Package ‘TraMineR’

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Description Toolbox for the manipulation, description and rendering of sequences, and more generally the mining of sequence data in the field of social sciences. Although the toolbox is primarily intended for analyzing state or event sequences that describe life courses such as family formation histories or professional careers, its features also apply to many other kinds of categorical sequence data. It accepts many different sequence representations as input and provides tools for converting sequences from one format to another. It offers several functions for describing and rendering sequences, for computing distances between sequences with different metrics (among which optimal matching), original dissimilarity-based analysis tools, and simple functions for extracting the most frequent subsequences and identifying the most discriminating ones among them. A user's guide can be found on the TraMineR web page.
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R topics documented:

- TraMineR-package ................................................. 4
- actcal ........................................................................ 5
- actcal.tse ................................................................... 6
- alphabet ..................................................................... 7
- bfspell ........................................................................ 8
- biofam ......................................................................... 9
- cpal ........................................................................... 11
- dissassoc .................................................................... 12
- disscenter ................................................................... 14
- dissmfacw ................................................................... 16
- dissrep ......................................................................... 18
- distree ........................................................................ 21
- disstree2dot ................................................................. 23
- disstreeleaf ................................................................. 26
- dissvr .......................................................................... 27
- ex1 ........................................................................... 28
- ex2 ........................................................................... 29
- famform ....................................................................... 30
- mvad ........................................................................... 30
- plot.seqdiff ................................................................. 32
- plot.stslist ................................................................. 33
- plot.stslist.freq .......................................................... 35
- plot.stslist.meant ........................................................ 37
- plot.stslist.modst ........................................................ 38
- plot.stslist.rep ............................................................ 39
- plot.stslist.statd .......................................................... 41
- plot.subseqelist ........................................................... 43
- plot.subseqelistchisq ................................................... 44
- read.tda.mdist ............................................................ 45
- seqalign ................................................................. 46
- seqcomp ................................................................. 47
- seqconc ................................................................. 48
- seqcost ................................................................. 49
- seqdecomp ............................................................... 52
- seqdef ................................................................. 53
- seqdiff ................................................................. 57
- seqdim ................................................................. 59
- seqdist ................................................................. 60
- seqdistmc ............................................................... 66
- seqdss ................................................................. 68
- seqdur ................................................................. 69
- seqeapplysub .............................................................. 70
- seqecmpgroup ............................................................ 72
- seqeconstraint ............................................................ 73
- seqecontain ............................................................... 75
- seqecreate ............................................................... 76
<table>
<thead>
<tr>
<th>R topics documented:</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>seqefsub</td>
<td>78</td>
</tr>
<tr>
<td>seqeid</td>
<td>81</td>
</tr>
<tr>
<td>seqelength</td>
<td>81</td>
</tr>
<tr>
<td>seqetm</td>
<td>82</td>
</tr>
<tr>
<td>seqeweight</td>
<td>84</td>
</tr>
<tr>
<td>seqfind</td>
<td>85</td>
</tr>
<tr>
<td>seqformat</td>
<td>86</td>
</tr>
<tr>
<td>seqfpos</td>
<td>91</td>
</tr>
<tr>
<td>seqgen</td>
<td>92</td>
</tr>
<tr>
<td>seqici</td>
<td>93</td>
</tr>
<tr>
<td>seqient</td>
<td>94</td>
</tr>
<tr>
<td>seqistatd</td>
<td>96</td>
</tr>
<tr>
<td>seqlegend</td>
<td>97</td>
</tr>
<tr>
<td>seqlength</td>
<td>98</td>
</tr>
<tr>
<td>seqLLCP</td>
<td>99</td>
</tr>
<tr>
<td>seqLLCS</td>
<td>100</td>
</tr>
<tr>
<td>seqlogp</td>
<td>101</td>
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<tr>
<td>seqmeant</td>
<td>102</td>
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<td>seqmodst</td>
<td>104</td>
</tr>
<tr>
<td>seqmpos</td>
<td>105</td>
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<tr>
<td>seqnum</td>
<td>106</td>
</tr>
<tr>
<td>seqpceplot</td>
<td>107</td>
</tr>
<tr>
<td>seqplot</td>
<td>112</td>
</tr>
<tr>
<td>seqpm</td>
<td>118</td>
</tr>
<tr>
<td>seqrecode</td>
<td>119</td>
</tr>
<tr>
<td>seqrep</td>
<td>121</td>
</tr>
<tr>
<td>seqsep</td>
<td>124</td>
</tr>
<tr>
<td>seqST</td>
<td>125</td>
</tr>
<tr>
<td>seqstad</td>
<td>126</td>
</tr>
<tr>
<td>seqstatf</td>
<td>128</td>
</tr>
<tr>
<td>seqstatl</td>
<td>129</td>
</tr>
<tr>
<td>seqsubsn</td>
<td>130</td>
</tr>
<tr>
<td>seqtab</td>
<td>131</td>
</tr>
<tr>
<td>seqtransn</td>
<td>133</td>
</tr>
<tr>
<td>seqtrate</td>
<td>134</td>
</tr>
<tr>
<td>seqtree</td>
<td>136</td>
</tr>
<tr>
<td>seqtreeedisplay</td>
<td>138</td>
</tr>
<tr>
<td>stlab</td>
<td>140</td>
</tr>
<tr>
<td>TraMineR.check.depr.args</td>
<td>141</td>
</tr>
<tr>
<td>TraMineR.checkupdates</td>
<td>142</td>
</tr>
<tr>
<td>TraMineRInternal</td>
<td>142</td>
</tr>
</tbody>
</table>
Description

(Version: 2.0-12) Toolbox for the manipulation, description and rendering of sequences, and more generally the mining of sequence data in the field of social sciences. Although the toolbox is primarily intended for analyzing state or event sequences that describe life courses such as family formation histories or professional careers, its features also apply to many other kinds of categorical sequence data. It accepts many different sequence representations as input and provides tools for converting sequences from one format to another. It offers several functions for describing and rendering sequences, for computing distances between sequences with different metrics (among which optimal matching), original dissimilarity-based analysis tools, and simple functions for extracting the most frequent subsequences and identifying the most discriminating ones among them. A user’s guide can be found on the TraMineR web page.

Details

TraMineR provides tools for both state sequences and event sequences. The first step when using the package is to define a state sequence object (with `seqdef`) if you want to explore state sequences, and an event sequence object (with `seqecreate`) if you are interested in event sequencing.

State sequences are defined from a series of variables giving the states at the successive positions, while event sequences are defined from (vertical) time stamped event data. The package, however, can handle many other different data organizations and provides tools to help converting state sequences into event sequences and vice versa.

Author(s)

Alexis Gabadinho, Matthias Studer, Nicolas S. Müller, Reto Bürgin, Pierre-Alexandre Fonta, and Gilbert Ritschard

References


Examples

```r
# load the mvad data
library(TraMineR)
data(mvad)

# create a state sequence object from columns 17 to 86
mvad.seq <- seqdef(mvad[,17:86])
```
```r
## distribution plot by sex (male)
seqdplot(mvad.seq, group=mvad$male, border=NA)

## compute the LCS pairwise distance matrix
## among the first 10 sequences
mvad.lcs <- seqdist(mvad.seq[1:10,], method="LCS")
```

### Description

This data set contains 2000 individual sequences of monthly activity statuses from January to December 2000.

### Usage

```r
data(actcal)
```

### Format

A data frame with 2000 rows, 12 state variables, 1 id variable and 11 covariates.

### Details

The data set is a subsample of the data collected by the Swiss Household Panel (SHP).

The state column (variable) names are ‘jan00’, ‘feb00’, etc... and correspond to columns 13 to 24.

There are four possible states:

- **A** = Full-time paid job (> 37 hours)
- **B** = Long part-time paid job (19-36 hours)
- **C** = Short part-time paid job (1-18 hours)
- **D** = Unemployed (no work)

The data set contains also the following covariates:

- **age00** (age in 2000)
- **educat00** (education level)
- **civsta00** (civil status)
- **nbadul00** (number of adults in household)
- **nbkid00** (number of children)
- **aoldki00** (age of oldest kid)
- **ayouki00** (age of youngest kid)
- **region00** (residence region)
- **com2.00** (residence commune type)
sex (sex of respondent)
birthy (birth year)

Source
Swiss Household Panel

References
https://forscenter.ch/projects/swiss-household-panel/

actcal.tse  Example data set: Activity calendar from the Swiss Household Panel (time stamped event format)

Description
This data set contains events defined from the state sequences in the actcal data set. It was created with the code shown in the examples section. It is provided to simplify example of event sequence mining.

Usage
data(actcal.tse)

Format
Time stamped events derived from state sequences in the actcal data set.

Source
Swiss Household Panel

See Also
seqformat, actcal

Examples
data(actcal)
actcal.seq <- seqdef(actcal[,1:24])

## Defining the transition matrix
transition <- seqetm(actcal.seq, method="transition")
transitions <- c("Increase,FullTime", "Increase,PartTime", "LowPartTime", "Stop")

### Converting STS data to TSE
actcal.tse <- seqformat(actcal, 13:24, from = "STS", to = "TSE",
tevent = transition)

### Defining the event sequence object
actcal.eseq <- seqecreate(id = actcal.tse$id,
time = actcal.tse$time, event = actcal.tse$event)

---

### alphabet

#### Get or set the alphabet of a state or event sequence object

**Description**

For state sequences, the function gets or sets the (short) labels associated to the states in the alphabet of a state sequence object (the list of all possible states). The get form also applies to event sequences, while the set form does not work with event sequences.

**Usage**

alphabet(seqdata)
alphabet(seqdata) <- value

**Arguments**

- seqdata: a state sequence object as defined with the `seqdef` function or, for the get form only, an event sequence object as defined with `seqecreate`, or a probabilistic suffix tree generated with the PST package.
- value: For state sequences only. Vector of characters of the same length as the vector returned by the `alphabet` function, i.e. one label for each state in the alphabet.

**Details**

A state sequence object—created with the `seqdef` function—stores sequences as a matrix where columns are factors. The levels of the factors include the alphabet plus the codes for missing values and void elements. The alphabet function retrieves or sets the “alphabet” attribute of the state sequence object. The state names composing the alphabet are preferably short labels, since they are used for printing sequences. Longer labels for describing more precisely each state in legend are stored in the “labels” attribute of the sequence object.

For an event sequence object—created with `seqecreate`—the get form of `alphabet` works as an alias for `levels`. The set form `alphabet` <- does not work and should not be used.
bfspell

Example data set: First 20 biofam sequences in SPELL form

Description

First 20 sequences of the biofam data set in SPELL form. The data serve to illustrate the use of seqformat for converting SPELL data into STS (horizontal) form.

Usage

data(bfspell)

Format

A data set with two data frames: bfspell20 with one row per spell and bfpdata20 with one row per id. The bfspell20 data frame contains the spell data themselves (4 variables id, begin, end, states) and bfpdata20 the year when aged 15 (2 variables id, when15).
Details

The states are coded with the following short labels

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See Also

biofam

---

biofam Example data set: Family life states from the Swiss Household Panel biographical survey

Description

2000 16 year-long family life sequences built from the retrospective biographical survey carried out by the Swiss Household Panel (SHP) in 2002.

Usage

data(biofam)
data(bfspell)

Format

A data frame with 2000 rows, 16 state variables, 1 id variable and 7 covariates and 2 weights variables.

Details

The biofam data set was constructed by Müller et al. (2007) from the data of the retrospective biographical survey carried out by the Swiss Household Panel (SHP) in 2002.

The data set contains (in columns 10 to 25) sequences of family life states from age 15 to 30 (sequence length is 16) and a series of covariates. The sequences are a sample of 2000 sequences of those created from the SHP biographical survey. It includes only individuals who were at least
30 years old at the time of the survey. The biofam data set describes family life courses of 2000 individuals born between 1909 and 1972.

The states numbered from 0 to 7 are defined from the combination of five basic states, namely Living with parents (Parent), Left home (Left), Married (Marr), Having Children (Child), Divorced:

0 = "Parent"
1 = "Left"
2 = "Married"
3 = "Left+Marr"
4 = "Child"
5 = "Left+Child"
6 = "Left+Marr+Child"
7 = "Divorced"

The covariates are:

- **sex**
- **birthyr** (birth year)
- **nat_1_02** (first nationality)
- **plingu02** (language of questionnaire)
- **p02r01** (religion)
- **p02r04** (religious participation)
- **cspfaj** (father’s social status)
- **cspmoj** (mother’s social status)

Two additional weights variables are inserted for illustrative purpose ONLY (since biofam is a subsample of the original data, these weights are not adapted to the actual data):

- **wp00tbgp** (weights inflating to the Swiss population)
- **wp00tbgs** (weights respecting sample size)

**Source**

Swiss Household Panel [https://forscenter.ch/projects/swiss-household-panel/](https://forscenter.ch/projects/swiss-household-panel/)

**References**

cpal

Get or set the color palette of a sequence object

Description

This function gets or sets the color palette of a sequence object, that is, the list of colors used to represent the states.

Usage

```r
cpal(seqdata)
```

```r
cpal(seqdata) <- value
```

Arguments

- `seqdata` a state sequence object as defined by the `seqdef` function.
- `value` a vector containing the colors, of length equal to the number of states in the alphabet. The colors can be passed as character strings representing color names such as returned by the `colors` function, as hexadecimal values or as RGB vectors using the `rgb` function. Each color is attributed to the corresponding state in the alphabet, the order being the one returned by the `alphabet`.

Details

In the plot functions provided for visualizing sequence objects, a different color is associated to each state of the alphabet. The color palette is defined when creating the sequence object, either automatically using the `brewer.pal` function of the `RColorBrewer` package or by specifying a user defined color vector. The cpal function can be used to get or set the color palette of a previously defined sequence object.

Value

For `cpal(seqdata)` a vector containing the colors.

For `cpal(seqdata) <-` the updated sequence object.

Author(s)

Alexis Gabadinho

See Also

`seqdef`
Examples

```r
## Creating a sequence object with the columns 13 to 24
## in the `actcal` example data set
## The color palette is automatically set
data(actcal)
actcal.seq <- seqdef(actcal,13:24)

## Retrieving the color palette
cpal(actcal.seq)
seqiplot(actcal.seq)

## Setting a user defined color palette
cpal(actcal.seq) <- c("blue","red","green","yellow")
seqiplot(actcal.seq)
```

---

dissassoc  

*Analysis of discrepancy from dissimilarity measures*

Description

Compute and test the share of discrepancy (defined from a dissimilarity matrix) explained by a categorical variable.

Usage

```r
dissassoc(diss, group, weights=NULL, R=1000,
          weight.permutation="replicate", squared=FALSE)
```

Arguments

- **diss**: A dissimilarity matrix or a dist object (see `dist`)
- **group**: A categorical variable. For a numerical variable use `dissmfacw`.
- **weights**: optional numerical vector containing weights.
- **R**: Number of permutations for computing the p-value. If equal to 1, no permutation test is performed.
- **weight.permutation**: Weighted permutation method: "diss" (attach weights to the dissimilarity matrix), "replicate" (replicate case using weights), "rounded-replicate" (replicate case using rounded weights), "random-sampling" (random assignment of covariate profiles to the objects using distributions defined by the weights.)
- **squared**: Logical. If TRUE the dissimilarities diss are squared.
Details

The dissassoc function assesses the association between objects characterized by their dissimilarity matrix and a discrete covariate. It provides a generalization of the ANOVA principle to any kind of distance metric. The function returns a pseudo R-square that can be interpreted as a usual R-square. The statistical significance of the association is computed by means of permutation tests. The function performs also a test of discrepancy homogeneity (equality of within variances) using a generalization of the Levene statistic and Bartlett’s statistics. There are print and hist methods (the latter producing an histogram of the permuted values used for testing the significance).

If a numeric group variable is provided, it will be treated as categorical, i.e., each different value will be considered as a different category. To measure the ‘linear’ effect of a numerical variable, use dissfmcw.

Value

An object of class dissassoc with the following components:

- groups: A data frame with the number of cases and the discrepancy of each group
- anova.table: The pseudo ANOVA table
- stat: The value of the statistics and their p-values
- perms: The permutation object, containing the values computed for each permutation

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

dissvar to compute the pseudo variance from dissimilarities and for a basic introduction to concepts of pseudo variance analysis.
disstree for an induction tree analyse of objects characterized by a dissimilarity matrix.
disscenter to compute the distance of each object to its group center from pairwise dissimilarities. dissmanova to perform multi-factor analysis of variance from pairwise dissimilarities.

Examples

```r
## Defining a state sequence object
data(mvad)
mvad.seq <- seqdef(mvad[, 17:86])

## Building dissimilarities (any dissimilarity measure can be used)
mvad.ham <- seqdist(mvad.seq, method="HAM")

## R=1 implies no permutation test
da <- dissassoc(mvad.ham, group=mvad$gcseSeq, R=1)
print(da)
print(da)

hist(da)
```

disscenter  

### Compute distances to the center of a group

**Description**

Computes the dissimilarity between objects and their group center from their pairwise dissimilarity matrix.

**Usage**

disscenter(diss, group=NULL, medoids.index=NULL, allcenter = FALSE, weights=NULL, squared=FALSE)

**Arguments**

- **diss**: a dissimilarity matrix such as generated by `seqdist`, or a dist object (see `dist`)
- **group**: if NULL (default), the whole data set is considered. Otherwise a different center is considered for each distinct value of the group variable
- **medoids.index**: if NULL, returns the dissimilarity to the center. If set to "first", returns the index of the first encountered most central sequence. If group is set, an index is returned per group. When set to "all", indexes of all medoids (one list per group) are returned.
- **allcenter**: logical. If TRUE, returns a data frame containing the dissimilarity between each object and its group center, each column corresponding to a group.
- **weights**: optional numerical vector containing weights.
- **squared**: Logical. If TRUE diss is squared.
Details

This function computes the dissimilarity between given objects and their group center. It is possible that the group center does not belong to the space formed by the objects (in the same way as the average of integer numbers is not necessarily an integer itself). This distance can also be understood as the contribution to the discrepancy (see \texttt{dissvar}). Note that when the dissimilarity measure does not respect the triangle inequality, the dissimilarity between a given object and its group center may be negative.

It can be shown that this dissimilarity is equal to (see Batagelj 1988):

\[
d_{x\bar{g}} = \frac{1}{n} \left( \sum_{i=1}^{n} d_{xi} - SS \right)
\]

where \(SS\) is the sum of squares (see \texttt{dissvar}).

Value

A vector with the dissimilarity to the group center for each object, or a list of medoid indexes.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

\texttt{dissvar} to compute the pseudo variance from dissimilarities and for a basic introduction to concepts of pseudo variance analysis

\texttt{dissassoc} to test association between objects represented by their dissimilarities and a covariate.

\texttt{disstree} for an induction tree analyse of objects characterized by a dissimilarity matrix.

\texttt{dissmfacw} to perform multi-factor analysis of variance from pairwise dissimilarities.
The `dissmfacw` function performs a multi-factor analysis of variance from a dissimilarity matrix.

**Description**

Perform a multi-factor analysis of variance from a dissimilarity matrix.

**Usage**

```r
dissmfacw(formula, data, R = 1000, gower = FALSE, squared = FALSE, weights = NULL)
```

**Arguments**

- `formula`: A regression-like formula. The left hand side term should be a dissimilarity matrix or a `dist` object.
- `data`: A data frame from which the variables in `formula` should be taken.
- `R`: Number of permutations used to assess significance.
- `gower`: Logical: Is the dissimilarity matrix already a Gower matrix?
- `squared`: Logical: Should we square the provided dissimilarities?
- `weights`: Optional numerical vector of case weights.
Details

This method is, in some way, a generalization of `dissassoc` to account for several explanatory variables. The function computes the part of discrepancy explained by the list of covariates specified in the formula. It provides for each covariate the Type-II effect, i.e. the effect measured when removing the covariate from the full model with all variables included.

(The returned F values may slightly differ from those obtained with TraMineR versions older than 1.8-9. Since 1.8-9, the within sum of squares at the denominator is divided by \( n - m \) instead of \( n - m - 1 \), where \( n \) is the sample size and \( m \) the total number of predictors and/or contrasts used to represent categorical factors.)

For a single factor `dissmfacw` is slower than `dissassoc`. Moreover, the latter performs also tests for homogeneity in within-group discrepancies (equality of variances) with a generalization of Levene’s and Bartlett’s statistics.

Part of the function is based on the Multivariate Matrix Regression with qr decomposition algorithm written in SciPy-Python by Ondrej Libiger and Matt Zapala (See Zapala and Schork, 2006, for a full reference.) The algorithm has been adapted for Type-II effects and extended to account for case weights.

Value

A `dissmultifactor` object with the following components:

- `mfac` The part of variance explained by each variable (comparing full model to model without the specified variable) and its significance using permutation test
- `call` Function call
- `perms` Permutation values as a boot object

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also
dissvar to compute a pseudo variance from dissimilarities and for a basic introduction to concepts of discrepancy analysis.
dissassoc to test association between objects represented by their dissimilarities and a covariate.
disstree for an induction tree analysis of objects characterized by a dissimilarity matrix.
disscenter to compute the distance of each object to its group center from pairwise dissimilarities.

Examples

```r
## Define the state sequence object
data(mvad)
mvad.seq <- seqdef(mvad[, 17:86])
## Here, we use only first 100 sequences
mvad.seq <- mvad.seq[1:100,]

## Compute dissimilarities (any dissimilarity measure can be used)
mvad.ham <- seqdist(mvad.seq, method="HAM")

## And now the multi-factor analysis
print(dissmfacw(mvad.ham ~ male + Grammar + funemp + gcseSeq + fmpr + livboth, data=mvad[1:100,], R=10))
```

---

dissrep

Extracting sets of representative objects using a dissimilarity matrix

### Description

The function extracts a set of representative objects that exhibits the key features of the whole data set, the goal being to get easy sounded interpretation of the latter. The user can set either the desired coverage level (the proportion of objects having a representative in their neighborhood) or the desired number of representatives.

