace mixture inference using Numerical Optimisation*

Description

An extension to the `*disclapmix*` method in the `*disclapmix*` package that supports duplicated loci and other non-standard haplotypes.

Description of your package

Usage

```
disclapmix2(  
  x,  
  number_of_clusters,  
  include_2_loci = FALSE,  
  remove_non_standard_haplotypes = TRUE,  
  use_stripped_data_for_initial_clustering = FALSE,  
  initial_y_method = "pam",  
  verbose = 0L  
)
```

Arguments

`x` DataFrame. Columns should be one character vector for each locus

`number_of_clusters` The number of clusters to fit the model for.

`include_2_loci` Should duplicated loci be included or excluded from the analysis?

`remove_non_standard_haplotypes` Should observations that are not single integer alleles be removed?

`use_stripped_data_for_initial_clustering` Should non_standard data be removed for the initial clustering?

`initial_y_method` Which cluster method to use for finding initial central haplotypes, y: pam (recommended) or clara.

`verbose` Set to 1 (or higher) to print optimisation details. Default is 0.

Value

List.

Author(s)

you

Examples

```

require(disclapmix)

data(danes)

x <- as.matrix(danes[rep(seq_len(nrow(danes))), danes$n], -ncol(danes)])
x2 <- as.data.frame(sapply(danes[rep(seq_len(nrow(danes))), danes$n], -ncol(danes)], as.character))

d1m_fit <- disclapmix(x, clusters = 3L)
d1m2_fit <- disclapmix2(x2, number_of_clusters = 3)

stopifnot(all.equal(d1m_fit$logL_marginal, d1m2_fit$log_lik))

```

haplotype_counts	<i>Count the number of times each haplotype occurs</i>
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Description

Count the number of times each haplotype occurs

Usage

```
haplotype_counts(x)
```

Arguments

`x` DataFrame (by locus) of character vectors containing haplotypes (rows) where alleles are separated by comma's, e.g. "13,14.2" is a haplotype

Value

Integer vector with count for each row in DataFrame

Examples

```

# read haplotypes
h <- readxl::read_excel(system.file("extdata", "South_Australia.xlsx",
package = "disclapmix2"),
col_types = "text")[-c(1,2)]

# obtain counts
counts <- disclapmix2::haplotype_counts(h)

# all haplotypes in the dataset are unique
stopifnot(all(counts == 1))

```

profile_pr_by_locus_and_cluster
Compute Profile Probability from fit

Description

Compute the profile probability for a new profile that was not used in the original fit.

Usage

```
profile_pr_by_locus_and_cluster(x, fit)
```

Arguments

x	DataFrame. Columns should be one character vector for each locus
fit	Output from disclapmix2

Value

Numeric.

Examples

```
require(disclapmix)

data(danes)

x <- as.data.frame(sapply(danes[rep(seq_len(nrow(danes))), danes$n], -ncol(danes)], as.character))

d1m2_fit <- disclapmix2(x, number_of_clusters = 3)

new_profile <- structure(list(DYS19 = "14", DYS389I = "13", DYS389II = "29",
  DYS390 = "22", DYS391 = "9", DYS392 = "15", DYS393 = "13",
  DYS437 = "14", DYS438 = "11", DYS439 = "12"),
  row.names = 1L, class = "data.frame")

profile_pr_by_locus_and_cluster(x = new_profile, d1m2_fit)
```

`unique_haplotype_counts`*List unique haplotypes with their counts*

Description

List unique haplotypes with their counts

Usage

```
unique_haplotype_counts(x)
```

Arguments

`x` DataFrame (by locus) of character vectors containing haplotypes (rows) where alleles are separated by comma's, e.g. "13,14.2" is a haplotype

Value

DataFrame with unique rows and a Count column added at the end

Examples

```
# read haplotypes
h <- readxl::read_excel(system.file("extdata", "South_Australia.xlsx",
  package = "disclapmix2"),
  col_types = "text")[-c(1,2)]

# obtain counts
unique_counts <- disclapmix2::unique_haplotype_counts(h)

# all haplotypes in the dataset are unique
stopifnot(all(unique_counts$Count == 1))
```

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