### Usage

```r
dissrep(diss, criterion = "density", score = NULL, decreasing = TRUE, coverage = NULL, nrep = NULL, pradius = NULL, dmax = NULL, weights = NULL, trep = NULL, tsim = NULL)
```

### Arguments

- **diss**: A dissimilarity matrix or a `dist` object (see `dist`)

---

Extracting sets of representative objects using a dissimilarity matrix

The function extracts a set of representative objects that exhibits the key features of the whole data set, the goal being to get easy sounded interpretation of the latter. The user can set either the desired coverage level (the proportion of objects having a representative in their neighborhood) or the desired number of representatives.

### Usage

```r
dissrep(diss, criterion = "density", score = NULL, decreasing = TRUE, coverage = 0.25, nrep = NULL, pradius = NULL, dmax = NULL, weights = NULL, trep = NULL, tsim = NULL)
```

### Arguments

- **diss**: A dissimilarity matrix or a `dist` object (see `dist`)
criterion  the representativeness criterion for sorting the candidate list. One of "freq" (frequency), "density" (neighborhood density) or "dist" (centrality). An optional vector containing the scores for sorting the candidate objects may also be provided. See below and details.

score  an optional vector containing the representativeness scores used for sorting the objects in the candidate list. The length of the vector must be equal to the number of rows/columns in the distance matrix, i.e the number of objects.

decreasing  if a score vector is provided, indicates whether the objects in the candidate list must be sorted in ascending or decreasing order of this score. The first object in the candidate list is supposed to be the most representative.

coverage  controls the size of the representative set by setting the desired coverage level, i.e the proportion of objects having a representative in their neighborhood. Neighborhood radius is defined by pradius.

nrep  number of representatives. If NULL (default), coverage argument is used to control the size of the representative set.

pradius  neighborhood radius as a percentage of the maximum (theoretical) distance dmax. Defaults to 0.1 (10%). Object y is redundant to object x when it is in the neighborhood of x, i.e., within a distance pradius*dmax from x.

dmax  maximum theoretical distance. Used to derive the neighborhood radius as pradius*dmax.

weights  vector of weights of length equal to the number of rows of the dissimilarity matrix. If NULL, equal weights are assigned.

trep  Deprecated. Use coverage instead.

ts1m  Deprecated. Use pradius instead.

Details

The representative set is obtained by an heuristic. Representatives are selected by successively extracting from the sequences sorted by their representativeness score those which are not redundant with already retained representatives. The selection stops when either the desired coverage or the wanted number of representatives is reached. Objects are sorted either by the values provided as score argument, or by specifying one of the following as criterion argument: "freq" (sequence frequency), "density" (neighborhood density), "dist" (centrality).

The frequency criterion uses the frequencies as representativeness score. The frequency of an object in the data is computed as the number of other objects with whom the dissimilarity is equal to 0. The more frequent an object the more representative it is supposed to be. Hence, objects are sorted in decreasing frequency order. Indeed, this criterion is the neighborhood (see below) criterion with the neighborhood diameter set to 0.

The neighborhood density is the number—density—of sequences in the neighborhood of the object. This requires to set the neighborhood radius pradius. Objects are sorted in decreasing density order.

The centrality criterion is the sum of distances to all other objects. The smallest the sum, the most representative the sequence.

Use criterion="dist" and nrep=1 to get the medoid and criterion="density" and nrep=1 to get the densest object pattern.

For more details, see Gabadinho et al., 2011.
Value

An object of class diss.rep. This is a vector containing the indexes of the representative objects with the following additional attributes:

Scores a vector with the representative score of each object given the chosen criterion.
Distances a matrix with the distance of each object to its nearest representative.
Statistics a data frame with quality measures for each representative: number of objects attributed to the representative, number of object in the representative’s neighborhood, mean distance to the representative.

Quality overall quality measure.

Print and summary methods are available.

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

References


See Also

seqrep, disscenter

Examples

```r
# Defining a sequence object with the data in columns 10 to 25
# (family status from age 15 to 30) in the biofam data set
data(biofam)
biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

# Computing the distance matrix
costs <- seqsubm(biofam.seq, method="TRATE")
biofam.om <- seqdist(biofam.seq, method="OM", sm=costs)

# Representative set using the neighborhood density criterion
biofam.rep <- dissrep(biofam.om)
biofam.rep
summary(biofam.rep)
```
**disstree**  
*Dissimilarity Tree*

**Description**

Tree structured discrepancy analysis of objects described by their pairwise dissimilarities.

**Usage**

```r
disstree(formula, data = NULL, weights = NULL, min.size = 0.05,  
max.depth = 5, R = 1000, pval = 0.01, object = NULL,  
weight.permutation = "replicate", squared = FALSE, first = NULL,  
min.size, maxdepth)
```

**Arguments**

- `formula`: Formula with a dissimilarity matrix as left hand side and the candidate partitioning variables on the right side.
- `data`: Data frame where variables in formula will be searched for.
- `weights`: Optional numerical vector of weights.
- `min.size`: Minimum number of cases in a node, will be treated as a proportion if less than 1.
- `max.depth`: Maximum depth of the tree
- `R`: Number of permutations used to assess the significance of the split.
- `pval`: Maximum allowed p-value for a split
- `object`: An optional R object represented by the dissimilarity matrix. This object may be used by the print method or `disstree2dot` to render specific object type.
- `weight.permutation`: Weight permutation method: "diss" (attach weights to the dissimilarity matrix), "replicate" (replicate cases using weights), "rounded-replicate" (replicate case using rounded weights), "random-sampling" (random assignment of covariate profiles to the objects using distributions defined by the weights).
- `squared`: Logical: Should the diss dissimilarities be squared?
- `first`: One of the variable in the right-hand side of the formula. This forces the first node of the tree to be split by this variable.
- `minSize`: Deprecated. Use `min.size` instead.
- `maxdepth`: Deprecated. Use `max.depth` instead.

**Details**

The procedure iteratively splits the data. At each step, the procedure selects the variable and split that explain the greatest part of the discrepancy, i.e., the split for which we get the highest pseudo R2. The significance of the retained split is assessed through a permutation test.

`seqtree` provides a simpler interface if you plan to use `disstree` for state sequence objects.
Value

An object of class disstree that contains the following components:

- **root**: A node object, root of the tree
- **info**: General information such as parameters used to build the tree
- **info$adjustment**: A `dissassoc` object providing global statistics for the tree.
- **formula**: The formula used to generate the tree
- **data**: Data used to build the tree
- **weights**: Weights

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

- `seqtree` to generate a specific `disstree` objects for analyzing state sequences.
- `seqtreedisplay` to generate graphic representation of `seqtree` objects when analyzing state sequences.
- `disstreedisplay` is a more general interface to generate such representation for other type of objects.
- `dissvar` to compute discrepancy using dissimilarities and for a basic introduction to discrepancy analysis.
- `dissassoc` to test association between objects represented by their dissimilarities and a covariate.
- `dissmfacw` to perform multi-factor analysis of variance from pairwise dissimilarities.
- `disscenter` to compute the distance of each object to its group center from pairwise dissimilarities.
disstree2dot

Examples

```r
# Defining a state sequence object
mvad.seq <- seqdef(mvad[, 17:86])

# Computing dissimilarities (any dissimilarity measure can be used)
mvad.ham <- seqdist(mvad.seq, method="HAM")
# Grow the tree using a low R value for illustration.
# For R=10, pval cannot be lower than 0.1
dt <- disstree(mvad.ham ~ male + Grammar + funemp + gcse5seq + fnmr + livboth,
               data=mvad, R = 10, pval = 0.1)
print(dt)

# Will only work if GraphViz is properly installed
# See seqtree for simpler way to plot a sequence tree.
# Not run:
disstreedisplay(dt, image.fun = seqdplot, image.data = mvad.seq,
                 # Additional parameters passed to seqdplot
                 with.legend = FALSE, axes = FALSE, ylab = "")

# End(Not run)
# Second method, using a specific function
myplotfunction <- function(individuals, seqs, ...) {
  par(font.sub=2, mar=c(3,0,0,0), mgp=c(0,0,0))
  # using mds to order sequence in seqiplot
  mds <- cmdscale(seqdist(seqs[individuals,], method="HAM"),k=1)
  seqiplot(seqs[individuals,], sortv=mds,...)
}

# If image.data is not set, index of individuals are sent to image.fun
# Not run:
disstreedisplay(dt, image.fun = myplotfunction, cex.main = 3,
                 # additional parameters passed to myplotfunction
                 seqs = mvad.seq,
                 # additional parameters passed to seqiplot (through myplotfunction)
                 with.legend = FALSE, axes = FALSE, idxs = 0, space = 0, ylab = ",", border = NA)

# End(Not run)
```

disstree2dot

Graphical representation of a dissimilarity tree

Description

Functions to generate a "dot" file and associated images files that can be used in GraphViz to get a graphical representation of the tree.
Usage

```r
disstree2dot(tree, filename, digits = 3, image.fun = NULL, image.data = NULL,
only.leaf = FALSE, device = "jpeg", image.format = "jpg",
device.args = list(), use.title = TRUE, label.pos = "main",
node.pos = "main", split.pos = "sub", cex.main = 1,
legend.text = NULL, image.legend = NULL, image.quality = NULL,
show.depth = FALSE, title.outer = FALSE,
imagefunc, imagedata, imgLeafOnly, devicefunc, imageext,
device.arg, label.loc, node.loc, split.loc, title.cex, legendtext,
legendimage, qualityimage, showdepth, ...)

disstree2dotp(tree, filename, image.data = NULL, only.leaf = FALSE,
image.fun = plot, cex.main = 3, with.quality = TRUE,
cex.quality = cex.main, title.outer = FALSE,
imagedata, imgLeafOnly, imagefunc, title.cex, withquality,
quality.fontsize, ...)

seqtree2dot(tree, filename, seqdata = tree$info$object, only.leaf = FALSE,
sortv = NULL, diss = NULL, cex.main = 3, with.legend = "auto",
cex.legend = cex.main, with.quality = FALSE,
cex.quality = cex.main, axes = FALSE,
imgLeafOnly, dist.matrix, title.cex,
withlegend, withquality, ...)
```

Arguments

- `tree` The tree to be plotted.
- `filename` A filename, without extension, that will be used to generate image and dot files.
- `digits` Number of significant digits to plot.
- `image.fun` A function to plot the individuals in a node, see details.
- `image.data` A data.frame that will be passed to `image.fun`, see details.
- `only.leaf` Logical: If TRUE, only terminal node will be plotted.
- `device` A device function, "jpeg" by default.
- `device.format` Extension for image files.
- `device.args` Argument passed to device.
- `use.title` Logical: If TRUE, node information will be printed using `title` command, see details.
- `label.pos` Location of the node label, see `title` for possible values.
- `node.pos` Node content location, see `title` for possible values.
- `split.pos` Split information location, see `title` for possible values.
- `cex.main` cex applied to all calls to title (see use.title).
- `title.outer` Logical: If TRUE, the title (see use.title) is printed in the outer margins.
- `legend.text` An optional text appearing in a distinct node.
image.legend An optional image file appearing in a distinct node.
image.quality An optional image file appearing in a distinct node.
show.depth Logical. If TRUE, information about depth of the tree is added to the plot.
with.quality If TRUE, a node displaying fitting measures of the tree is added to the plot.
cex.quality Numeric. Size of the font of the fitting measures node.
seqdata a sequence object as defined by the the seqdef function.
sortv The name of an optional variable used to sort the data before plotting, see seqplot.
diss The name of an optional dissimilarity matrix used to find representative sequences, seqrplot.
with.legend defines if and where the legend of the state colors is plotted. The default value "auto" sets the position of the legend automatically. Other possible value is "right". Obsolete value TRUE is equivalent to "auto".
cex.legend Size of the font of the legend.
axes if set to "all" (default value) x axes are drawn for each plot in the graphic. If set to "bottom" and group is used, axes are drawn only under the plots located at the bottom of the graphic area. If FALSE, no x axis is drawn.
imagefunc Deprecated. Use image.fun instead.
imagemdata Deprecated. Use image.data instead.
imgLeafOnly Deprecated. Use only.leaf instead.
devicefunc Deprecated. Use device instead.
imageext Deprecated. Use image.format instead.
device.arg Deprecated. Use device.args instead.
label.loc Deprecated. Use label.pos instead.
node.loc Deprecated. Use node.pos instead.
split.loc Deprecated. Use split.pos instead.
title.cex Deprecated. Use cex.main instead.
legendtext Deprecated. Use legend.text instead.
legendimage Deprecated. Use image.legend instead.
qualityimage Deprecated. Use image.quality instead.
showdepth Deprecated. Use show.depth instead.
withquality Deprecated. Use with.quality instead.
quality.fontsize Deprecated. Use cex.quality instead.
dist.matrix Deprecated. Use diss instead.
withlegend Deprecated. Use with.legend instead.
... other parameters that will be passed to image.fun or seqplot (for seqtree2dot).
Details

These functions generate a "dot" file that can be used in GraphViz (http://www.graphviz.org). It also generates one image per node through a call to image.fun passing the selected lines of image.data if present or otherwise a list of indexes (of individuals belonging to a node). These functions are not intended to be used by end-user. See seqtreedisplay and disstreedisplay for a much simpler way to generate a graphical representation of a tree (seqtree or disstree).

seqtree2dot is a shortcut for sequences objects using the plot function seqplot. For each node, it calls seqplot with the corresponding subset of rows of seqdata and the provided seqplot’s arguments. You should at least specify the type of the plot (e.g. type="d", see seqplot for more details).

If use.title is TRUE, image.fun should take care to leave enough space for the title.
disstree2dotp is a simplified interface of disstree2dot which automatically leaves enough space for the title and subtitles. These functions are intended to be generic.

Value

Nothing but generates a "dot" and several image files (one per node) in the current working directory (see getwd and setwd).

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

See Also

seqtree and seqtreedisplay, disstree and disstreedisplay.

### disstreeleaf

**Terminal node membership**

Return a factor with the terminal node membership of each case.

#### Usage

```r
disstreeleaf(tree, label=FALSE)
```

#### Arguments

- `tree` The tree, a disstree or DissTreeNode object.
- `label` Logical. If TRUE, the returned leaf memberships are labelled with the corresponding classifications rules.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)
dissvar

See Also
disstree for examples

---

dissvar Dissimilarity based discrepancy

Description
Compute the discrepancy from the pairwise dissimilarities between objects. The discrepancy is a measure of dispersion of the set of objects.

Usage
dissvar(diss, weights=NULL, squared = FALSE)

Arguments
diss A dissimilarity matrix or a dist object (see dist)
weights optional numerical vector containing weights.
squared Logical. If TRUE diss is squared.

Details
The discrepancy is an extension of the concept of variance to any kind of objects for which we can compute pairwise dissimilarities. The discrepancy $s^2$ is defined as:

$$ s^2 = \frac{1}{2n^2} \sum_{i=1}^{n} \sum_{j=1}^{n} d_{ij} $$

**Mathematical ground:** In the Euclidean case, the sum of squares can be expressed as:

$$ SS = \sum_{i=1}^{n} (y_i - \bar{y})^2 = \frac{1}{2n} \sum_{i=1}^{n} \sum_{j=1}^{n} (y_i - y_j)^2 $$

The concept of discrepancy generalizes the equation by allowing to replace the $(y_i - y_j)^2$ term with any measure of dissimilarity $d_{ij}$.

Value
The discrepancy.

Author(s)
Matthias Studer (with Gilbert Ritschard for the help page)
References


See Also

dissassoc to test association between objects represented by their dissimilarities and a covariate.
disstree for an induction tree analyse of objects characterized by a dissimilarity matrix.
disscenter to compute the distance of each object to its group center from pairwise dissimilarities.
dissmfacw to perform multi-factor analysis of variance from pairwise dissimilarities.

Examples

```r
## Defining a state sequence object
data(mvad)
mvad.seq <- seqdef(mvad[, 17:86])

## Building dissimilarities (any dissimilarity measure can be used)
mvad.ham <- seqdist(mvad.seq, method="HAM")

## Pseudo variance of the sequences
print(dissvar(mvad.ham))
```

---

**ex1**  
Example data set with missing values and weights

---

Description

Example data set used to demonstrate the handling of missing values and weights.

The state columns (variable) are named 'P1' to 'P13'.

The alphabet is made of four possible states: A, B, C and D.

The data set contains also case weights (variable weights). The sum of the weights is 60.
Usage
data(ex1)

Format
A data frame with 7 rows, 13 state variables, 1 weight variable.

Source
The brain of the TraMineR package team.

Description
Example data sets used to demonstrate the handling of weights. The 'ex2.weighted' data set contains 6 sequences with weights inflating to 100 sequences (sum of weights is 100). The second data frame 'ex2.unweighted' contains the corresponding 100 sequences.
The sequences are, in both data frames, in the 'seq' column, and weights in the 'weight' column of 'ex2.weighted'.
The alphabet is made of four possible states: A, B, C and D.

These data sets are mainly intended to test and illustrate the handling of weights in TraMineR's functions. Weighted results obtained with 'ex2.weighted' data set should be exactly the same as unweighted results obtained with the 'ex2.unweighted' data set.

Usage
data(ex2)

Format
The command data(ex2) generates two data frames:
ex2.weighted: a data frame with 6 rows, 1 variable containing sequences as character strings, 1 weight variable.
ex2.unweighted: a data frame with 100 rows, 1 variable containing sequences as character strings.

Source
The brain of the TraMineR package team.

Examples
data(ex2)

ex2w.seq <- seqdef(ex2.weighted, 1, weights=ex2.weighted$weight)
ex2u.seq <- seqdef(ex2.unweighted)
Example data set: sequences of family formation

Description
This data set contains 5 sequences of family formation histories, used by Elzinga (2008) to introduce several metrics for computing distances between sequences. These sequences don’t contain information about the duration spent in each state, they contain only distinct successive states.

Usage

```
data(famform)
```

Format
A data frame with 5 rows and 1 variable.

Details
The sequences are in ‘STS’ format and stored in character strings with states separated with ‘-’.
This data set is used in TraMineR’s manual to crosscheck some results with those presented by Elzinga.

Source
Elzinga (2008)

References

Example data set: Transition from school to work

Description
The data comes from a study by McVicar and Anyadike-Danes on transition from school to work. The data consist of static background characteristics and a time series sequence of 72 monthly labour market activities for each of 712 individuals in a cohort survey. The individuals were followed up from July 1993 to June 1999. The monthly states are recorded in columns 15 (Ju1. 93) to 86 (Jun. 99).
States are:
employment (EM)
FE further education (FE)
HE higher education (HE)
joblessness (JL)
school (SC)
training (TR)

The data set contains also ids (id) and sample weights (weight) as well as the following binary covariates:

male
catholic
Belfast, N.Eastern, Southern, S.Eastern, Western (location of school, one of five Education and Library Board areas in Northern Ireland)
Grammar (type of secondary education, 1=grammar school)
funemp (father's employment status at time of survey, 1=father unemployed)
gcse5eq (qualifications gained by the end of compulsory education, 1=5+ GCSEs at grades A-C, or equivalent)
fmpr (SOC code of father's current or most recent job, 1=SOC1 (professional, managerial or related))
livboth (living arrangements at time of first sweep of survey (June 1995), 1=living with both parents)

Usage
data(mvad)

Format
A data frame containing 712 rows, 72 state variables, 1 id variable and 13 covariates.

Source
McVicar and Anyadike-Danes (2002)

References
Description

Plot method for the sliding values returned by seqdiff. Plots a statistic (the Pseudo R2 by default) along the position axis.

Usage

```
## S3 method for class 'seqdiff'
plot(x, stat = "Pseudo R2", type = "l", ylab = stat,
     xlab = ",", legend.pos = "top", ylim = NULL, xaxis = TRUE, col = NULL,
     xstep = NULL, tick.last = NULL, legendposition, xaxt, ...)
```

Arguments

- `x`: an object produced by seqdiff
- `stat`: character. Name of the statistic to be plotted. Can be any of the statistics returned by seqdiff or "discrepancy". See details.
- `type`: the line type, see lines
- `ylab`: character: y-axis label.
- `xlab`: character: x-axis label.
- `legend.pos`: character: position of the line legend, see legend
- `ylim`: numeric: if not NULL, range of the y-axis.
- `xaxis`: logical: if TRUE an x-axis is plotted.
- `col`: list of colors to use for each line.
- `xstep`: integer: optional step between tick-marks and labels on the x-axis. If unspecified, the xstep attribute of the sequence object x is used. (see seqdef)
- `tick.last`: Logical. Should a tick mark be enforced at the last position on the x-axis? If unspecified, the tick.last attribute of the x object is used.
- `legendposition`: Deprecated. Use legend.pos instead.
- `xaxt`: Deprecated. Use xaxis instead.
- `...`: Additional parameters passed to lines

Details

The function plots the sliding values of the requested statistic.

You can plot the evolution of two statistics by providing for instance `stat=c("Pseudo R2", "Levene")`. Use `stat="discrepancy"` to plot the within-discrepancies.

For "discrepancy", a separate line is drawn for the whole set of sequences and for each group. Those two values cannot be paired with another statistic.
Author(s)
Matthias Studer (with Gilbert Ritschard for the help page)

See Also
seqdiff

plot.stslist  Plot method for state sequence objects

Description
This is the plot method for state sequence objects of class stslist created by the seqdef function. It produces a sequence index plot.

Usage
## S3 method for class 'stslist'
plot(x, idxs = NULL, weighted = TRUE, sortv = NULL, cpal = NULL, missing.color = NULL, ylab = NULL, yaxis = TRUE, xaxis = TRUE, ytlab = NULL, ylas = 0, xlab = NULL, xtstep = NULL, tick.last = NULL, cex.axis = 1, tlim, cex.plot, ...)

Arguments

- **x**
  A state sequence object created with the seqdef function.

- **idxs**
  Indexes of the sequences to be plotted (default value is 1:10), for instance 20:50 to plot sequences 20 to 50, c(2, 8, 12, 25) to plot sequences 2, 8, 12 and 25 in seqdata. If set to NULL, all sequences in seqdata are plotted.

- **weighted**
  Logical: Should the bar representing each sequence be proportional to its weight? Ignored when no weights are assigned to sequences (see seqdef).

- **sortv**
  A sorting variable or a sort method (one of "from.start" or "from.end"). See details.

- **cpal**
  alternative color palette to use for the states. If user specified, a vector of colors with number of elements equal to the number of states in the alphabet. By default, the cpal attribute of the seqdata sequence object is used (see seqdef).

- **missing.color**
  alternative color for representing missing values inside the sequences. By default, this color is taken from the "missing.color" attribute of the x sequence object.

- **ylab**
  An optional label for the y axis. If set to NA, no label is drawn.

- **yaxis**
  Controls whether the y axis is plotted or not. When set to TRUE, sequence indexes are displayed.

- **xaxis**
  if TRUE (default), the x (time) axis is plotted.
ytlab  
    the labels of the plotted sequences to display on the y axis. Default is the indexes of the sequences as defined by the `idxs` argument. Can be set to "id" for displaying the row names (id) of the sequences instead of their indexes; row names can be assigned to the sequence object with the `id` argument of the `seqdef` function or afterwards with `rownames`. Otherwise `ytlab` can be set to a vector of length equal to the number of sequences to be plotted.

ylas  
    sets the orientation of the sequence labels appearing on the y axis. Accepted values are the same as for the `las` standard option
    0: always parallel to the axis (default),
    1: always horizontal,
    2: always perpendicular to the axis,
    3: always vertical.

xtlab  
    optional labels for the x axis ticks labels. If unspecified, the column names of the `seqdata` sequence object are used (see `seqdef`).

xtstep  
    optional interval at which the tick-marks and labels of the x-axis are displayed. For example, with `xtstep=3` a tick-mark is drawn at position 1, 4, 7, etc... The display of the corresponding labels depends on the available space and is dealt with automatically. If unspecified, the `xtstep` attribute of the `x` object is used.

tick.last  
    Logical. Should a tick mark be enforced at the last position on the x-axis? If unspecified, the `tick.last` attribute of the `x` object is used.

cex.axis  
    Axis annotation magnification. See `par`.

tlim  
    Deprecated. Use `idxs` instead.

cex.plot  
    Deprecated. Use `cex.axis` instead.

...  
    arguments to be passed to the plot function or other graphical parameters.

Details

This is the default plot method for state sequence objects (produced by the `seqdef` function), i.e., for objects of class `stslist`. It produces a sequence index plot, where individual sequences are rendered with stacked bars depicting the states over time.

This method is called by the generic `seqplot` function (if `type="i"`). The latter produces more sophisticated plots, allowing grouping and automatic display of the state color legend. The `seqiplot` function is a shortcut for calling `seqplot` with `type="i"`.

When a `sortv` variable is provided to `seqiplot` or `seqplot`, its values define the order in which the sequences are plotted. With `sortv = "from.start"`, sequence are sorted by the elements of the alphabet at the successive positions starting from the beginning of the sequences. The "from.end" method proceeds similarly, but backward from the last position.

The interest of sequence index plots has for instance been stressed by Scherer (2001) and Brzinsky-Fay et al. (2006). Notice that such index plots for thousands of sequences result in very heavy graphic files if they are stored in PDF or POSTSCRIPT format. To reduce the size, we suggest saving the figures in bitmap format by using for instance `png` instead of `postscript` or `pdf`.

See Also

`seqplot`
Examples

```r
## Defining a sequence object with the data in columns 10 to 25
## (family status from age 15 to 30) in the biofam data set
biofam <- biofam[500:600,]  ## using a subsample only
data(biofam)

biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

## Plot of the 10 most frequent sequences
## with bar width proportional to the frequency
plot(biofam.seq)

## Plotting the whole data set
## with no borders
plot(biofam.seq, idxs=0, space=0, border=NA)

## Weights
##
data(ex1)
ex1.seq <- seqdef(ex1, 1:13, weights=ex1$weights)
plot(ex1.seq)
plot(ex1.seq, weighted=FALSE)
```

### Description

Plot method for output produced by the `seqtab` function, i.e objects of class `stslist.freq`.

### Usage

```r
## S3 method for class 'stslist.freq'
plot(x, cpal = NULL, missing.color = NULL, pbarw = TRUE,
     ylab = NULL, yaxis = TRUE, xaxis = TRUE,
     xtlab = NULL, xtstep = NULL, tick.last = NULL, cex.axis = 1,
     cex.plot, ...)  
```

### Arguments

- **x**: an object of class `stslist.freq` as produced by the `seqtab` function.
- **cpal**: alternative color palette to be used for the states. If user specified, a vector of colors with number of elements equal to the number of states in the alphabet. By default, the ‘cpal’ attribute of the x object is used.
- **missing.color**: alternative color for representing missing values inside the sequences. By default, this color is taken from the `missing.color` attribute of the x object.
pbarw if pbarw=TRUE (default), the width of the bars are proportional to the sequence frequency in the dataset.

ylab an optional label for the y axis. If set to NA, no label is drawn.

yaxis if TRUE or "cum", the y axis is plotted with a label showing the cumulated percentage frequency of the displayed sequences. If "pct", the percentage value for each sequence is displayed.

xaxis if TRUE (default) the x-axis is plotted.

xtlab optional labels for the ticks of the x-axis. If unspecified, the names attribute of the x object is used.

xtstep optional interval at which the tick-marks and labels of the x-axis are displayed. For example, with xtstep=3 a tick-mark is drawn at position 1, 4, 7, etc... The display of the corresponding labels depends on the available space and is dealt with automatically. If unspecified, the xtstep attribute of the x object is used.

tick.last Logical. Should a tick mark be enforced at the last position on the x-axis? If unspecified, the tick.last attribute of the x object is used.

cex.axis Axis annotation magnification. See par.

... further graphical parameters. For example border=NA to remove the bars borders, space=0 to remove space between sequences. For more details about the graphical parameter arguments, see barplot and par.

cex.plot Deprecated. Use cex.axis instead.

Details

This is the plot method for the output produced by the seqtab function, i.e. objects of class stslist.freq. It produces a plot showing the sequences sorted bottom up according to their frequency in the data set.

This method is called by the generic seqplot function (if type="f") that produces more sophisticated plots, allowing grouping and automatic display of the state color legend. The seqfplot function is a shortcut for calling seqplot with type="f".

Author(s)

Alexis Gabadinho

Examples

```r
## Loading the 'actcal' example data set
data(actcal)

## Defining a sequence object with data in columns 13 to 24
## (activity status from January to December 2000)
actcal.lab <- c("> 37 hours", "19-36 hours", "1-18 hours", "no work")
actcal.seq <- seqdef(actcal, 13:24, labels=actcal.lab)

## 10 most frequent sequences in the data
actcal.freq <- seqtab(actcal.seq)
```
## Plot method for objects produced by the seqmeant function

### Description

This is the plot method for objects of class `stslist.meant` produced by the `seqmeant` function.

### Usage

```r
## S3 method for class 'stslist.meant'
plot(x, cpal = NULL, ylab = NULL, yaxis = TRUE,
     xaxis = TRUE, cex.axis = 1, ylim = NULL, cex.plot, ...)
```

### Arguments

- **x**
  - an object of class `stslist.meant` as produced by the `seqmeant` function.
- **cpal**
  - alternative color palette to use for the states. If user specified, a vector of colors with number of elements equal to the number of states in the alphabet. By default, the `cpal` attribute of the `seqdata` sequence object is used (see `seqdef`).
- **ylab**
  - an optional label for the y axis. If set to `NA`, no label is drawn.
- **yaxis**
  - controls whether the y axis is plotted. Default is `TRUE`.
- **xaxis**
  - if `TRUE` (default) the xaxis is plotted.
- **cex.axis**
  - Axis annotation magnification. See `par`.
- **ylim**
  - an optional vector setting the limits for the y axis. If `NULL` (default), limits are set to (0, max. sequence length).
- **cex.plot**
  - Deprecated. Use `cex.axis` instead.
- **...**
  - further graphical parameters. For more details about the graphical parameter arguments, see `barplot` and `par`.

### Details

This is the plot method for the output produced by the `seqmeant` function, i.e., objects of class `stslist.meant`. It produces a plot showing the mean times spent in each state of the alphabet.

When the "se" attribute of `x` is `TRUE`, i.e., when `x` contains also the standard errors of the mean times, error bars are automatically displayed on the plot. See the `serr` argument of `seqmeant`.

This method is called by the generic `seqplot` function (if `type=``mt``") that produces more sophisticated plots, allowing grouping and automatic display of the states legend. The `seqmtplot` function is a shortcut for calling `seqplot` with `type=``mt``".

## Example

```r
## Plotting the object
plot(actcal.freq, main="Sequence frequencies - actcal data set")

## Plotting all the distinct sequences without borders
## and space between sequences
actcal.freq2 <- seqtab(actcal.seq, idxs=0)
plot(actcal.freq2, main="Sequence frequencies - actcal data set",
     border=NA, space=0)
```
Examples

```r
## Loading the mvad data set and creating a sequence object
data(mvad)
mvad.labels <- c("employment", "further education", "higher education",
                "joblessness", "school", "training")
mvad.scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad.seq <- seqdef(mvad, 15:86, states=mvad.scodes, labels=mvad.labels)

## Computing the mean times
mvad.meant <- seqmeant(mvad.seq)

## Plotting
plot(mvad.meant, main="Mean durations in each state of the alphabet")

## Changing the y axis limits
plot(mvad.meant, main="Mean durations in each state of the alphabet",
     ylim=c(0,40))

## Displaying error bars
mvad.meant.e <- seqmeant(mvad.seq, serr=TRUE)
plot(mvad.meant.e, main="Mean durations in each state of the alphabet",
     ylim=c(0,40))
```

Description

Plot method for output produced by the seqmodst function, i.e objects of class stslist.modst.

Usage

```r
## S3 method for class 'stslist.modst'
plot(x, cpal = NULL, ylab = NULL, yaxis = TRUE,
     xaxis = TRUE, xtlab = NULL, xtstep = NULL, tick.last = NULL,
     cex.axis = 1, cex.plot, ...)
```

Arguments

- `x`: an object of class stslist.modst as produced by the seqmodst function.
- `cpal`: alternative color palette to use for the states. If user specified, a vector of colors with number of elements equal to the number of states in the alphabet. By default, the 'cpal' attribute of the x object is used.
- `ylab`: an optional label for the y axis. If set to NA, no label is drawn.
- `yaxis`: if TRUE (default) the y axis is plotted.
- `xaxis`: if TRUE (default) the x axis is plotted.
xtlab

optional labels for the x axis ticks. If unspecified, the names attribute of the x object is used.

xtstep

optional interval at which the tick-marks and labels of the x-axis are displayed. For example, with xtstep=3 a tick-mark is drawn at position 1, 4, 7, etc... The display of the corresponding labels depends on the available space and is dealt with automatically. If unspecified, the xtstep attribute of the x object is used.

tick.last

Logical. Should a tick mark be enforced at the last position on the x-axis? If unspecified, the tick.last attribute of the x object is used.

cex.axis

Axis annotation magnification. See \texttt{par}.

cex.plot

Deprecated. Use \texttt{cex.axis} instead.

... further graphical parameters. For more details about the graphical parameter arguments, see \texttt{barplot} and \texttt{par}.

Details

This is the plot method for the output produced by the \texttt{seqmodst} function, i.e. objects of class \texttt{stslist.modst}. It produces a plot showing the sequence of modal states with bar width proportional to the state frequencies.

This method is called by the generic \texttt{seqplot} function (if \texttt{type="ms"}) that produces more sophisticated plots, allowing grouping and automatic display of the states legend. The \texttt{seqmsplot} function is a shortcut for calling \texttt{seqplot} with \texttt{type="ms"}.

Examples

```r
## Defining a sequence object with the data in columns 10 to 25
## (family status from age 15 to 30) in the biofam data set
data(biofam)
biofam.lab <- c("Parent", "Left", "Married", "Left+Marr",
    "Child", "Left+Child", "Left+Marr+Child", "Divorced")
biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

## Modal state sequence
biofam.modst <- seqmodst(biofam.seq)
plot(biofam.modst)
```

---

Plot method for representative sequence sets

Description

This is the plot method for output produced by the \texttt{seqrep} function, i.e, for objects of class \texttt{stslist.rep}. It produces a representative sequence plot.
Usage

```r
## S3 method for class 'stslist.rep'
plot(x, cpal = NULL, missing.color = NULL, pbarw = TRUE,
    dmax = NULL, stats = TRUE, ylab = NULL, xaxis = TRUE, xtlab = NULL,
    xtstep = NULL, tick.last = NULL, cex.with.axis = 1, cex.plot, ...)
```

Arguments

- **x**: an object of class `stslist.rep` as produced by the `seqrep` function.
- **cpal**: alternative color palette to use for the states. If user specified, a vector of colors with number of elements equal to the number of states in the alphabet. By default, the ’cpal’ attribute of the `x` object is used.
- **missing.color**: alternative color for representing missing values inside the sequences. By default, this color is taken from the ”missing.color” attribute of the sequence object being plotted.
- **pbarw**: when `TRUE`, the bar heights are set proportional to the number of represented sequences.
- **dmax**: maximal theoretical distance, used for the x axis limits.
- **stats**: if `TRUE` (default), mean discrepancy in each subset defined by all sequences attributed to one representative sequence and the mean distance to this representative sequence are displayed.
- **ylab**: an optional label for the y axis. If set to `NA`, no label is drawn.
- **xaxis**: controls whether a x axis is plotted.
- **xtlab**: optional labels for the x axis ticks labels. If unspecified, the column names of the object being plotted.
- **xtstep**: optional interval at which the tick-marks and labels of the x-axis are displayed. For example, with `xtstep=3` a tick-mark is drawn at position 1, 4, 7, etc... The display of the corresponding labels depends on the available space and is dealt with automatically. If unspecified, the `xtstep` attribute of the `x` object is used.
- **tick.last**: Logical. Should a tick mark be enforced at the last position on the x-axis? If unspecified, the `tick.last` attribute of the `x` object is used.
- **cex.with.axis**: Axis annotation and plotting text and symbols magnification. See `par`.
- **cex.plot**: Deprecated. Use `cex.with.axis` instead.
- **...**: further graphical parameters. For more details about the graphical parameter arguments, see `barplot` and `par`.

Details

This is the plot method for the output produced by the `seqrep` function, i.e. objects of class `stslist.rep`. It produces a plot where the representative sequences are displayed as horizontal bars with width proportional to the number of sequences assigned to them. Sequences are plotted bottom-up according to their representativeness score. Above the plot, two parallel series of symbols associated to each representative are displayed horizontally on a scale ranging from 0 to the maximal theoretical distance $D_{max}$. The location of the
symbol associated to the representative \( r_i \) indicates on axis \( A \) the (pseudo) variance \( V_i \) within the subset of sequences assigned to \( r_i \) and on the axis \( B \) the mean distance \( MD_i \) to the representative.

This method is called by the generic seqplot function (if type="r") that produces more sophisticated plots with group splits and automatic display of the color legend. The seqrplot function is a shortcut for calling seqplot with type="r".

Author(s)
Alexis Gabadinho (with Gilbert Ritschard for the help page)

Examples

```r
## Loading the mvad data set and creating a sequence object
data(mvad)
mvad.labels <- c("employment", "further education", "higher education",
"joblessness", "school", "training")
mvad.scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")

## First 36 months trajectories
mvad.seq <- seqdef(mvad, 15:50, states=mvad.scodes, labels=mvad.labels)

## Computing Hamming distances
##
dist.ham <- seqdist(mvad.seq, method="HAM")

## Extracting a representative set using the sequence frequency
## as a representativeness criterion
mvad.rep <- seqrep(mvad.seq, diss=dist.ham)

## Plotting the representative set
plot(mvad.rep)
```

Description
This is the plot method for output produced by the seqstatd function, i.e for objects of class stslist.statd.

Usage

```r
## S3 method for class 'stslist.statd'
plot(x, type = "d", cpal = NULL, ylab = NULL,
yaxis = TRUE, xaxis = TRUE, xlab = NULL, xtstep = NULL, tick.last = NULL,
cex.axis = 1, space = 0, xlab = NULL, cex.plot, ...)
```
Arguments

- **x**: an object of class `stslist.statd` as produced by the `seqstatd` function.
- **type**: if "d" (default), a state distribution plot is produced. If "Ht" an entropy index plot is produced.
- **cpal**: alternative color palette to be used for the states. If user specified, a vector of colors with number of elements equal to the number of states in the alphabet. By default, the 'cpal' attribute of the x object is used.
- **ylab**: an optional label for the y axis. If set to NA, no label is drawn.
- **yaxis**: if TRUE or "cum", the y axis is plotted with a label showing the cumulated percentage frequency of the displayed sequences. If "pct", the percentage value for each sequence is displayed.
- **xaxis**: if TRUE (default) the x-axis is plotted.
- **xtlab**: optional labels for the ticks of the x-axis. If unspecified, the names attribute of the input x object is used.
- **xtstep**: optional interval at which the tick-marks and labels of the x-axis are displayed. For example, with xtstep=3 a tick-mark is drawn at position 1, 4, 7, etc... The display of the corresponding labels depends on the available space and is dealt with automatically. If unspecified, the xtstep attribute of the x object is used.
- **tick.lsat**: Logical. Should a tick mark be enforced at the last position on the x-axis? If unspecified, the tick.lsat attribute of the x object is used.
- **cex.axis**: Axis annotation magnification. See `par`.
- **space**: the space between the stacked bars. Default is 0, i.e. no space.
- **xlab**: Optional title for the x axis. See `title`.
- **cex.plot**: Deprecated. Use cex.axis instead.
- **...**: further graphical parameters such as border=NA to remove the borders of the bars. For more details about the graphical parameter arguments, see `barplot` and `par`.

Details

This is the plot method for the output produced by the `seqstatd` function, i.e. for objects of class `stslist.statd`. If type="d" it produces a state distribution plot presenting the sequence of the transversal state frequencies at each successive (time) position, as computed by the `seqstatd` function. With type="Ht", the series of entropies of the transversal state distributions is plotted.

This method is called by the generic `seqplot` function (if type="d" or type="Ht") that produces more sophisticated plots, allowing grouping and automatic display of the state color legend. The `seqdplot` and `seqhtplot` functions are shortcuts for calling `seqplot` with type="d" or type="Ht" respectively.

Value

The plotted values, i.e. for type="d" the cross-sectional distributions, and for type="Ht" the cross-sectional entropies.
Examples

```r
## Defining a sequence object with the data in columns 10 to 25
## (family status from age 15 to 30) in the biofam data set
data(biofam)
biofam.lab <- c("Parent", "Left", "Married", "Left+Marr",
"Child", "Left+Child", "Left+Marr+Child", "Divorced")
biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

## State distribution
biofam.statd <- seqstatd(biofam.seq)

## State distribution plot (default type="d" option)
plot(biofam.statd)

## Entropy index plot
plot(biofam.statd, type="Ht")
```

plot.subseqelist  

---

**Description**

Plot frequencies of subsequences.

**Usage**

```r
## S3 method for class 'subseqelist'
plot(x, freq=NULL, cex=1,...)
```

**Arguments**

- `x`  
The subsequences to plot (a `subseqelist` object)
- `freq`  
The frequencies to plot, support if `NULL`
- `cex`  
Plotting text and symbols magnification. See `par`.
- `...`  
arguments passed to `barplot`

**Author(s)**

Matthias Studer (with Gilbert Ritschard for the help page)

**See Also**

`seqefsub`
Examples

```r
## loading data
data(actcal.tse)

## creating sequences
actcal.eseq <- seqecreate(actcal.tse)

## Looking for frequent subsequences
fsubseq <- seqefsub(actcal.eseq, pmin.support=0.01)

## Frequency of first ten subsequences
plot(fsubseq[1:10], cex=2)
plot(fsubseq[1:10])
```

---

**plot.subseqlistchisq**  
*Plot discriminant subsequences*

Description

Plot the result of `seqecmpgroup`

Usage

```r
## S3 method for class 'subseqlistchisq'
plot(x, ylim = "uniform", rows = NA, cols = NA,
     resid.levels = c(0.05, 0.01),
     cpal = brewer.pal(1 + 2 * length(resid.levels), "RdBu"),
     vlegend = NULL,
     cex.legend = 1, ptype = "freq", legend.title = NULL, residlevels, legendcol,
     legend.cex, ...)
```

Arguments

- **x**: The subsequences to plot (a subseqlist object).
- **ylim**: if "uniform" all axes have same limits.
- **rows**: Number of graphic rows
- **cols**: Number of graphic columns
- **resid.levels**: Significance levels used to colorize the Pearson residual
- **cpal**: Color palette used to color the results
- **vlegend**: When TRUE the legend is printed vertically, when FALSE it is printed horizontally. If NULL (default) the best position will be chosen.
- **cex.legend**: Scale parameters for text legend.
- **ptype**: If set to "resid", Pearson residuals are plotted instead of frequencies
- **legend.title**: Legend title.
- **residlevels**: Deprecated. Use resid.levels instead.
Description

This function reads a distance matrix produced by TDA into an R object. When computing OM
distances in TDA, the output is a ‘half’ matrix stored in a text file as a vector.

Usage

read.tda.mdist(file)

Arguments

file the path to the file containing TDA output.

Value

a R matrix containing the distances.
Description
The function provides details about a pairwise alignment.

Usage
```r
seqalign(seqdata, indices, indel=1, sm, with.missing = FALSE)
```

```r
## S3 method for class 'seqalign'
plot(x, cpal = NULL, missing.color = NULL, ylab = NULL,
     yaxis = TRUE, xaxis = TRUE, ytlab = NULL, ylas = 0, xtlab = NULL,
     cex.axis = 1, cex.plot, ...)
```

```r
## S3 method for class 'seqalign'
print(x, digits=3, ...)
```

Arguments
- `seqdata` a state sequence object defined with the `seqdef` function.
- `indices` a vector of length 2 giving the indexes of the two sequences
- `indel` indel cost (see `seqdist`)
- `sm` matrix of substitution costs or a method for computing the costs (see `seqdist`)
- `with.missing` logical: Should the missing state be considered as an element of the alphabet?
- `x` an object of class `seqalign`
- `cpal` color palette
- `missing.color` color for missing elements
- `ylab` y label
- `yaxis` yaxis
- `xaxis` xaxis
- `ytlab` ytlab
- `ylas` ylas
- `xtlab` xtlab
- `cex.axis` Axis annotation magnification. See `par`.
- `digits` number of digits for printed output
- `cex.plot` Deprecated. Use `cex.axis` instead.
- `...` additional arguments passed to other functions

Details
There are print and plot methods for `seqalign` objects.
seqcomp

Value
Object of class seqalign

Author(s)
Alexis Gabadinho (plot.seqalign) and Matthias Studer (seqalign) (with Gilbert Ritschard for the help page)

See Also
seqdist

Examples
data(biofam)
biofam.seq <- seqdef(biofam, 10:25)
costs <- seqsubm(biofam.seq, method="TRATE")
sa <- seqalign(biofam.seq, 1:2, indel=1, sm=costs)
print(sa)
plot(sa)
sa <- seqalign(biofam.seq, c(1,5), indel=0.5, sm=costs)
print(sa)
plot(sa)

seqcomp(x, y)

Arguments
x a state sequence object containing a single sequence (typically the row of a main sequence object, see seqdef)
y a state sequence object containing a single sequence (typically the row of a main sequence object, see seqdef)

Value
TRUE if sequences are identical, FALSE otherwise

See Also
seqfind, seqfpos, seqpm
Examples

```r
data(mvad)
mvad.shortlab <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad.seq <- seqdef(mvad, states=mvad.shortlab, 15:86)

## Comparing sequences 1 and 2 in mvad.seq
seqcomp(mvad.seq[1,],mvad.seq[2,])

## Comparing sequences 176 and 211 in mvad.seq
seqcomp(mvad.seq[176,],mvad.seq[211,])
```

---

### seqconc

**Concatenate vectors of states or events into a character string**

**Description**

Concatenate vectors of states or events into a character string. In the string, each state is separated by `sep`. The void elements in the input sequences are eliminated.

**Usage**

```r
seqconc(data, var=NULL, sep="-", vname="Sequence", void=NA)
```

**Arguments**

- **data**
  - A dataframe or matrix containing sequence data.
- **var**
  - List of the columns containing the sequences. Default is NULL in which case all columns are retained. Whether the sequences are in the compressed (character strings) or extended format is automatically detected by counting the number of columns.
- **sep**
  - Character used as separator. By default, ",-".
- **vname**
  - an optional name for the variable containing the sequences. By default, "Sequence".
- **void**
  - the code used for void elements appearing in the sequences (see Gabadinho et al. (2009) for more details on missing values and void elements in sequences). Default is NA.

**Value**

- a vector of character strings, one for each row in the input data.

**Author(s)**

Alexis Gabadinho
seqcost

References


See Also

seqdecomp.

Examples

data(actcal)
actcal.string <- seqconc(actcal,13:24)
head(actcal.string)

seqcost  Generate substitution and indel costs

Description

The function seqcost proposes different ways to generate substitution costs (supposed to reflect state dissimilarities) and possibly indel costs. Proposed methods are: "CONSTANT" (same cost for all substitutions), "TRATE" (derived from the observed transition rates), "FUTURE" (Chi-squared distance between conditional state distributions lag positions ahead), "FEATURES" (Gower distance between state features), "INDELS", "INDELSLOG" (based on estimated indel costs). The substitution-cost matrix is intended to serve as sm argument in the seqdist function that computes distances between sequences. seqsubm is an alias that returns only the substitution cost matrix, i.e., no indel.

Usage

seqcost(seqdata, method, cval = NULL, with.missing = FALSE, miss.cost = NULL, time.varying = FALSE, weighted = TRUE, transition = "both", lag = 1, miss.cost.fixed = NULL, state.features = NULL, feature.weights = NULL, feature.type = list(), proximities = FALSE)

seqsubm(...)

Arguments

seqdata  A sequence object as returned by the seqdef function.
method  String. How to generate the costs. One of "CONSTANT" (same cost for all substitutions), "TRATE" (derived from the observed transition rates), "FUTURE" (Chi-squared distance between conditional state distributions lag positions ahead), "FEATURES" (Gower distance between state features), "INDELS", "INDELSLOG" (based on estimated indel costs).
cval  Scalar. For method "CONSTANT", the single substitution cost. For method "TRATE", a base value from which transition probabilities are subtracted. If NULL, cval=2 is used, unless transition is "both" and time.varying is TRUE, in which case cval=4.

with.missing  Logical. Should an additional entry be added in the matrix for the missing states? If TRUE, the ‘missing’ state is also added to the state alphabet. Use this if you want to compute distances with gaps (non deleted missing values) inside the sequences. See Gabadinho et al. (2010) for more details on the options for handling missing values when creating state sequence objects.

miss.cost  Scalar or vector. Cost for substituting the missing state. Default is cval.

miss.cost.fixed  Logical. Should the substitution cost for missing be set as the miss.cost value. When NULL (default) it will be set as FALSE when method = "INDELS" or "INDELSLOG", and TRUE otherwise.

time.varying  Logical. If TRUE return an array with a distinct matrix for each time unit. Time is the third dimension (subscript) of the returned array.

weighted  Logical. Should weights in seqdata be used when applicable?

transition  String. Only used if method=“TRATE” and time.varying=TRUE. On which transition are rates based? Should be one of "previous" (from previous state), "next" (to next state) or "both".

lag  Integer. For methods TRATE and FUTURE only. Time ahead to which transition rates are computed (default is lag=1).

state.features  Data frame with features values for each state.

feature.weights  Vector of feature weights with length equal to the number of columns of state.features.

feature.type  List of feature types. See daisy for details.

proximities  Logical: should state proximities be returned instead of substitution costs?

Arguments passed to seqcost

Details

The substitution-cost matrix has dimension ns*ns, where ns is the number of states in the alphabet of the sequence object. The element (i,j) of the matrix is the cost of substituting state i with state j. It defines the dissimilarity between

With method CONSTANT, the substitution costs are all set equal to the cval value, the default value being 2.

With method TRATE (transition rates), the transition probabilities between all pairs of states is first computed (using the seqtrate function). Then, the substitution cost between states i and j is obtained with the formula

$$SC(i, j) = cval - P(i|j) - P(j|i)$$

where $P(i|j)$ is the probability of transition from state j to i lag positions ahead.
With method FUTURE, the cost between \(i\) and \(j\) is the Chi-squared distance between the vector \(d(alphabet|i)\) of probabilities of transition from states \(i\) and \(j\) to all the states in the alphabet \(\text{lag}\) positions ahead:

\[
SC(i, j) = \text{ChiDist}(d(alphabet|i), d(alphabet|j))
\]

With method FEATURES, each state is characterized by the variables state.features, and the cost between \(i\) and \(j\) is computed as the Gower distance between their vectors of state.features values.

With methods INDELS and INDELSLOG, values of indels are first derived from the state relative frequencies \(f_i\). For INDELS, \(\text{indel}_i = 1/f_i\) is used, and for INDELSLOG, \(\text{indel}_i = \log[2/(1 + f_i)]\). Substitution costs are then set as \(SC(i, j) = \text{indel}_i + \text{indel}_j\).

For all methods but INDELS and INDELSLOG, the indel is set as \(\max(sm)/2\) when time.varying=FALSE and as 1 otherwise.

**Value**

For seqcost, a list of two elements, indel and sm or prox:

- **indel**
  - The indel cost. Either a scalar or a vector of size \(ns\).
- **sm**
  - The substitution-cost matrix when proximities = FALSE (default).
- **prox**
  - The state proximity matrix when proximities = TRUE.

\(sm\) and \(prox\) are a matrix of size \(ns \times ns\), where \(ns\) is the number of states in the alphabet of the sequence object.

For seqsubm, only one element, the matrix sm.

**Author(s)**

Gilbert Ritschard and Matthias Studer (and Alexis Gabadinho for first version of seqsubm)

**References**


**See Also**

seqtrate, seqdef, seqdist.
Examples

```r
# Defining a sequence object with columns 10 to 25
# of a subset of the 'biofam' example data set.
data(biofam)
biofam.seq <- seqdef(biofam[501:600,10:25])

# Optimal matching using transition rates based substitution-cost matrix
# and insertion/deletion costs of 3
trcost <- seqcost(biofam.seq, method="TRATE")
biofam.om <- seqdist(biofam.seq, method="OM", indel=3, sm=trcost$sm)

# Using the insertion/deletion cost returned by seqcost
biofam.om <- seqdist(biofam.seq, method="OM", indel=trcost$indel, sm=trcost$sm)

# Using costs based on FUTURE with a forward lag of 4
fucost <- seqcost(biofam.seq, method="FUTURE", lag=4)
biofam.om <- seqdist(biofam.seq, method="OM", indel=fucost$indel, sm=fucost$sm)

# Optimal matching using a unique substitution cost of 2
# and an insertion/deletion cost of 3
ccost <- seqsubm(biofam.seq, method="CONSTANT", cval=2)
biofam.om.c2 <- seqdist(biofam.seq, method="OM", indel=3, sm=ccost)

# Displaying the distance matrix for the first 10 sequences
biofam.om.c2[1:10,1:10]

# Example with weights and missings
# -------------------------------
data(ex1)
ex1.seq <- seqdef(ex1,1:13, weights=ex1$weights)

# Unweighted
subm <- seqcost(ex1.seq, method="TRATE", with.missing=TRUE, weighted=FALSE)
ex1.om <- seqdist(ex1.seq, method="OM", sm=subm$sm, with.missing=TRUE)

# Weighted
subm.w <- seqcost(ex1.seq, method="TRATE", with.missing=TRUE, weighted=TRUE)
ex1.omw <- seqdist(ex1.seq, method="OM", sm=subm.w$sm, with.missing=TRUE)

ex1.om == ex1.omw
```

---

**seqdecomp**

*Convert a character string into a vector of states or events*

**Description**

For the moment, each character in the string will be considered to be one state or event = this function will not give accurate results if the character string representing the sequence contains events or states coded with more than one character.
**seqdef**

Create a state sequence object

**Description**

Create a state sequence object with attributes such as alphabet, color palette and state labels. Most TraMineR functions for state sequences require such a state sequence object as input argument. There are specific methods for plotting, summarizing and printing state sequence objects.

**Usage**

```r
seqdef(data, var=NULL, informat="STS", stsep=NULL, alphabet=NULL, states=NULL, id=NULL, weights=NULL, start=1, left=NA, right="DEL", gaps=NA, missing=NA, void="%", nr="*", cnames=NULL, xtstep=1, tick.last=FALSE, cpal=NULL, missing.color="darkgrey", labels=NULL, ...)
```

**Examples**

```r
## Converts 'seq' into a vector of states of length 10
seq <- "A-A-A-A-B-B-B-C-C-C"
seqdecomp(seq)
```
Arguments

data  
a data frame or matrix containing sequence data.

var  
the list of columns containing the sequences. Default is NULL, i.e. all the columns. The function detects automatically whether the sequences are in the compressed (successive states in a character string) or extended format.

informat  
format of the original data. Default is "STS". Other available formats are: "SPS" and "SPELL", in which case the seqformat function is called to convert the data into the "STS" format (see TraMineR user’s manual (Gabadinho et al., 2010) for a description of these formats). A better solution is nonetheless to convert first your data with seqformat, so as to have better control over the conversion process and visualize the intermediate "STS" formatted data.

stsep  
the character used as separator in the original data if input format is successive states in a character string. If NULL (default value), the seqfcheck function is called for detecting automatically a separator among "-" and ":". Other separators must be specified explicitly.

alphabet  
optional vector containing the alphabet (the list of all possible states). Use this option if some states in the alphabet don’t appear in the data or if you want to reorder the states. The specified vector MUST contain AT LEAST all the states appearing in the data. It may possibly contain additional states not appearing in the data. If NULL, the alphabet is set to the distinct states appearing in the data as returned by the seqstatl function. See details.

states  
an optional vector containing the short state labels. Must have a length equal to the size of the alphabet and the labels must be ordered conformably with alphanumeric ordered values returned by the seqstatl function, or, when alphabet is set, with the thus newly defined alphabet.

id  
optional argument for setting the rownames of the sequence object. If NULL (default), the rownames are taken from the input data. If set to "auto", sequences are numbered from 1 to the number of sequences. A vector of rownames of length equal to the number of sequences may be specified as well.

weights  
optional numerical vector containing weights, which are accounted for by plotting and statistical functions when applicable.

start  
starting time. For instance, if sequences begin at age 15, you can specify 15. At this stage, used only for labelling column names.

left  
the behavior for missing values appearing before the first (leftmost) valid state in each sequence. See Gabadinho et al. (2010) for more details on the options for handling missing values when defining sequence objects. By default, left missing values are treated as ‘real’ missing values and converted to the internal missing value code defined by the nr option. Other options are "DEL" to delete the positions containing missing values or a state code (belonging to the alphabet or not) to replace the missing values.

right  
the behavior for missing values appearing after the last (rightmost) valid state in each sequence. Same options as for the left argument.

gaps  
the behavior for missing values appearing inside the sequences, i.e. after the first (leftmost) valid state and before the last (rightmost) valid state of each sequence. Same options as for the left argument.
seqdef

missing  the code used for missing values in the input data. When specified, all cells containing this value will be replaced by NA's, the internal R code for missing values. If 'missing' is not specified, cells containing NA's are considered as missing values.

void  the internal code used by TraMineR for representing void elements in the sequences. Default is "%". Must be different from left, gaps, and right.

nr  the internal code used by TraMineR for representing real missing elements in the sequences. Default is"*".

cnames  optional names for the columns composing the sequence data. Those names will be used by default in the graphics as axis labels. If NULL (default), names are taken from the original column names in the data.

xtstep  step between displayed tick-marks and labels on the time x-axis of state sequence plots. If not overridden by the user, plotting functions retrieve this parameter from the xtstep attribute of the sequence object. For example, with xtstep=3 a tick-mark is displayed at positions 1, 4, 7, etc... Default value is 1; i.e., a tick mark is displayed at each position. The display of the corresponding labels depends on the available space and is dealt with automatically.

tick.last  Logical. Should a tick mark be enforced at the last position on the time x-axis?

cpal  an optional color palette for representing the states in the graphics. If NULL (default), a color palette is created by calling the brewer.pal function of the RColorBrewer package. If number of states is less or equal than 8, the "Accent" palette is used. If number of states is between 8 and 12, the "Set3" palette is used. If the number of states in the data is greater than 12 you have to specify your own palette. The list of available colors is displayed by the colors function. You can also use alternatively some other palettes from the RColorBrewer package.

missing.color  alternative color for representing missing values inside the sequences. Defaults to "darkgrey".

labels  optional state labels used for the color legend of TraMineR’s graphics. If NULL (default), the state names in the alphabet are used as state labels as well.

...  options passed to the seqformat function for handling input data that is not in STS format.

Details

Applying subscripts to sequence objects (eg. seq[,1:5] or seq[1:10,]) returns a state sequence object with some attributes preserved (alphabet, missing) and some others (start, column names) adapted to the selected column or row subset. If only one column is specified, a factor is returned.

For reordering the states use the alphabet argument. This may for instance be of interest when you want to compare data from different sources with different codings of similar states. Using alphabet permits to order the states conformably in all sequence objects. Otherwise, the default state order is the alpha-numeric order returned by the seqstatl function which may differ when you have different original codings.
Value

An object of class \texttt{stslist}. There are \texttt{print}, \texttt{plot} and \texttt{summary} methods for such objects. State sequence objects are required as argument to other functions such as plotting functions (\texttt{seqdplot}, \texttt{seqiplot} or \texttt{seqfplot}), functions to compute distances (\texttt{seqdist}), etc...

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for help page)

References


See Also

\texttt{plot.stslist} to plot state sequence objects,
\texttt{seqplot} for high level plots of state sequence objects,
\texttt{seqcreate} to create an event sequence object,
\texttt{seqformat} for converting between various longitudinal data formats.

Examples

```r
## Creating a sequence object with the columns 13 to 24
## in the 'actcal' example data set
data(actcal)
actcal.seq <- seqdef(actcal,13:24,
labels=c("37 hours", "19-36 hours", "1-18 hours", "no work"))

## Displaying the first 10 rows of the sequence object
actcal.seq[1:10,]

## Displaying the first 10 rows of the sequence object
## in SPS format
print(actcal.seq[1:10,], format="SPS")

## Plotting the first 10 sequences
plot(actcal.seq)

## Re-ordering the alphabet
actcal.seq <- seqdef(actcal,13:24,alphabet=c("B","A","D","C"))
alphabet(actcal.seq)

## Adding a state not appearing in the data to the
## alphabet
actcal.seq <- seqdef(actcal,13:24,alphabet=c("A","B","C","D","E"))
alphabet(actcal.seq)
```
## Description

The function analyses how the differences between groups of sequences evolve along the positions. It runs a sequence of discrepancy analyses on sliding windows.

## Usage

```r
dist <- seqdiff(seqdata = seqdef, group = c(1, 2), cmprange = c(0, 1),
                seqdist.args = list(method = "LCS", norm = TRUE), with.missing = FALSE,
                weighted = TRUE, squared = FALSE, seqdist_arg = 1)
```

## Arguments

- **seqdata**: A state sequence object created with the `seqdef` function.
- **group**: The group variable.
- **cmprange**: The time range of the sliding window on which subsequences are compared.
seqdist.args List of arguments passed to seqdist for computing the distances.

with.missing Logical. If TRUE, missing values are considered as an additional state. If FALSE subsequences with missing values are removed from the analysis.

weighted Logical. If TRUE, seqdiff uses the weights specified in seqdata.

squared Logical. If TRUE the dissimilarities are squared for computing the discrepancy.

seqdist_arg Deprecated. Use seqdist.args instead.

Details

The function analyses how the part of discrepancy explained by the group variable evolves along the position axis. It runs successively discrepancy analyses within a sliding time-window of range cmprange). At each position, the method uses seqdist to compute a distance matrix over the time-window and then derives the explained discrepancy on that window with dissassoc.

There are print and plot methods for the returned value.

Value

A seqdiff object, with the following items:

stat A data.frame with three statistics (PseudoF, PseudoR2 and PseudoT) for each time stamp of the sequence, see dissassoc.

discrepancy A data.frame with, at each time stamp, the discrepancy within each group defined by the group variable and for the whole population.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

dissassoc to analyse the association of the group variable with the whole sequence
### Examples

```r
## Define a state sequence object
data(mvad)
## First 12 months of first 200 trajectories
mvad.seq <- seqdef(mvad[1:200, 17:28])

## Position-wise discrepancy analysis
mvad.diff <- seqdiff(mvad.seq, group=mvad$gcseSeq[1:200])
print(mvad.diff)
plot(mvad.diff, stat=c("Pseudo R2", "Levene"), xtstep=6)
plot(mvad.diff, stat="discrepancy")
```

---

### seqdim

**Dimension of a set of sequences**

### Description

Returns the number of sequences (rows) and the maximum length of a set of sequences.

### Usage

```r
seqdim(seqdata)
```

### Arguments

- `seqdata` a set of sequences.

### Details

The function will first search for separators '-' or ':' in the sequences in order to detect whether they are in the compressed or extended format.

### Value

- a vector with the number of sequences and the maximum sequence length.

### Author(s)

Alexis Gabadinho
**seqdist**

**Distances (dissimilarities) between sequences**

**Description**

Computes pairwise dissimilarities between sequences or dissimilarity from a reference sequence. Several dissimilarity measures can be chosen, including optimal matching (OM) and many of its variants, distance based on the count of common attributes, and distances between sequence state distributions.

**Usage**

```
seqdist(seqdata, method, refseq = NULL, norm = "none", indel = 1.0, sm = NULL,
with.missing = FALSE, full.matrix = TRUE, kweights = rep(1.0, ncol(seqdata)),
tpow = 1.0, expcost = 0.5, context, link = "mean", h = 0.5, nu,
transindel = "constant", otto, previous = FALSE, add.column = TRUE,
breaks = NULL, step = 1, overlap = FALSE, weighted = TRUE,
global.pdotj = NULL, prox = NULL)
```

**Arguments**

- **seqdata**: State Sequence Object. The sequence data to use. It can be created with the `seqdef` function.
- **method**: String. The dissimilarity measure to use. It can be "OM", "OMloc", "OMslen", "OMspell", "OMstran", "HAM", "DHD", "CHI2", "EUCLID", "LCS", "LCP", "RLCP", "NMS", "NMSMST", "SVRspell", or "TWED". See the Details section.
- **refseq**: NULL, Integer, or State Sequence Object. Default: NULL. The baseline sequence to compute the distances from.
  The most frequent sequence (\(\emptyset\)) or a sequence in seqdata at a specified index (strictly greater than \(\emptyset\)) when an integer and method is one "OM", "OMloc", "OMslen", "OMspell", "OMstran", "HAM", "DHD", "LCS", "LCP", "RLCP", "NMS", "NMSMST", "SVRspell", or "TWED".
  An external sequence when a state sequence object and method is one of "OM", "HAM", "DHD", "LCS", "LCP", or "RLCP". It must have a single row and the same alphabet as seqdata.
- **norm**: String. Default: "none". The normalization to use when method is one of "OM", "HAM", "DHD", "LCS", "LCP", "RLCP", "CHI2", "EUCLID", or "NMS", "NMSMST", "SVRspell", or "TWED".
  It can be "none", "auto", or, except for "CHI2" and "EUCLID", "maxlength", "gmean", "maxdist", or "YujiianBo". "auto" is equivalent to "maxlength" when method is one of "OM", "HAM", or "DHD", or "gmean" when method is one of "LCS", "LCP", or "RLCP". See the Details section.
- **indel**: Double or Vector of Doubles. Default: 1.0. Insertion/deletion cost(s).
  The single state-independent insertion/deletion cost when a double and method is one of "OM", "OMslen", "OMspell", "OMstran", or "TWED".
The state-dependent insertion/deletion costs when a vector of doubles and method = "OM" or method = "OMstran". It contains an indel cost for each state in the same order as the alphabet.

sm

NULL, Matrix, Array, or String. Substitution costs. Default: NULL.

The substitution-cost matrix when a matrix and method is one of "OM", "OMloc", "OMslen", "OMspell", "OMstran", "HAM", or "TWED". It contains an indel cost for each state in the same order as the alphabet.

The series of the substitution-cost matrices when an array and method = "DHD". They are grouped in a 3-dimensional array with the third index referring to the position in the sequence.

The name of a seqcost method when a string and method is one of "OM", "OMloc", "OMslen", "OMspell", "OMstran", "HAM", or "TWED". The method is used to build sm. It can be "INDELS" or "INDELSLOG" for "OM", "OMloc", "OMslen", "OMspell", "OMstran", "HAM", and "TWED", "CONSTANT" for "OM" and "HAM", "TRATE" for "OM", "HAM", and "DHD".

sm is mandatory when method is one of "OM", "OMloc", "OMslen", "OMspell", "OMstran", or "TWED".

sm is autogenerated when method is one of "HAM" or "DHD" and sm = NULL. See the Details section.

Note: With method = "NMS" or method = "SVRspell", see prox instead.

with.missing

Logical. Default: FALSE. When method isn't "OMslen" or "OMstran", should the non-deleted gap (missing value) be added to the alphabet as an additional state? If FALSE and seqdata or refseq contains such gaps, an error is raised.

full.matrix

Logical. Default: TRUE. When refseq = NULL, if TRUE, the full distance matrix is returned, if FALSE, an object of class dist is returned, that is, a vector containing only values from the upper triangle of the distance matrix. Objects of class dist are smaller and can be passed directly as arguments to most clustering functions.

kweights

Vector of Doubles. Default: vector of 1.0. The weights applied to subsequences when method is one of "NMS", "NMSMST", or "SVRspell". It contains at position k the weight applied to the subsequences of length k. It must be positive. Its length must be equal to the number of columns of seqdata.

tpow

Double. Default: 1.0. The exponential weight of spell length when method is one of "OMspell", "NMSMST", or "SVRspell".

expcost

Double. Default: 0.5. The cost of spell length transformation when method = "OMloc" or method = "OMspell". It must be positive. The exact interpretation is distance-dependent.

context

Double. Default: 1-2*expcost. The cost of local insertion when method = "OMloc". It must be positive.

link

String. Default: "mean". The function used to compute substitution costs when method = "OMslen". One of "mean" (arithmetic average) or "gmean" (geometric mean as in the original proposition of Halpin 2010).

h

Double. Default: 0.5. It must be greater than or equal to 0.

The exponential weight of spell length when method = "OMslen".

The gap penalty when method = "TWED". It corresponds to the lambda in Halpin (2014), p 88.
nu Double. Stiffness when method = "TWED". It must be strictly greater than 0. See Halpin (2014), p 88.

transindel String. Default: "constant". Method for computing transition indel costs when method = "OMstran". One of "constant" (single indel of 1.0), "subcost" (based on substitution costs), or "prob" (based on transition probabilities).

otto Double. The origin-transition trade-off weight when method = "OMstran". It must be in [0, 1].

previous Logical. Default: FALSE. When method = "OMstran", should we also account for the transition from the previous state?

add.column Logical. Default: TRUE. When method = "OMstran", should the last column (and also the first column when previous = TRUE) be duplicated?

breaks NULL, List of pairs Integers. Default: NULL. The list of the possibly overlapping intervals when method = "CHI2" or method = "EUCLID".

step Integer. Default: 1. The length of the intervals when method = "CHI2" or method = "EUCLID" and breaks = NULL. It must be positive. It must also be even when overlap = TRUE.

overlap Logical. Default: FALSE. When method = "CHI2" or method = "EUCLID" and breaks = NULL, should the intervals overlap?

weighted Logical. Default: TRUE. When method is "CHI2", should the distributions of the states account for the sequence weights in seqdata? See seqdef.

global.pdotj Numerical vector, "obs", or NULL. Default: NULL. Only for method = "CHI2". The vector of state proportions to be used as marginal distribution. When NULL, the state distribution on the corresponding interval is used. When "obs", the overall state distribution in seqdata is used for all intervals. When a vector of proportions, it is used as marginal distribution for all intervals.

prox NULL or Matrix. Default: NULL. The matrix of state proximities when method = "NMS" or method = "SVRspell".

Details

The seqdist function returns a matrix of distances between sequences or a vector of distances from the reference sequence when refseq is set. The available metrics (see method option) include:

- **Edit distances**: optimal matching ("OM"), localized OM ("OMloc"), spell-length-sensitive OM ("OMslen"), OM of spell sequences ("OMspell"), OM of transition sequences ("OMstran"), Hamming ("HAM"), dynamic Hamming ("DHD"), and the time warp edit distance ("TWED").

- **Metrics based on counts of common attributes**: distance based on the longest common subsequence ("LCS"), on the longest common prefix ("LCP"), on the longest common suffix ("RLCP"), on the number of matching subsequences ("NMS"), on the number of matching subsequences weighted by the minimum shared time ("NMSMST") and, the subsequence vectorial representation distance ("SVRspell").

- **Distances between state distributions**: Euclidean ("EUCLID"), Chi-squared ("CHI2").

See Studer and Ritschard (2014, 2016) for a description and the comparison of the above dissimilarity measures except "TWED" for which we refer to Marteau (2009) and Halpin (2014).

Each method can be controlled with the following parameters:
method | parameters
---|---
OM | sm, indel, norm, refseq
OMloc | sm, expcost, context, refseq
OMslen | sm, indel, link, h, refseq
OMspell | sm, indel, tpow, expcost, refseq
OMstran | sm, indel, transindel, otto, previous, add.column
HAM, DHD | sm, norm, refseq
CHI2 | breaks, step, overlap, norm, weighted, global.pdotj
EUCLID | breaks, step, overlap, norm
LCS, LCP, RLCP | norm, refseq
NMS | prox, kweights, refseq
NMSMST | kweights, tpow, refseq
SVRspell | prox, kweights, tpow, refseq
TWED | sm, indel, h, nu, refseq

"LCS" is "OM" with a substitution cost of 2 (sm = "CONSTANT", cval = 2) and an indel of 1.0. "HAM" is "OM" without indels. "DHD" is "HAM" with specific substitution costs at each position.

"HAM" and "DHD" apply only to sequences of equal length.

When sm = NULL, the substitution-cost matrix is automatically created for "HAM" with a single substitution cost of 1 and for "DHD" with the costs derived from the transition rates at the successive positions.

Some distances can optionally be normalized by means of the norm argument. If set to "auto", Elzinga’s normalization (similarity divided by geometrical mean of the two sequence lengths) is applied to "LCS", "LCP" and "RLCP" distances, while Abbott’s normalization (distance divided by length of the longer sequence) is used for "OM", "HAM" and "DHD". Elzinga’s method can be forced with "gmean" and Abbott’s rule with "maxlength". With "maxdist" the distance is normalized by its maximal possible value. For more details, see Gabadinho et al. (2009, 2011). Finally, "YujianBo" is the normalization proposed by Yujian and Bo (2007) that preserves the triangle inequality. The square of the "CHI2" and "EUCLID" distances are normalized by the number of intervals and by the maximal distance on each interval. Note that for 'CHI2' the maximal distance on each interval depends on the state distribution on the interval.

When sequences contain gaps and the gaps = NA option was passed to seqdef (i.e. when there are non deleted missing values), the with.missing argument should be set as TRUE. If left as FALSE the function stops when it encounters a gap. This is to make the user aware that there are gaps in the sequences. For methods that need an sm value, seqdist expects a substitution-cost matrix with a row and a column entry for the missing state (symbol defined with the nr option of seqdef). Substitution-cost matrices returned by seqcost (and so seqsubm) include these additional entries when the function is called with with.missing = TRUE. More details on how to compute distances with sequences containing gaps can be found in Gabadinho et al. (2009).

**Value**

When refseq is NULL (default), the whole matrix of pairwise distances between sequences or, if full.matrix = FALSE, the corresponding dist object of pairwise distances between sequences is returned. Otherwise, a vector with distances between the sequences in the state sequence object and the reference sequence specified with refseq is returned.
Author(s)
Matthias Studer, Pierre-Alexandre Fonta, Alexis Gabadinho, Nicolas S. Müller, Gilbert Ritschard.

References
Gabadinho, A., G. Ritschard, M. Studer and N. S. Müller (2009). Mining Sequence Data in R with the TraMineR package: A user’s guide Department of Econometrics and Laboratory of Demography, University of Geneva
See also all references in Studer and Ritschard (2014, 2016)

See Also
seqcost, seqsubm, seqdef, and for multichannel distances seqdistmc.

Examples
## -------------------------------
## Example without missings
## -------------------------------

## Defining a sequence object with columns 10 to 25 of a
## subset of the 'biofam' data set
data(biofam)
biofam.seq <- seqdef(biofam[501:600, 10:25])

## OM distances with a substitution-cost matrix derived
## from transition rates
biofam.om <- seqdist(biofam.seq, method = "OM", indel = 3,
                      sm = "TRATE")

## OM distances using the vector of estimated indels and
## substitution costs derived from the estimated indels
```r
costs <- seqcost(biofam.seq, method = "INDELSLOG")
biofam.om <- seqdist(biofam.seq, method = "OM",
    indel = costs$indel, sm = costs$sm)

## Normalized LCP distances
biofam.lcp.n <- seqdist(biofam.seq, method = "LCP",
    norm = "auto")

## Normalized LCS distances to the most frequent sequence
biofam.dref1 <- seqdist(biofam.seq, method = "LCS",
    refseq = 0, norm = "auto")

## LCS distances to an external sequence
ref <- seqdef(as.matrix("(0,5)\-(3,5)\-(4,6)"), informat = "SPS",
    alphabet = alphabet(biofam.seq))

biofam.dref2 <- seqdist(biofam.seq, method = "LCS",
    refseq = ref)

## Chi-squared distance over the full observed timeframe
biofam.chi.full <- seqdist(biofam.seq, method = "CHI2",
    step = max(seqlength(biofam.seq)))

## Chi-squared distance over successive overlapping intervals of length 4
biofam.chi.ostep <- seqdist(biofam.seq, method = "CHI2",
    step = 4, overlap = TRUE)

# Example with missings
# Example with missings

data(ex1)
ex1.seq <- seqdef(ex1[, 1:13])

## OM with substitution costs based on transition probabilities and indel set as half the maximum
## substitution cost
costs.tr <- seqcost(ex1.seq, method = "TRATE",
    with.missing = TRUE)

biofam.om <- seqdist(ex1.seq, method = "OM",
    indel = costs.tr$indel, sm = costs.tr$sm,
    with.missing = TRUE)

## Localized OM

biofam.omloc <- seqdist(ex1.seq, method = "OMloc",
    indel = costs.tr$indel, sm = costs.tr$sm,
    with.missing = TRUE)

## OM of spells

biofam.omspell <- seqdist(ex1.seq, method = "OMspell",
    sm = costs.tr$sm, indel = costs.tr$indel,
    with.missing = TRUE)
```
seqdistmc

Multichannel distances between sequences

Description

Compute multichannel pairwise optimal matching (OM) distances between sequences by deriving the substitution costs from the costs of the single channels. Works with OM and its following variants: distance based on longest common subsequence (LCS), Hamming distance (HAM), and Dynamic Hamming distance (DHD).

Usage

```r
seqdistmc(channels, method, norm="none", indel=1, sm=NULL,
           with.missing=FALSE, full.matrix=TRUE, link="sum", cval=2,
           miss.cost=2, cweight=0)
```

Arguments

- **channels**: A list of state sequence objects defined with the `seqdef` function, each state sequence object corresponding to a "channel".
- **method**: A character string indicating the metric to be used. One of "OM" (Optimal Matching), "LCS" (Longest Common Subsequence), "HAM" (Hamming distance), "DHD" (Dynamic Hamming distance).
- **norm**: String. Default: "none". The normalization method to use. See `seqdist`.
- **indel**: A vector with an insertion/deletion cost for each channel (OM method).
- **sm**: A list with a substitution-cost matrix for each channel (OM, HAM and DHD method) or a list of method names for generating the substitution-costs (see `seqsubm`).
- **with.missing**: Must be set to TRUE when sequences contain non deleted gaps (missing values) or when channels are of different length. See details.
- **full.matrix**: If TRUE (default), the full distance matrix is returned. If FALSE, an object of class `dist` is returned.
- **link**: One of "sum" or "mean". Method to compute the "link" between channels. Default is to sum the substitution costs.
- **cval**: Substitution cost for "CONSTANT" matrix, see `seqsubm`.
- **miss.cost**: Missing values substitution cost, see `seqsubm`.
- **cweight**: A vector of channel weights. Default is 1 (same weight for each channel).
Details

The seqdistmc function returns a matrix of multichannel distances between sequences. The available metrics (see 'method' option) are optimal matching ("OM"), longest common subsequence ("LCS"), Hamming distance ("HAM") and Dynamic Hamming Distance ("DHD"). See seqdist for more information about distances between sequences.

The seqdistmc function computes a multichannel distance in two steps following the strategy proposed by Pollock (2007). First it builds a new sequence object derived from the combination of the sequences of each channel. Second, it derives the substitution cost matrix by summing (or averaging) the costs of substitution across channels. It then calls seqdist to compute the distance matrix.

Normalization may be useful when dealing with sequences that are not all of the same length. For details on the applied normalization, see seqdist.

Value

A matrix of pairwise distances between multichannel sequences is returned.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

seqsubm, seqdef, seqdist.

Examples

data(biofam)

## Building one channel per type of event left, children or married
bf <- as.matrix(biofam[, 10:25])
children <- bf==4 | bf==5 | bf==6
married <- bf == 2 | bf== 3 | bf==6
left <- bf==1 | bf==3 | bf==5 | bf==6

## Building sequence objects
child.seq <- seqdef(children)
marr.seq <- seqdef(married)
left.seq <- seqdef(left)

## Using transition rates to compute substitution costs on each channel
mcdist <- seqdistmc(channels=list(child.seq, marr.seq, left.seq),
method="OM", sm=list("TRATE", "TRATE", "TRATE"))
## seqdss

### Extract distinct states sequence from a sequence object

**Description**

Extract distinct states sequence from a sequence object.

**Usage**

```r
seqdss(seqdata, with.missing=FALSE)
```

**Arguments**

- `seqdata`: a sequence object as defined by the `seqdef` function.
- `with.missing`: if set to `TRUE`, missing statuses (gaps in sequences) also appear in the DSS. See `seqdef` on options for handling missing values when creating sequence objects.

**Details**

Returns a sequence object containing the distinct states sequences, i.e., the durations are not taken into account. The DSS contained in `'D=D=D=D=A=A=A=A=A=A=A=A'` is `'D=A'`. Associated durations can be extracted with the `seqdur` function.

If called with the `{with.missing=TRUE}` argument, a missing state in a sequence is considered as the occurrence of an additional symbol of the alphabet, and two or more consecutive missing states are considered as two or more occurrences of the same state. Hence the DSS of `A=A+*+*+*+*+*+B+B+C+C+D` is `A+*+B+C+D`.

**Value**

A sequence object containing the distinct state sequence (DSS) for each sequence in the object given as argument.

**Author(s)**

Alexis Gabadinho

**See Also**

- `seqdur`
**seqdur**

*Extract state durations from a sequence object.*

**Description**


**Usage**

`seqdur(seqdata, with.missing=FALSE)`

**Arguments**

- **seqdata**
  a sequence object as defined by the *seqdef* function.

- **with.missing**
  if set to TRUE, durations are also computed for missing statuses (gaps in sequences). See *seqdef* on options for handling missing values when creating sequence objects.

**Value**

a matrix containing the states durations for each distinct state in each sequence.

**Author(s)**

Alexis Gabadinho
seqeapplysub

Checking for the presence of given event subsequences

Description

Checks occurrences of the subsequences subseq among the event sequences and returns the result according to the selected method.

Usage

seqeapplysub(subseq, method = NULL, constraint = NULL, rules=FALSE)

Arguments

- subseq: list of subsequences (an event subsequence object) such as created by seqefsub
- method: type of result, should be one of "count", "presence" or "age"
- constraint: Time constraints overriding those used to compute subseq. See seqeconstraint
- rules: If set to TRUE, instead of checking occurrences of the subsequences among the event sequences, check the occurrence of the subsequences inside the subsequences (internally used by seqerules)

Details

There are three methods implemented: "count" counts the number of occurrence of each given subsequence in each event sequence; "presence" returns 1 if the subsequence is present, 0 otherwise; "age" returns the age of appearance of each subsequence in each event sequence. In case of multiple possibilities, the age of the first occurrence is returned. When the subsequence is not in the sequence, -1 is returned.
Value

The return value is a matrix where each row corresponds to a sequence (row names are set accordingly) and each column corresponds to a subsequence (col names are set accordingly). The cells of the matrix contain the requested values (count, presence-absence indicator or age).

Author(s)

Matthias Studer and Reto Bürgin (alternative counting methods) (with Gilbert Ritschard for the help page)

References


See Also

seqecreate for more information on event sequence object and Gabadinho et al. (2009) on how to use the event sequence analysis module.

Examples

```r
## Loading data
data(actcal.tse)

## Creating the event sequence object
actcal.eseq <- seqecreate(actcal.tse)

## Printing sequences
actcal.eseq[1:10]

## Looking for frequent subsequences
fsubseq <- seqefsub(actcal.eseq,pmin.support=0.01)

## Counting the number of occurrences of each subsequence
msubcount <- seq eapplysub(fsubseq,method="count")
## First lines...
msubcount[1:10,1:10]

## Presence-absence of each subsequence
msubpres <- seq eapplysub(fsubseq,method="presence")
## First lines...
msubpres[1:10,1:10]

## Age at first appearance of each subsequence
msubage <- seq eapplysub(fsubseq,method="age")
## First lines...
msubage[1:10,1:10]
```
seqecmpgroup

Identifying discriminating subsequences

Description

Identify and sort the most discriminating subsequences by their discriminating power.

Usage

seqecmpgroup(subseq, group, method="chisq", pvalue.limit=NULL, weighted = TRUE)

Arguments

subseq A subseqelist object (list of subsequences) such as produced by seqefsub

group Group membership, i.e., a variable or factor defining the groups which we want to discriminate

method The discrimination method; one of "bonferroni" or "chisq"

pvalue.limit Can be used to filter the results. Only subsequences with a p-value lower than this parameter are selected. If NULL all subsequences are returned (regardless of their p-values).

weighted Logical. If TRUE, seqecmpgroup uses the weights specified in subseq, (see seqefsub).

Details

The following discrimination test functions are implemented: chisq, the Pearson Independence Chi-squared test, and bonferroni, the Pearson Independence Chi-squared test with Bonferroni correction.

Value

An objet of type subseqelistchisq (subtype of subseqelist) with the following elements

subseq Sorted list of found discriminating subsequences
eseq The event sequence object on which the tests were computed
constraint Time constraints used for searching the subsequences (see seqeconstraint)
labels Levels (value labels) of the target group variable
type Type of test used
data A data frame with columns support, index (original rank of the subsequence, i.e., its position in the inputted subseq) and a pair of frequency and Pearson residual columns for each group

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)
References


See Also

See also plot.subseqlistchisq to plot the results

Examples

data(actcal.tse)
actcal.eseq <- seqecreate(actcal.tse)

#Searching for frequent subsequences, that is, appearing at least 20 times
fsubseq <- seqefsub(actcal.eseq, pmin.support=0.01)

#Searching for subsequences discriminating the most men and women
data(actcal)
discr <- seqecmpgroup(fsubseq, group=actcal$sex, method="bonferroni")

#Printing the six most discriminating subsequences
print(discr[1:6])

#Plotting the six most discriminating subsequences
plot(discr[1:6])

seteconstraint

Setting time constraints and the counting method

Description

Function used to set time constraints and the counting method in methods (seqe...) for event sequences such as seqefsub for searching frequent subsequences or seqeapplysub for checking occurrences of subsequences.

Usage

seqeconstraint(max.gap = -1, window.size = -1, age.min = -1, age.max = -1, age.max.end = -1, count.method = 1, maxGap, windowSize, ageMin, ageMax, ageMaxEnd, countMethod)
Arguments

max.gap  The maximum time gap between two events
window.size  The maximum time span accepted for subsequences
age.min  Minimal start time position allowed for subsequences. Ignored when equal to -1 (default).
age.max  Maximal start time position allowed for subsequences. Ignored when equal to -1 (default).
age.max.end  Maximal end time position allowed for subsequences. Ignored when equal to -1 (default).
count.method  By default, subsequences are counted only one time by sequence ('COBJ' method). Alternative counting methods are 'CDIST_O', 'CWIN', 'CMINWIN' or 'CDIST' respectively. See details.

maxGap  Deprecated. Use max.gap instead.
WindowSize  Deprecated. Use window.size instead.
ageMin  Deprecated. Use age.min instead.
ageMax  Deprecated. Use age.max instead.
ageMaxEnd  Deprecated. Use age.max.end instead.
countMethod  Deprecated. Use count.method instead.

Details

max.gap, window.size, age.min, age.max and age.max.end. If so, two events should not be separated by more than max.gap and the whole subsequence should not exceed a window.size time span. The other parameters specify the start and end age of the subsequence, it should start between age.min and age.max and finish before age.max.end. Parameters age.min, age.max and age.max.end are interpreted as the number of positions (time units) from the beginning of the sequence.

There are 5 options for the count.method argument. (1) By default, the count is the number of sequences that contain the subsequence ('COBJ' method). Alternatives are (2) "CDIST_O" (counts all distinct occurrences in each sequence including possibly overlapping occurrences, i.e., occurrences sharing a same event occurrence), (3) "CWIN" (number of slidden windows of length window.size that contain an occurrence of the subsequence), (4) "CMINWIN" (number of minimal windows of occurrence) and (5) "CDIST" (distinct occurrences without event occurrences overlap). See references.

Value

A constraint object containing one item per constraint type.

Author(s)

Matthias Studer, Nicolas S. Müller and Reto Bürgin (alternative counting methods) (with Gilbert Ritschard for the help page)
References


See Also

seqefsub, seqeapplysub

seqecontain

Check if sequence contains events

Description

Check if an event sequence or subsequence contains given events

Usage

seqecontain(eseq, event.list, unknown.exclude = FALSE, seq, eventlist, exclude)

Arguments

eseq A event sequence object (seqelist) or a an event subsequence object (subseqelist)

event.list A list of events

unknown.exclude if TRUE the search is exclusive and returns FALSE for any subsequence containing an event that is not in event.list

seq Deprecated. Use eseq instead.

eventList Deprecated. Use event.list instead.

exclude Deprecated. Use unknown.exclude instead.

Details

Checks, for each provided event sequence, if it contains one of the events in event.list. If unknown.exclude is TRUE, seqecontain looks if all events of the subsequence are in event.list.

Value

A logical vector.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)
seqecreate

See Also

seqecreate for creating event sequence objects and seqefsub for creating event subsequence objects.

Examples

data(actcal.tse)
actcal.eseq <- seqecreate(actcal.tse)

# Searching for frequent subsequences, that is appearing at least 20 times
fsubseq <- seqefsub(actcal.eseq, min.support = 20)

# looking for subsequence with FullTime
seqecontain(fsubseq, c("FullTime"))

seqecreate

Create event sequence objects.

Description

Create an event sequence object either from time stamped events or from a state sequence object.

Usage

seqecreate(data = NULL, id = NULL, timestamp = NULL, event = NULL, end.event = NULL, tevent = "transition", use.labels = TRUE, weighted = TRUE, endEvent)

Arguments

data A state sequence object (see seqdef) or a data frame
id The sequence 'id' (integer) column when data are provided in TSE format (ignored if data argument is provided).
timestamp The event 'timestamp' (double) column when data are provided in TSE format, i.e., the time at which events occur (ignored if data argument is provided).
event The 'event' column when data are provided in TSE format, i.e., the events occurring at the specified time stamps (ignored if data argument is provided).
end.event If specified this event indicates the end of observation time (total length of event sequences) when it is not followed by any other valid event. The event is ignored when occurring in between two valid events.
tevent Either a transition matrix or a method to generate events from state sequences (see seqetm). Used only when data is a state sequence object.
use.labels If TRUE, transitions names are built from long state labels rather than from the short state names of the alphabet.
weighted If TRUE and data is a state sequence object, use the weights specified in data (see `seqdef`)
endEvent Deprecated. Use end.event instead.

Details

There are several ways to create an event sequence object. The first one is by providing the events in TSE format (see `seqformat`), i.e. by providing three paired lists: id, timestamp and event, such that each triplet (id, timestamp, event) defines the event that occurs at time timestamp for case id. Several events at the same time for a same id are allowed. The lists can be provided with the arguments id, timestamp and event. An alternative is by providing a data frame as data argument in which case the function takes the required information from the "id", "timestamp" and "event" columns of that data frame. In any case with TSE format, listed events should be grouped by id and an error will be thrown otherwise. Such grouping can be achieved by ordering the data according to the id column using the `order` function (e.g., `data[order(data$id), ]`).

The other way is to pass a state sequence object (as data argument) and to perform an automatic state-to-event conversion. The simplest way to make a conversion is by means of a predefined method (see `seqetm`), such as "transition" (one distinct event per possible transition), "state" (a new event for each entering in a new state) and "period" (a pair of events, one start-state event and one end-state event for each found transition). For a more customized conversion, you can specify a transition matrix in the same way as in `seqformat`. Function `seqetm` can help you in creating your transition matrix.

Event sequence objects as created by `seqecreate` are required by most other `seqe` methods, such as `seqefsub` or `seqeapplysub` for example.

Value

An object of class `seqelist`. There are print, and plot methods for such objects.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

`seqformat` for converting between sequence formats, `seqeweight` for retrieving or assigning weights, `seqefsub` for searching frequent subsequences, `seqecmpgroup` to search for discriminant subsequences, `seqeapplysub` for counting subsequence occurrences, `seqelength` for information about length (observation time) of event sequences, `seqdef` to create a state sequence object.
Examples

```r
## Starting with states sequences
## Loading data
data(biofam)
## Creating state sequences
biofam.seq <- seqdef(biofam, 10:25, informat='STS')
## Creating event sequences from biofam
biofam.eseq <- seqecreate(biofam.seq)

## Loading data
data(actcal.tse)
## Creating sequences
actcal.eseq <- seqecreate(id=actcal.tse$id, timestamp=actcal.tse$timestamp, event=actcal.tse$event)
## Printing sequences
actcal.eseq[1:10]
## Using the data argument
actcal.eseq <- seqecreate(data=actcal.tse)

## Example with misses
data(ex1) ## STS data with missing values

## Creating the state sequence object with by default
## the left missings and gaps coded as 'x' and
## end missings coded as void ('%')
sqex1 <- seqdef(ex1[,1:13])
## and without ignoring right missings (coded as 'x')
sqex1b <- seqdef(ex1[,1:13], right=NA)

## Compare the outcome
seqecreate(sqex1)
seqecreate(sqex1, tevent='state')
seqecreate(sqex1, tevent='state', end.event=attr(sqex1,'void'))
seqecreate(sqex1b, tevent='state')
```

Description

Returns the list of subsequences with minimal support sorted in decreasing order of support. Various
time constraints can be set to restrict the search to specific time periods or subsequence durations.
The function permits also to get information on specified subsequences.

Usage

```r
seqefsub(eseq, str.subseq = NULL, min.support = NULL, pmin.support = NULL, constraint = seqeconstraint(), max.k = -1, weighted = TRUE, seq, strsubseq, minSupport, pMinSupport, maxK)
```
Arguments
eseq
- A list of event sequences
str.subseq
- A list of specific subsequences to look for. See details.
min.support
- The minimum support (in number of sequences)
pmin.support
- The minimum support (in percentage, corresponding count will be rounded)
constraint
- A time constraint object as returned by seqeconstraint
max.k
- The maximum number of events allowed in a subsequence
weighted
- Logical. Should seqefsub use the weights specified in eseq (see seqeweight).
seq
- Deprecated. Use eseq instead.
str.subseq
- Deprecated. Use str.subseq instead.
minSupport
- Deprecated. Use min.support instead.
pMinSupport
- Deprecated. Use pmin.support instead.
maxK
- Deprecated. Use max.k instead.

Details

There are two usages of this function. The first is for searching subsequences satisfying a support condition. By default, the support is counted per sequence and not per occurrence, i.e. when a sequence contains several occurrences of a same subsequence it is counted only once. Use the count.method argument of seqeconstraint to change that. The minimal required support can be set with pmin.support as a proportion (between 0 and 1) in which case the support will be rounded, or through min.support as a number of sequences. Time constraints can also be imposed with the constraint argument, which must be the outcome of a call to the seqeconstraint function.

The second possibility is for searching sequences that contain specified subsequences. This is done by passing the list of subsequences with the str.subseq argument. The subsequences must contain only events from the alphabet of events of eseq and must be in the same format as that used to display subsequences (see str.seqelist). Each transition (group of events) should be enclosed in parentheses () and separated with commas, and the succession of transitions should be denoted by a '-' indicating a time gap. For instance "(FullTime)-(PartTime, Children)" stands for the subsequence "FullTime" followed by the transition defined by the two simultaneously occurring events "PartTime" and "Children".

To get information such as the number of occurrences of the subsequences returned by seqefsub or the sequences that contain each subsequence use the function seqeapplysub.

Subsets of the returned subseqelist can be accessed with the [] operator (see example). There are print and plot methods for subseqelist.

Value

A subseqelist object with at least the following attributes:
eseq
- The list of sequences in which the subsequences were searched (a seqelist event sequence object).
subseq
- A list of subsequences (a seqelist event sequence object).
data
- A data frame containing details (support, frequency, ...) about the subsequences
constraint  The constraint object used when searching the subsequences.

type     The type of search: 'frequent' or 'user'

Author(s)
Matthias Studer and Reto Bürgin (alternative counting methods) (with Gilbert Ritschard for the help page)

References

See Also
See plot.subseqelist to plot the result. See seqecreate for creating event sequences. See seqeapplysub to count the number of occurrences of frequent subsequences in each sequence. See is.seqelist about seqelist.

Examples

data(actcal.tse)
actcal.eseq <- seqecreate(actcal.tse)

## Searching for subsequences appearing at least 20 times
fsubseq <- seqefsub(actcal.eseq, min.support=20)

## The same using a percentage
fsubseq <- seqefsub(actcal.eseq, pmin.support=0.01)

## Getting a string representation of subsequences
## First ten most frequent subsequences
fsubseq[1:10]

## Using time constraints
## Looking for subsequences starting in Summer (between June and September)
fsubseq <- seqefsub(actcal.eseq, min.support=10,
                      constraint=seqeconstraint(age.min=6, age.max=9))
fsubseq[1:10]

## Looking for subsequences occurring in Summer (between June and September)
fsubseq <- seqefsub(actcal.eseq, min.support = 10,
                      constraint=seqeconstraint(age.min=6, age.max=9, age.max.end=9))
fsubseq[1:10]

## Looking for subsequence enclosed in a 6 month period
## and with a maximum gap of 2 month
fsubseq <- seqefsub(actcal.eseq, min.support=10,
                      constraint=seqeconstraint(max.gap=2, window.size=6))
fsubseq[1:10]
seqeid

Retrieve unique ids from an event sequence object.

Description

Retrieve the unique ids from an event sequence object or from a list of event sequence object.

Usage

seqeid(eseq, s)

Arguments

eseq An event sequence object (as created with seqecreate) or a list of event sequence objects
s Deprecated. Use eseq instead.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

Examples

data(actcal.tse)
actcal.eseq <- seqecreate(actcal.tse)
sid <- seqeid(actcal.eseq)
length(sid)
head(sid)

seqelength

Lengths of event sequences

Description

The length of an event sequence is its time span, i.e., the total time of observation. This information is useful to perform for instance a survival analysis. The function seqelength retrieves the lengths of the provided sequences, while seqelength <- sets the length of the sequences.

Usage

seqelength(eseq, s)
seqelength(eseq, s) <- value
Arguments

eseq  An event sequence object (seqelist).
value  A list of sequence lengths.
s    Deprecated. Use eseq instead.

Value

A numeric vector with the lengths of the sequences.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

Examples

data(actcal.tse)
actcal.eseq <- seqecreate(actcal.tse)
## Since end.event is not specified, contains no sequence lengths
## We set them manually as 12 for all sequences
sl <- numeric()
sl[1:2000] <- 12
seqelength(actcal.eseq) <- sl
actcal.eseq[1:10]
## Retrieve lengths
slen <- seqelength(actcal.eseq)
summary(slen)

seqetm  Create a transition-definition matrix

Description

This function automatically creates a transition-definition matrix from a state sequence object to transform the state sequences into time stamped event sequences (in TSE format).

Usage

seqetm(seqdata, method = "transition", use.labels = TRUE, sep = ">",
       bp = "", ep = "end", seq)

Arguments

seqdata  State sequence object from which transition events will be determined
method   The method to use. One of "transition", "period" or "state".
use.labels  If TRUE, transition names are built from state labels rather than from the alphabet.
sep    Separator to be used between the from-state and to-state that define the transition ("transition" method).
**seqetm**

bp       Prefix for beginning of period event names ("period" method)
ep       Prefix for end of period event names ("period" method)
seq      Deprecated. Use seqdata instead.

**Details**

Warning!!!: State labels should not contain commas ",", which are reserved for separating multiple events of a same transition!

One of three methods can be selected with the method argument:

"transition" generates a single (from-state > to-state) event for each found transition and a distinct start-state event for each different sequence start;

"period" generates a pair of events (end-state-event, start-state-event) for each found transition, a start-state event for the beginning of the sequence and an end-state event for the end of the sequence; names used for end-state and start-state names can be controlled with the bp and ep arguments;

"state" generates only the to-state event of each found transition (useful for analysing state sequences with methods for event sequences);

**Value**

The transition-definition matrix.

**Author(s)**

Matthias Studer (with Gilbert Ritschard for the help page)

**See Also**

seqformat for converting to TSE format, seqecreate for creating an event sequence object, seqdef for creating a state sequence object.

**Examples**

```r
## Creating a state sequence object from columns 13 to 24
## in the 'actcal' example data set
data(actcal)
actcal.seq <- seqdef(actcal,13:24,
  labels=c("FullTime", "PartTime", "LowPartTime", "NoWork"))
## Creating a transition matrix, one event per transition
seqtm(actcal.seq,method = "transition")

## Creating a transition matrix, single to-state events
seqtm(actcal.seq,method = "state")

## Creating a transition matrix, two events per transition
seqtm(actcal.seq,method = "period")

## changing the prefix of period start event.
seqtm(actcal.seq,method = "period", bp="begin")
```
seqeweight  

Setting or retrieving weights of an event sequence object.

Description

Event sequence objects can be weighted. Weights are used by other functions such as seqefsub or seqecmpgroup to compute weighted statistics.

Usage

```
seqeweight(eseq, s)
seqeweight(eseq, s) <- value
```

Arguments

- `eseq`: An event sequence object (seqelist).
- `value`: Numerical vector containing weights
- `s`: Deprecated. Use `eseq` instead.

Value

seqeweight returns a numerical vector containing the weights associated to each event sequence.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

Examples

```
## Starting with states sequences
## Loading data
data(biofam)
## Creating state sequences
biofam.seq <- seqdef(biofam,10:25,informat='STS')

## Creating event sequences from biofam
biofam.eseq <- seqecreate(biofam.seq, weighted=FALSE)

## Using the weights
seqeweight(biofam.eseq) <- biofam$wp00tbgs

## Now seqefsub accounts for weights unless weighted is set to FALSE
fsubseq <- seqefsub(biofam.eseq, pmin.support=0.01)

## Searching for weighted subsequences which best
## discriminate the birth cohort
discr <- seqecmpgroup(fsubseq, group=biofam$birthyr>=1940)
plot(discr[1:15])
```
Description

Finds the row indexes of state sequence(s) x in the state sequence object y.

Usage

\[
\text{seqfind}(x, y)
\]

Arguments

\[
\begin{align*}
\text{x} & \quad \text{a state sequence object containing one or more sequences (seqdef).} \\
\text{y} & \quad \text{a state sequence object.}
\end{align*}
\]

Value

row index(es) of sequence(s) x in the set of sequences y.

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

See Also

.

Examples

\[
\begin{align*}
data(\text{mvad}) \\
\text{mvad}\text{.shortlab} & \leftarrow \text{c("EM", "FE", "HE", "JL", "SC", "TR")} \\
\text{mvad}\text{.seq} & \leftarrow \text{seqdef(mvad, states=mvad\text{.shortlab, 15:86})}
\end{align*}
\]

\[
\begin{align*}
\# \text{ Finding occurrences of sequence 176 in mvad\text{.seq}} \\
\text{seqfind(mvad\text{.seq}[176,],mvad\text{.seq})}
\end{align*}
\]

\[
\begin{align*}
\# \text{ Finding occurrences of sequence 1 to 8 in mvad\text{.seq}} \\
\text{seqfind(mvad\text{.seq}[1:8,],mvad\text{.seq})}
\end{align*}
\]
seqformat

Conversion between sequence formats

Description

Convert a sequence data set from one format to another.

Usage

```r
seqformat(data, var = NULL, from, to, compress = FALSE, nrep = NULL, tevent,
stsep = NULL, covar = NULL, SPS.in = list(xfix = "()", sdsep = ","),
SPS.out = list(xfix = "()", sdsep = ","), id = 1, begin = 2, end = 3,
status = 4, process = TRUE, pdata = NULL, pvar = NULL, limit = 100,
overwrite = TRUE, fillblanks = NULL, tmin = NULL, tmax = NULL, missing = ";\*",
with.missing = TRUE, right="DEL", compressed, nr)
```

Arguments

data
- **Data Frame, Matrix, or State Sequence Object.** The data to use.
  - A data frame or a matrix with sequence data in one or more columns when `from = "STS"` or `from = "SPS"`. If sequence data are in a single column, they are assumed to be in the compressed form (see `stsep`).
  - A data frame with sequence data in one or more columns when `from = "SPELL"`. If sequence data has not four columns ordered as individual ID, spell start time, spell end time, and spell state status, use `var` or `id / begin / end / status`.
  - A state sequence object when `from = "STS"` or `from` is not specified.

var
- **NULL, List of Integers or Strings.** Default: `NULL`. The indexes or the names of the columns with the sequence data in `data`. If `NULL`, all columns are considered.

from
- **String.** The format of the input sequence data. It can be "STS", "SPS", or "SPELL". It is not needed if `data` is a state sequence object.

to
- **String.** The format of the output data. It can be "STS", "DSS", "SPS", "SRS", "SPELL", or "TSE".

compress
- **Logical.** Default: `FALSE`. When `to = "STS"`, `to = "DSS"`, or `to = "SPS"`, should the sequences (row vector of states) be concatenated into strings? See `seqconc`.

nrep
- **Integer.** The number of shifted replications when `to = "SRS"`.

tevent
- **Matrix.** The transition-definition matrix when `to = "TSE"`. It should be of size `d * d` where `d` is the number of distinct states appearing in the sequences. The cell `(i,j)` lists the events associated with a transition from state `i` to state `j`. It can be created with `seqetm`.

stsep
- **NULL, Character.** Default:`NULL`. The separator between states in the compressed form (strings) when `from = "STS"` or `from = "SPS"`. If `NULL`, `seqfcheck` is called for detecting automatically a separator among ";-" and ";.". Other separators must be specified explicitly. See `seqdecomp`.
covar

List of Integers or Strings. The indexes or the names of additional columns in data to include as covariates in the output when to = "SRS". The covariates are replicated across the shifted replicated rows.

SPS.in

List. Default: list(xfix = "()", sdsep = ","). The specifications for the state-duration couples in the input data when from = "SPS". The first specification, xfix, specifies the prefix/suffix character. Use a two-character string if the prefix and the suffix differ. Use xfix = "" when no prefix/suffix are present. The second specification, sdsep, specifies the state/duration separator.

SPS.out

List. Default: list(xfix = "()", sdsep = ","). The specifications for the state-duration couples in the output data when to = "SPS". See SPS.in above.

id

NULL, Integer, String. List of Integers or Strings. Default: 1. When from = "SPELL", the index or the name of the column containing the individual IDs in data (after var filtering).

When to = "TSE", the index or the name of the column containing the individual IDs in data (after var filtering) or the unique individual IDs. If id is not manually specified, id is set as NULL for backward compatibility with TraMineR 1.8-13 behaviour. If id is manually or automatically set as NULL, the original individual IDs are ignored and replaced by the indexes of the sequences in the input data.

When from = "SPELL" and to = "TSE", the index or the name of the column containing the individual IDs in data (after var filtering). The TSE output will use the original individual IDs.

begin

Integer or String. Default: 2. The index or the name of the column containing the spell start times in data (after var filtering) when from = "SPELL". Start times should be positive integers.

end

Integer or String. Default: 3. The index or the name of the column containing the spell end times in data (after var filtering) when from = "SPELL". End times should be positive integers.

status

Integer or String. Default: 4. The index or the name of the column containing the spell statuses in data (after var filtering) when from = "SPELL".

process

Logical. Default: TRUE. When from = "SPELL", if TRUE, create sequences on a process time axis, if FALSE, create sequences on a calendar time axis.

This process argument as well as the associated pdata and pvar arguments are intended for data containing spell data with calendar begin and end times. When those times are ages, use process = FALSE with pdata=NULL to use those ages as process times. Option process = TRUE does currently not work for age times.

pdata

NULL, "auto", or Data Frame. Default: NULL.

If NULL, the start and end times of each spell are supposed to be, if process = TRUE, ages, if process = FALSE, years when from = "SPELL".

If "auto", ages are computed using the start time of the first spell of each individual as her/his birthdate when from = "SPELL" and process = TRUE. For from = "SPELL" and process = FALSE, "auto" is equivalent to NULL.

A data frame containing the ID and the birth time of the individuals when from = "SPELL" or to = "SPELL". Use pvar to specify the column names.
The ID is used to match the birth time of each individual with the sequence data. The birth time should be integer. It is the start time from which the positions on the time axis are computed. It also serves to compute $t_{\min}$ and to guess $t_{\max}$ when the latter are NULL, from = "SPELL", and process = FALSE.

**pvar**
List of Integers or Strings. The indexes or names of the columns of the data frame `pdata` that contain the ID and the birth time of the individuals in that order.

**limit**
Integer. Default: 100. The maximum age of age sequences when from = "SPELL" and process = TRUE. Age sequences will be considered to start at 1 and to end at `limit`.

**overwrite**
Logical. Default: TRUE. When from = "SPELL", if TRUE, the most recent episode overwrites the older one when they overlap each other, if FALSE, in case of overlap, the most recent episode starts after the end of the previous one.

**fillblanks**
Character. The value to fill gaps between episodes when from = "SPELL".

**tmin**
NULL or Integer. Default: NULL. The start time of the axis when from = "SPELL" and process = FALSE. If NULL, the value is the minimum of the spell start times (see `begin`) or the minimum of the birth time of the individuals (see `pdata` when it is a data frame and process = FALSE).

**tmax**
NULL or Integer. Default: NULL. The end time of the axis when from = "SPELL" and process = FALSE. If NULL, the value is the maximum of the spell end times (see `end`) or the sum of the maximum of the spell end times and of the maximum of the birth time of the individuals (see `pdata` when it is a data frame and process = FALSE).

**missing**
String. Default: ".". The code for missing states in data. It will be replaced by NA in the output data. Ignored when data is a state sequence object (see `seqdef`), in which case the attribute `nr` is used as missing value code.

**with.missing**
Logical. Default: TRUE. When to = "SPELL", should the spells of missing states be included?

**right**
One of "DEL" or NA. Default: "DEL". When to = "SPELL" and with.missing=TRUE, set right=NA to include the end spells of missing states.

**compressed**
Deprecated. Use `compress` instead.

**nr**
Deprecated. Use `missing` instead.

**Details**

The `seqformat` function is used to convert data from one format to another. The input data is first converted into the STS format and then converted to the output format. Depending on input and output formats, some information can be lost in the conversion process. The output is a matrix or a data frame, NOT a sequence object to be passed to TraMineR functions for plotting and mining sequences (use the `seqdef` function for that). See Gabadinho et al. (2009) and Ritschard et al. (2009) for more details on longitudinal data formats and converting between them.

When data are in "SPELL" format (from = "SPELL"), the begin and end times are expected to be positions in the sequences. Therefore they should be strictly positive integers. With process=TRUE, the outcome sequences will be aligned on ages (process duration since birth), while with process=FALSE they will be aligned on dates (position on the calendar time). If TRUE, values in the `begin` and `end`
columns of data are assumed to be integer dates when \( \text{pdata} \) is not NULL, and ages otherwise. If FALSE, begin and end values are assumed to be dates when \( \text{pdata} \) is NULL and ages otherwise.

**Value**

A data frame for SRS, TSE, and SPELL, a matrix otherwise.

**Author(s)**

Alexis Gabadinho, Pierre-Alexandre Fonta, Nicolas S. Müller, Matthias Studer, and Gilbert Ritschard.

**References**


**See Also**

\texttt{seqdef}

**Examples**

```r
## Examples with raw STS sequences as input
##
## Loading a data frame with sequence data in the columns 13 to 24
data(actcal)

## Converting to SPS format
actcal.SPS.A <- seqformat(actcal, 13:24, from = "STS", to = "SPS")
head(actcal.SPS.A)

## Converting to compressed SPS format with no
## prefix/suffix and with "/" as state/duration separator
actcal.SPS.B <- seqformat(actcal, 13:24, from = "STS", to = "SPS",
                          compress = TRUE, SPS.out = list(xfix = ",", sdsep = "/"))
head(actcal.SPS.B)

## Converting to compressed DSS format
actcal.DSS <- seqformat(actcal, 13:24, from = "STS", to = "DSS",
                         compress = TRUE)
head(actcal.DSS)

## Examples with a state sequence object as input
```
## Loading a data frame with sequence data in the columns 10 to 25
data(biofam)

## Limiting the number of considered cases to the first 20
biofam <- biofam[1:20,]

## Creating a state sequence object
biofam.labs <- c("Parent", "Left", "Married", "Left/Married", "Child", "Left/Child", "Left/Married/Child", "Divorced")
biofam.seq <- seqdef(biofam, 10:25, alphabet = 0:7,
                      states = biofam.short.labs, labels = biofam.labs)

## Converting to SPELL format
bf.spell <- seqformat(biofam.seq, from = "STS", to = "SPELL",
                      pdata = biofam, pvar = c("idhous", "birthyr"))
head(bf.spell)

## Examples with SPELL sequences as input

## Loading two data frames: bfspell20 and bfpdata20
## bfspell20 contains the first 20 biofam sequences in SPELL format
## bfpdata20 contains the IDs and the years at which the
## considered individuals were aged 15
data(bfspell)

## Converting to STS format with alignment on calendar years
bf.sts.y <- seqformat(bfspell20, from = "SPELL", to = "STS",
                      id = "id", begin = "begin", end = "end", status = "states",
                      process = FALSE)
head(bf.sts.y)

## Converting to STS format with alignment on ages
bf.sts.a <- seqformat(bfspell20, from = "SPELL", to = "STS",
                      id = "id", begin = "begin", end = "end", status = "states",
                      process = TRUE, pdata = bfpdata20, pvar = c("id", "when15"),
                      limit = 16)
names(bf.sts.a) <- paste0("a", 15:30)
head(bf.sts.a)

## Examples for TSE and SPELL output
## in presence of missing values

## STS data with missing values
## creating the state sequence object with by default
seqfpos

Search for the first occurrence of a given element in a sequence

Description

Returns a vector containing the position of the first occurrence of the given element in each of the sequences in the data set.

Usage

seqfpos(seqdata, state)

Arguments

seqdata a sequence object (see seqdef function).
state the state element to search in the sequences

Details

The state to search for has to be passed as a character string, and must be one of the state returned by the alphabet function. If the state is not contained in a sequence, NA is returned for this sequence.

Author(s)

Alexis Gabadinho
Examples

```r
seqgen <- seqdef(biofam, 10:25)

## Searching for the first occurrence of state 1
## in each of the 5 first sequence of the biofam data set.
seqfpos(biofam.seq[1:5], "1")
```

---

**Description**

Generates random sequences.

**Usage**

```r
seqgen(n, length, alphabet, p)
```

**Arguments**

- `n`: number of sequences to generate
- `length`: sequences length
- `alphabet`: the alphabet from which the sequences are generated
- `p`: an optional vector of probabilities for the states in the alphabet. Must be of the same length as the alphabet. If not specified, equal probabilities are used.

**Details**

Each sequence is generated by choosing a set of random numbers (with min=1 and max=length of the alphabet) using the `runif` function. When the probability distribution is not specified, the uniform probability distribution giving same probability to each state is used to generate the sequences.

**Value**

a sequence object.

**Author(s)**

Alexis Gabadinho (with Gilbert Ritschard for the help page)

**Examples**

```r
seq <- seqgen(1000, 10, 1:4, c(0.2, 0.1, 0.3, 0.4))
seqstatd(seqdef(seq))
```
seqici  Complexity index of individual sequences

Description
Computes the complexity index, a composite measure of sequence complexity. The index uses the number of transitions in the sequence as a measure of the complexity induced by the state ordering and the longitudinal entropy as a measure of the complexity induced by the state distribution in the sequence.

Usage
seqici(seqdata, with.missing=FALSE, silent=TRUE)

Arguments
seqdata  a sequence object as returned by the seqdef function.
with.missing if set to TRUE, missing status (gaps in sequences) is handled as an additional state when computing the state distribution and the number of transitions in the sequence.
silent  logical: should messages about running operations be displayed?

Details
The complexity index $C(s)$ of a sequence $s$ is

$$C(s) = \sqrt{\frac{q(s)}{q_{\text{max}}} \frac{h(s)}{h_{\text{max}}}}$$

where $q(s)$ is the number of transitions in the sequence, $q_{\text{max}}$ the maximum number of transitions, $h(s)$ the within entropy, and $h_{\text{max}}$ the theoretical maximum entropy which is $h_{\text{max}} = -\log 1/|A|$ with $|A|$ the size of the alphabet.

The index $C(s)$ is the geometric mean of its two normalized components and is, therefore, itself normalized. The minimum value of 0 can only be reached by a sequence made of one distinct state, thus containing 0 transitions and having an entropy of 0. The maximum 1 of $C(s)$ is reached when the two following conditions are fulfilled: i) Each of the state in the alphabet is present in the sequence, and the total durations are uniform, i.e. each state occurs $\ell/|A|$ times, and ii) the number of transitions in the sequence is $\ell - 1$, meaning that the length $\ell_d$ of the DSS is equal to the length of the sequence $\ell$.

Value
a single-column matrix of length equal to the number of sequences in seqdata containing the complexity index value of each sequence.
Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

References


See Also

`seqdss`, `seqstatd`. For other measures of sequence complexity see `seqient` and `seqST`.

Examples

```r
### Creating a sequence object from the mvad data set
data(mvad)
mvad.labels <- c("employment", "further education", "higher education", "joblessness", "school", "training")
mvad.scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad.seq <- seqdef(mvad, 15:86, states=mvad.scodes, labels=mvad.labels)

###
mvad.ci <- seqici(mvad.seq)
summary(mvad.ci)
hist(mvad.ci)

### Example using with.missing argument
data(ex1)
ex1.seq <- seqdef(ex1, 1:13)
seqici(ex1.seq)
seqici(ex1.seq, with.missing=TRUE)
```

---

**seqient**  
*Within sequence entropies*

Description

Computes normalized or non-normalized within sequence entropies

Usage

`seqient(seqdata, norm=TRUE, base=exp(1), with.missing=FALSE, silent=TRUE)`
seqient

Arguments

seqdata: a sequence object as returned by the `seqdef` function.
norm: logical: should the entropy be normalized? TRUE by default. (see details)
base: real positive value: base of the logarithm used in the entropy formula (see details). Default is `exp(1)`, i.e., the natural logarithm is used.
with.missing: logical: if TRUE, the missing state (gap in sequences) is handled as an additional state when computing the state distribution in the sequence.
silent: logical: should messages about running operations be displayed?

Details

The seqient function returns the Shannon entropy of each sequence in seqdata. The entropy of a sequence is computed using the formula

$$ h(\pi_1, \ldots, \pi_s) = - \sum_{i=1}^{s} \pi_i \log \pi_i $$

where $s$ is the size of the alphabet and $\pi_i$ the proportion of occurrences of the $i$th state in the considered sequence. The base of the log is controlled with the base argument. By default the natural logarithm, i.e. the logarithm in base $e$, is used. The entropy can be interpreted as the ‘uncertainty’ of predicting the states in a given sequence. If all states in the sequence are the same, the entropy is equal to 0. For example, the maximum entropy for a sequence of length 12 with an alphabet of 4 states is 1.386294 and is attained when each of the four states appears 3 times.

Normalization can be requested with the `norm=TRUE` option, in which case the returned value is the entropy divided by the entropy of the alphabet. The latter is an upper bound for the entropy of sequences made from this alphabet. It is exactly the maximal entropy when the sequence length is a multiple of the alphabet size. The value of the normalized entropy is independent of the chosen logarithm base.

Value

a single-column matrix with an entropy value for each sequence in seqdata; the column length is equal to the number of sequences.

Author(s)

Alexis Gabadinho

References


See Also

seqistatd for the entropy of the transversal state distributions by positions in the sequence, and the composite measures of complexity seqici and seqST.

Examples

data(actcal)
actcal.seq <- seqdef(actcal, 13:24)

## Summarize and plots an histogram
## of the within sequence entropy
actcal.lent <- seqient(actcal.seq)
summary(actcal.lent)
hist(actcal.lent)

## Examples using with.missing argument
data(ex1)
ex1.seq <- seqdef(ex1, 1:13, weights=ex1$weights)

seqient(ex1.seq)
seqient(ex1.seq, with.missing=TRUE)

seqistatd State frequencies in each individual sequence

Description

Returns the state frequencies (total durations) for each sequence in the sequence object.

Usage

seqistatd(seqdata, with.missing=FALSE, prop=FALSE)

Arguments

seqdata a sequence object (see seqdef function).
with.missing logical: if set as TRUE, total durations are also computed for the missing status (gaps in the sequences). See seqdef on options for handling missing values when creating sequence objects.
prop logical: if TRUE, proportions of time spent in each state are returned instead of absolute values. This option is specially useful when sequences contain missing states, since the sum of the state durations may not be the same for all sequences.

Author(s)

Alexis Gabadinho
References


Examples

```r
data(actcal)
actcal.seq <- seqdef(actcal,13:24)
seqistatd(actcal.seq[1:10,])

## Example using "with.missing" argument
data(ex1)
ex1.seq <- seqdef(ex1, 1:13, weights=ex1$weights)

seqistatd(ex1.seq)
seqistatd(ex1.seq, with.missing=TRUE)
```

---

**seqlegend**

*Plot a legend for the states in a sequence object*

**Description**

Plots a legend for the states in a sequence object. Useful if several graphics are plotted together and only one legend is necessary. Unless specified by the user, the *cpal* and *labels* attributes of the sequence object are used for the colors and text appearing in the legend (see *seqdef*).

**Usage**

```r
seqlegend(seqdata, with.missing = "auto", cpal = NULL, missing.color = NULL,
ltext = NULL, position = "topleft", cex = 1, fontsize, ...)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>seqdata</code></td>
<td>a sequence object as returned by the <code>seqdef</code> function.</td>
</tr>
<tr>
<td><code>with.missing</code></td>
<td>if set to &quot;auto&quot; (default), a legend for the missing state is added automatically if one or more of the sequences in <code>seqdata</code> contains a missing state. If TRUE a legend for the missing state is added in any case. Setting to FALSE omits the legend for the missing state.</td>
</tr>
<tr>
<td><code>cpal</code></td>
<td>alternative color palette to use for the states. If user specified, a vector of colors with number of elements equal to the number of distinct states. By default, the 'cpal' attribute of the 'seqdata' sequence object is used (see <code>seqdef</code>).</td>
</tr>
<tr>
<td><code>missing.color</code></td>
<td>alternative color for representing missing values inside the sequences. By default, this color is taken from the &quot;missing.color&quot; attribute of the sequence object being plotted.</td>
</tr>
</tbody>
</table>
seqlength

Description

Returns the length of sequences.

Usage

seqlength(seqdata)

Arguments

seqdata a sequence object created with the seqdef function.
Details

The length of a sequence is computed by eliminating the missing values at the end (right) and counting the number of states or events. The seqlength function returns a vector containing the length of each sequence in the sequence object given as argument.

Author(s)

Alexis Gabadinho

Examples

```r
## Loading the 'famform' example data set
data(famform)

## Defining a sequence object with the 'famform' data set
ff.seq <- seqdef(famform)

## Retrieving the length of the first 10 sequences
## in the ff.seq sequence object
seqlength(ff.seq)
```

seqLLCP

| seqLLCP | Compute the length of the longest common prefix of two sequences |

Description

Returns the length of the longest common prefix of two sequences. This attribute is described in Elzinga (2008).

Usage

```
seqLLCP(seq1, seq2)
```

Arguments

- `seq1` a sequence from a sequence object.
- `seq2` a sequence from a sequence object.

Value

an integer being the length of the longest common prefix of the two sequences.

References

seqLLCS

Compute the length of the longest common subsequence of two sequences

Description

Returns the length of the longest common subsequence of two sequences. This attribute is described in Elzinga (2008).

Usage

seqLLCS(seq1, seq2)

Arguments

seq1 a sequence from a sequence object
seq2 a sequence from a sequence object

Value

an integer being the length of the longest common subsequence of the two sequences.

References


See Also

seqdist

Examples

LCS.ex <- seqdef(LCS.ex)
seqLLCS(LCS.ex[1,], LCS.ex[3,])
seqlogp

Logarithm of the probabilities of state sequences

Description

Compute the logarithm of the probability of each state sequence obtained from a state transition model. The probability of a sequence is equal to the product of each state probability of the sequence. There are several methods to compute a state probability.

Usage

seqlogp(seqdata, prob="trate", time.varying=TRUE,
       begin="freq", weighted=TRUE)

Arguments

seqdata The sequence to compute the probabilities.
prob either the name ("trate" or "freq") of the probability model to use to compute
       the state probabilities, or an array specifying the transition probabilities at each
       position t (see details).
time.varying Logical. If TRUE, the probabilities (transitions or frequencies) are computed
       separately for each time t point.
begin Model used to compute the probability of the first state. Either "freq" to use the
       observed frequencies on the first period or a vector specifying the probability of
       each state of the alphabet.
weighted Logical. If TRUE, uses the weights specified in seqdata when computing the
       observed transition rates.

Details

The sequence likelihood \( P(s) \) is defined as the product of the probability with which each of its observed successive state is supposed to occur at its position. Let \( s = s_1 s_2 \cdots s_\ell \) be a sequence of length \( \ell \). Then

\[
P(s) = P(s_1, 1) \cdot P(s_2, 2) \cdots P(s_\ell, \ell)
\]

with \( P(s_t, t) \) the probability to observe state \( s_t \) at position \( t \).

The question is how to determinate the state probabilities \( P(s_t, t) \). Several methods are available and can be set using the prob argument.

One commonly used method for computing them is to postulate a Markov model, which can be of various order. We can consider probabilities derived from the first order Markov model, that is, each \( P(s_t, t), t > 1 \) is set as the transition rate \( p(s_t | s_{t-1}) \). This is available in seqlogp by setting prob="trate".

The transition rates may be considered constant over time/positions (time.varying=FALSE), that is estimated across sequences from the observations at positions \( t \) and \( t - 1 \) for all \( t \) together. Time varying transition rates may also be considered (time.varying=TRUE), in which case they are computed separately for each position, that is estimated across sequences from the observations.
at positions $t$ and $t-1$ for each $t$, yielding an array of transition matrices. The user may also specify
his own transition rates array or matrix.

Another method is to use the frequency of a state at each position to set $P(s_i, t)$ ($\text{prob}="\text{freq}"$). In the latter case, the probability of a sequence is independent of the probability of the transitions. Here again, the frequencies can be computed all together ($\text{time\_varying=}\text{FALSE}$) or separately for each position $t$ ($\text{time\_varying=}\text{TRUE}$).

For $t=1$, we set $P(s_1, 1)$ to the observed frequency of the state $s_1$ at position 1. Alternatively, the
begin argument allows to specify the probability of the first state.

The likelihood $P(s)$ being generally very small, `seqlogp` return $-\log P(s)$. The latter quantity is
minimal when $P(s)$ is equal to 1.

**Value**

A vector containing the logarithm of each sequence probability.

**Author(s)**

Matthias Studer and Alexis Gabadinho (with Gilbert Ritschard for the help page)

**Examples**

```r
## Creating the sequence objects using weights
biofam.seq <- seqdef(biofam, 10:25, weights=biofam$wp0@thgs)

## Computing sequence probabilities
biofam.prob <- seqlogp(biofam.seq)
## Comparing the probability of each cohort
cohort <- biofam$birthyr>1940
boxplot(biofam.prob~cohort)
```

---

**seqmeant**

*Mean durations in each state*

**Description**

Compute the mean total time spent in each state of the alphabet for the set of sequences given as input.

**Usage**

`seqmeant(seqdata, weighted=TRUE, with.missing=FALSE, prop=FALSE, serr=FALSE)`
Arguments

seqdata a sequence object as defined by the seqdef function.

weighted logical: if TRUE, the weights (weights attribute) attached to the sequence object are used for computing weighted mean total time.

with.missing logical: if set to TRUE, cumulated durations are also computed for the missing status (gaps in the sequences). See seqdef on options for handling missing values when creating sequence objects.

prop logical: if TRUE, proportions of time spent in each state are returned instead of absolute values. This option is especially useful when sequences contain missing states, since the sum of the state durations may not be the same for all sequences.

serr logical: if TRUE, the variance and standard deviation of the total time spent in the states, as well as the standard error of the mean are also computed.

Value

An object of class stslist.meant. There are print and plot methods for such objects.

Author(s)

Alexis Gabadinho

References


See Also

plot.stslist.meant for basic plots of stslist.meant objects and seqmtplot (seqplot with type="mt") argument for more sophisticated plots of the mean durations allowing grouping and legend.

Examples

```r
## Defining a sequence object with columns 13 to 24
## in the actcal example data set
data(actcal)
actcal.lab <- c("> 37 hours", "19-36 hours", "1-18 hours", "no work")
actcal.seq <- seqdef(actcal,13:24,labels=actcal.lab)

## Computing the mean time in the different states
seqmeant(actcal.seq)

## Mean times with their standard error
seqmeant(actcal.seq, serr=TRUE)
```
seqmodst  

Sequence of modal states

Description

Sequence made of the modal state at each position.

Usage

seqmodst(seqdata, weighted=TRUE, with.missing=FALSE)

Arguments

seqdata  
a state sequence object as defined by the seqdef function.

weighted  
if TRUE, distributions account for the weights assigned to the state sequence object (see seqdef). Set as FALSE if you want ignore the weights.

with.missing  
If FALSE (default value), returned distributions ignore missing values.

Details

In case of multiple modal states at a given position, the first one is taken. Hence, the result may vary with the alphabet order.

Value

an object of class stslist.modst. This is actually a state sequence object (containing a single state sequence) with additional attributes, among which the frequencies attribute containing the transversal frequency of each state in the sequence. There are print and plot methods for such objects. More sophisticated plots can be produced with the seqplot function.

Author(s)

Alexis Gabadinho

References


See Also

plot.stslist.modst for default plot method, seqplot for higher level plots.
Examples

```r
## Defining a sequence object with the data in columns 10 to 25
## (family status from age 15 to 30) in the biofam data set
data(biofam)
biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

## Modal state sequence
seqmodst(biofam.seq)

## Examples using weights and with.missing arguments
data(exl)
exl.seq <- seqdef(exl, 1:13, weights=exl$weights)

seqmodst(exl.seq)
seqmodst(exl.seq, weighted=FALSE)
seqmodst(exl.seq, weighted=FALSE, with.missing=TRUE)
```

---

**seqmpos**

*Number of matching positions between two sequences.*

**Description**

Returns the number of common elements, i.e., same states appearing at the same position in the two sequences.

**Usage**

```r
seqmpos(seq1, seq2, with.missing=FALSE)
```

**Arguments**

- `seq1` a sequence from a sequence object.
- `seq2` a sequence from a sequence object.
- `with.missing` if TRUE, gaps appearing at the same position in both sequences are also considered as common elements.

**Author(s)**

Alexis Gabadinho (with Gilbert Ritschard for help page)

**See Also**

`seqLLCP, seqLLCS`
Examples

```r
data(famform)
famform.seq <- seqdef(famform)

seqnpos(famform.seq[1,], famform.seq[2,])
seqnpos(famform.seq[2,], famform.seq[4,])

## Example with gaps in sequences
a <- c(NA, "A", NA, "B", "C")
b <- c(NA, "C", NA, "B", "C")
ex1.seq <- seqdef(rbind(a, b))

seqnpos(ex1.seq[1,], ex1.seq[2,])
seqnpos(ex1.seq[1,], ex1.seq[2,], with.missing=TRUE)
```

---

**seqnum**

*Transform into a sequence object with numerical alphabet.*

**Description**

The function `seqnum` transforms the provided state sequence object into an equivalent sequence object in which the original alphabet is replaced with an alphabet of numbers ranging from 0 to (nbstates-1).

**Usage**

```r
seqnum(seqdata, with.missing=FALSE)
```

**Arguments**

- `seqdata` a state sequence object as defined by the `seqdef` function.
- `with.missing` logical: Should missing elements in the sequences be turned into numerical values as well? The code for missing values in the sequences is retrieved from the 'nr' attribute of `seqdata`.

**Details**

The first state (for example 'A') is coded with the value 0, the second state (for example 'B') is coded with the value 1, etc... The function returns a sequence object containing the original sequences coded with the new numerical alphabet ranging from 0 to (nbstates-1).

**Author(s)**

Alexis Gabadinho
seqpcplot

See Also

seqdef, alphabet

Examples

data(actcal)
actcal.seq <- seqdef(actcal, 13:24)

## The first 10 sequences in the actcal.seq
## sequence object
actcal.seq[1:10,]
alphabet(actcal.seq)

## The first 10 sequences in the actcal.seq
## sequence object with numerical alphabet
seqnum(actcal.seq[1:10,])

## states A,B,C,D are now coded 0,1,2,3
alphabet(seqnum(actcal.seq))

Description

A decorated parallel coordinate plot to render the order of the successive elements in sequences. The
sequences are displayed as jittered frequency-weighted parallel lines. The plot is also embedded as
the type="pc" option of the seqplot function and serves as plot method for eseq and seqelist
objects.

Usage

seqpcplot(seqdata, group = NULL, weights = NULL, cex = 1, lwd = 1/4,
cpal = NULL, grid.scale = 1/5, ltype = "unique",
embedding = "most-frequent", lorder = NULL, lcourse = "upwards",
filter = NULL, hide.col = "grey80", alphabet = NULL,
missing = "auto", order.align = "first", main = NULL, xlab = NULL,
ylab = NULL, xaxis = TRUE, yaxis = TRUE, axes = "all", xlab = NULL,
cex.lab = 1, rows = NA, cols = NA, plot = TRUE, seed = NULL,
weighted = TRUE, with.missing = TRUE,
title, cex.plot, ...)

sepcfilter(method = c("minfreq", "cumfreq", "linear"), level = 0.05)
Arguments

seqdata  The sequence data. Either an event sequence object of class `seqelist` (see `seqecreate`) or a state sequence object of class `stslist` (see `seqdef`).

group    a vector (numeric or factor) of group memberships of length equal the number of sequences. When specified, one plot is generated for each different membership value.

weights  a numeric vector of weights of length equal the number of sequences. When `NULL`, the weights are taken from the `seqdata` object.

cex       Plotting text and symbols magnification. See `par`.

lwd       expansion factor for line widths. The expansion is relative to the size of the squared symbols.

cpal      color palette vector for line coloring.

grid.scale Expansion factor for the translation zones.

ltype     the type of sequence that is drawn. Either "unique" to render unique patterns or "non-embeddable" to render non-embeddable sequences.

embedding The method for embedding sequences embeddable in multiple non-embeddable sequences. Either "most-frequent" (default) or "uniformly". Relevant only with `ltype = "non-embeddable"`.

lorder    line ordering. Either "background" or "foreground".

lcourse   Method to connect simultaneous elements with the preceding and following ones. Either "upwards" (default) or "downwards".

filter    list of line coloring options. See details.

hide.col  Color for sequences filtered-out by the filter specification.

alphabet  a vector of response levels in the order they should appear on the y-axis. This argument is solely relevant for `seqelist` objects.

missing   character. Whether and how missing values should be displayed. Available are "auto", "show" and "hide". If "auto", the plot will show missings only if present. "hide" will fade out missings and "show" will always show missings. If with.missing=FALSE, missing is turned into "hide". If with.missing=TRUE and missing="hide", missing is turned into "auto".

order.align Aligning method. For aligning on order positions use either "first" (default) or "last". Option "first" numbers the positions from the beginning while "last" numbers them from the end. With `order.align = "time"`, the elements in the sequences are aligned on their rounded timestamps.

main      title for the graphic.

xlab      label for the x axis

ylab      label for the y axis

xaxis     logical: Should x-axis be plotted?

yaxis     logical: Should y-axis be plotted?

axes      if set as "all" (default value) x-axes are drawn for each plot in the graphic. If set as "bottom" and group is used, axes are drawn only under the plots at the bottom of the graphic area. If `FALSE`, no x-axis is drawn.
xtlab labels for the x-axis ticks.
cex.lab x and y labels magnification. See par.
rows,cols integers. Number of rows and columns of the plot panel.
plot logical. Should the plot be displayed? Set as FALSE to retrieve the seqpcplot object without plotting it.
seed integer. Start seed value.
weighted logical. Should weights be accounted for? Default is TRUE.
with.missing logical. Should we care about possible missings? Default is TRUE. See also the missing argument.
method character string. Defines the filtering function. Available are "minfreq", "cumfreq" and "linear".
level numeric scalar between 0 and 1. The frequency threshold for the filtering methods "minfreq" and "cumfreq".
title Deprecated. Use main instead.
cex.plot Deprecated. Use cex.lab instead.
... arguments to be passed to other methods, such as graphical parameters (see par).

Details

For plots by groups specified with the group argument, plotted line widths and point sizes reflect relative frequencies within group.

The filter argument serves to specify filters to gray less interesting patterns. The filtered-out patterns are displayed in the hide.col color. The filter argument expects a list with at least elements type and value. The following types are implemented:

Type "sequence": colors a specific pattern, for example assign
filter = list(type = "sequence", value = "(Leaving Home, Union)-(Child)").

Type "subsequence": colors patterns which include a specific subsequence, for example
filter = list(type = "subsequence", value = "(Child)-(Marriage)").

Type "value": gradually colors the patterns according to the numeric vector (of length equal to the number of sequences) provided as "value" element in the list. You can give something like filter = list(type = "value", value = c(0.2, 1, ...)) or provide the distances to the medoid as value vector for example.

Type "function": colors the patterns depending on the values returned by a [0,1] valued function of the frequency x of the pattern. Three native functions can be used: "minfreq", "cumfreq" and "linear". Use filter = list(type = "function", value = "minfreq", level = 0.05) to color patterns with a support of at least 5% (within group). Use filter = list(type = "function", value = "cumfreq", level = 0.5) to highlight the 50% most frequent patterns (within group). Or, use filter = list(type="function", value="linear") to use a linear gradient for the color intensity (the most frequent trajectory gets 100% intensity).

Other user-specified functions can be provided by giving something like
filter = list(type="function", value=function(x, arg1, arg2) {return(x/max(x) * arg1/arg2)}, arg1 = 1, arg2 = 1). This latter function adjusts gradually the color intensity of patterns according to the frequency of the pattern.
The function `seqpcfilter` is a convenience function for type "function". The three examples above can be imitated by `seqpcfilter("minfreq", 0.05)`, `seqpcfilter("cumfreq", 0.5)` and `seqpcfilter("linear")`. If a numeric scalar is assigned to `filter`, the "minfreq" filter is used.

Value

An object of class "seqpcplot" with various information necessary for constructing the plot, e.g. coordinates. There is a summary method for such objects.

Author(s)

Reto Bürgin (with Gilbert Ritschard for the help page)

References


See Also

`seqplot`, `seqdef`, `seqecreate`

Examples

```r
## ===========
## plot biofam data
## ===========

data(biofam)
lab <- c("Parent","Left","Married","Left+Marr","Child","Left+Child",
         "Left+Marr+Child","Divorced")

## plot state sequences in STS representation
## ================

## creating the weighted state sequence object.
biofam.seq <- seqdef(data = biofam[,10:25], labels = lab, weights = biofam$wp00tbg)

## select the first 20 weighted sequences (sum of weights = 18)
biofam.seq <- biofam.seq[1:20, ]

par(mar=c(4,8,2,2))
seqpcplot(seqdata = biofam.seq, order.align = "time")

## .. or
seqplot(seqdata = biofam.seq, type = "pc", order.align = "time")

## Distinct successive states (DSS)
## seqpcplot

```r
## seqpcplot

# plot event sequences
seqplot(seqdata = biofam.seq, type = "pc", order.align = "first")

# or (equivalently)
biofam.DSS <- seqdss(seqdata = biofam.seq) # prepare format
seqpcplot(seqdata = biofam.DSS)

## plot the time in the x-axis
seqpcplot(seqdata = biofam.eseq, order.align = "time", alphabet = lab)

## ordering of events
seqpcplot(seqdata = biofam.eseq, order.align = "first", alphabet = lab)

# plot non-embeddable sequences
seqpcplot(seqdata = biofam.eseq, ltype = "non-embeddable", order.align = "first", alphabet = lab)

# align on last event
par(mar=c(4,8,2,2))
seqpcplot(seqdata = biofam.eseq, order.align = "last", alphabet = lab)

# use group variables
seqpcplot(seqdata = biofam.eseq, group = biofam$sex[1:20], order.align = "first", alphabet = lab)

# color patterns (Parent)-(Married) and (Parent)-(Left+Marriage+Child)
par(mfrow = c(1, 1))
seqpcplot(seqdata = biofam.eseq,
    filter = list(type = "sequence",
                  value=c("(Parent)-(Married)", "(Parent)-(Left+Marriage+Child)"),
                  alphabet = lab, order.align = "first")

# color subsequence pattern (Parent)-(Left)
seqpcplot(seqdata = biofam.eseq,
    filter = list(type = "subsequence",
                  value = "(Parent)-(Left)"),
                  alphabet = lab, order.align = "first")

# color sequences over 10% (within group) (function method)
```
seqpcplot(seqdata = biofam.eseq,  
    filter = list(type = "function",  
                  value = "minfreq",  
                  level = 0.1),  
    alphabet = lab, order.align = "first", seed = 1)

## .. same result using the convenience functions
seqpcplot(seqdata = biofam.eseq,  
    filter = 0.1,  
    alphabet = lab, order.align = "first", seed = 1)

seqpcplot(seqdata = biofam.eseq,  
    filter = seqpcfilter("minfreq", 0.1),  
    alphabet = lab, order.align = "first", seed = 1)

## highlight the 50% most frequent sequences
seqpcplot(seqdata = biofam.eseq,  
    filter = list(type = "function",  
                  value = "cumfreq",  
                  level = 0.5),  
    alphabet = lab, order.align = "first", seed = 2)

## .. same result using the convenience functions
seqpcplot(seqdata = biofam.eseq,  
    filter = seqpcfilter("cumfreq", 0.5),  
    alphabet = lab, order.align = "first", seed = 2)

## linear gradient
seqpcplot(seqdata = biofam.eseq,  
    filter = list(type = "function",  
                  value = "linear"),  
    alphabet = lab, order.align = "first", seed = 2)

seqpcplot(seqdata = biofam.eseq,  
    filter = seqpcfilter("linear"),  
    alphabet = lab, order.align = "first", seed = 1)

---

**seqplot**

*Plot state sequence objects*

**Description**

High level plot functions for state sequence objects that can produce state distribution (chrono-grams), frequency, index, transversal entropy, sequence of modes, meant time, and representative plots.
seqplot

Usage

seqplot(seqdata, group = NULL, type = "i", main = NULL, cpal = NULL, missing.color = NULL, ylab = NULL, yaxis = TRUE, axes = "all", xtlab = NULL, cex.axis = 1, with.legend = "auto", ltext = NULL, cex.legend = 1, use.layout = (!is.null(group) | with.legend != FALSE), legend.prop = NA, rows = NA, cols = NA, title, cex.plot, withlegend, ...)

seqdplot(seqdata, group = NULL, main = NULL, ...)
seqfplot(seqdata, group = NULL, main = NULL, ...)
seqiplot(seqdata, group = NULL, main = NULL, ...)
seqhtplot(seqdata, group = NULL, main = NULL, ...)
seqmsplot(seqdata, group = NULL, main = NULL, ...)
seqmtplot(seqdata, group = NULL, main = NULL, ...)
seqrplot(seqdata, group = NULL, main = NULL, ...)

Arguments

seqdata State sequence object created with the seqdef function.

group Grouping variable of length equal to the number of sequences. When not NULL, a distinct plot is generated for each level of group.

type the type of the plot. Available types are "d" for state distribution plots (chrono-

grams), "f" for sequence frequency plots, "ht" for transversal entropy plots, "i" for selected sequence index plots, "I" for whole set index plots, "ms" for plotting the sequence of modal states, "mt" for mean times plots, "pc" for parallel coordinate plots and "r" for representative sequence plots.

main Character string. Title for the graphic. Default is NULL.

cpal Color palette used for the states. By default, the cpal attribute of the seqdata

sequence object is used (see seqdef). If user specified, a vector of colors of

length and order corresponding to alphabet(seqdata).

missing.color Color for representing missing values inside the sequences. By default, this

color is taken from the missing.color attribute of seqdata.

ylab Character string. an optional label for the y-axis. If set to NA, no label is drawn.

yaxis Logical. Should the y-axis be plotted? When set as TRUE (default value), se-
quence indexes are displayed for "i" and "I", mean time values for "mt", per-

centages for "d" and "f", and state/event labels for "pc". Ignored for "r".

axes Character string or logical. If set as "all" (default value) x axes are drawn for
each plot in the graphic. If set as "bottom" and group is used, axes are drawn only under the plots located at the bottom of the graphic area. If FALSE, no x-axis is drawn.

xtlab Vector of length equal to the number of columns of seqdata. Optional labels for

the x-axis tick labels. If unspecified, the column names of the seqdata sequence

object are used (see seqdef).

cex.axis Real value. Axis annotation magnification. When type = "r" and for seqrplot(),
it also determines the magnification of the plotted text and symbols. See par.
with.legend Character string or logical. Defines if and where the legend of the state colors is plotted. The default value "auto" sets the position of the legend automatically. Other possible value is "right". Obsolete value TRUE is equivalent to "auto".

ltext Vector of character strings of length and order corresponding to alphabet(seqdata). Optional description of the states to appear in the legend. If unspecified, the label attribute of the seqdata sequence object is used (see seqdef).

cex.legend Real. Legend magnification. See legend.

use.layout Logical. Should layout be used to arrange plots when using the group option or plotting a legend? When layout is activated, the standard 'par(mfrow=....)' for arranging plots does not work. With with.legend=FALSE and group=NULL, layout is automatically deactivated and 'par(mfrow=....)' can be used.

legend.prop Real in range [0,1]. Proportion of the graphic area devoted to the legend plot when use.layout=TRUE and with.legend=TRUE. Default value is set according to the place (bottom or right of the graphic area) where the legend is plotted.

rows,cols Integers. Number of rows and columns of the plot panel when use.layout=TRUE.

title Deprecated. Number of rows and columns of the plot panel when use.layout=TRUE.

cex.plot Deprecated. Use cex.axis instead.

with.legend Deprecated. Use with.legend instead.

... arguments to be passed to the function called to produce the appropriate statistics and the associated plot method (see details), or other graphical parameters. For example the weighted argument can be passed to control whether (un)weighted statistics are produced or with.missing argument to take missing values into account when computing transversal or longitudinal state distributions.

Details

seqplot is the generic function for high level plots of state sequence objects with group splits and automatic display of the color legend. Many different types of plots can be produced by means of the type argument. Except for sequence index plots, seqplot first calls the specific function producing the required statistics and then the plot method for objects produced by this function (see below). For sequence index plots, the state sequence object itself is plotted by calling the plot stslist method. When splitting by groups and/or displaying the color legend, the layout function is used for arranging the plots.

The seqdplot, seqfplot, seqiplot, seqhtplot, seqmsplot, seqmtplot, seqpcplot and seqrplot functions are aliases for calling seqplot with type argument set respectively to "d", "f", "i", "t", "Ht", "ms", "mt", "pc" or "r".

State distribution plot (type="d") represent the sequence of the cross-sectional state frequencies by position (time point) computed by the seqstatd function. Such plots are also known as chronograms.

Sequence frequency plots (type="f") display the most frequent sequences, each one with an horizontal stack bar of its successive states. Sequences are displayed bottom-up in decreasing order of their frequencies (computed by the seqtab function). The plot stslist.freq plot method is called for producing the plot.

The idxs optional argument may be specified for selecting the sequences to be plotted (default is 1:10, i.e. the 10 most frequent sequences). The width of the bars representing the sequences is by
default proportional to their frequencies, but this can be disabled with the pbarw=FALSE optional argument. If weights have been specified when creating seqdata, weighted frequencies will be returned by seqtab since the default option is weighted=TRUE. See examples below, the seqtab and plot.stslist.freq manual pages for a complete list of optional arguments and Müller et al., (2008) for a description of sequence frequency plots.

In sequence index plots (type="i" or type="I"), the requested individual sequences are rendered with horizontal stacked bars depicting the states over successive positions (time). Optional arguments are idxs for specifying the indexes of the sequences to be plotted (when type="i" defaults to the first ten sequences, i.e. idxs=1:10). For plotting nicely a (big) whole set one can use type="I" which is the same as using idxs=0 together with the additional graphical parameters border=NA and space=0 to suppress bar borders and space between bars. The sortv argument can be used to pass a vector of numerical values for sorting the sequences or to specify a sorting method. See plot.stslist for a complete list of optional arguments and their description.

The interest of sequence index plots has, for instance, been stressed by Scherer (2001) and Brzinsky-Fay et al. (2006). Notice that index plots for thousands of sequences result in very heavy PDF or POSTSCRIPT graphic files. Dramatic file size reduction may be achieved by saving the figures in bitmap format with using for instance the png graphic device instead of postscript or pdf.

The transversal entropy plot (type="Ht") displays the evolution over positions of the transversal entropies (Billari, 2001). Transversal entropies are computed by calling seqstatd function and then plotted by calling the plot.stslist.statd plot method.

The modal state sequence plot (type="ms") displays the sequence of the modal states with each mode proportional to its frequency at the given position. The seqmodst function is called which returns the sequence and the result is plotted by calling the plot.stslist.modst plot method.

The mean time plot (type="mt") displays the mean time spent in each state of the alphabet as computed by the seqmeant function. The plot.stslist.meant plot method is used to plot the resulting statistics. Set serr=TRUE to display error bars on the mean time plot.

The representative sequence plot (type="r") displays a reduced, non redundant set of representative sequences extracted from the provided state sequence object and sorted according to a representativeness criterion. The seqrep function is called to extract the representative set which is then plotted by calling the plot.stslist.rep method. A distance matrix is required that is passed with the diss argument or by calling the seqdist function if diss=NULL. The criterion argument sets the representativeness criterion used to sort the sequences. Refer to the seqrep and plot.stslist.rep manual pages for a complete list of optional arguments. See Gabadinho and Ritschard (2013) for more details on the extraction of representative sets. Also look at the examples below.

For decorated parallel coordinate plots (type="pc") see the specific manual page of seqpcplot.

Author(s)
Alexis Gabadinho (with Gilbert Ritschard for the help page)

References


See Also

plot.stslist.statd, plot.stslist.freq, plot.stslist, plot.stslist.modst, plot.stslist.meant, plot.stslist.rep, seqrep, seqpcplot.

Examples

## --------------------------------------------------------------
## Creating state sequence objects from example data sets
## --------------------------------------------------------------

## biofam data set
data(biofam)
## We use only a sample of 300 cases
set.seed(10)
biofam <- biofam[sample(nrow(biofam),300),]
biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

## actcal data set
data(actcal)
## We use only a sample of 300 cases
set.seed(1)
actcal <- actcal[sample(nrow(actcal),300),]
actcal.lab <- c(">37 hours", "19-36 hours", "1-18 hours", "no work")
actcal.seq <- seqdef(actcal,13:24,labels=actcal.lab)

## ex1 using weights
data(ex1)
ex1.seq <- seqdef(ex1, 1:13, weights=ex1$weights)

## Sequence frequency plots
## seqplot

### Plot of the 10 most frequent sequences

```r
seqplot(biofam.seq, type="f")
```

### Grouped by sex

```r
seqplot(actcal.seq, group=actcal$sex)
```

### Unweighted vs weighted frequencies

```r
seqfplot(ex1.seq, weighted=FALSE)
seqfplot(ex1.seq, weighted=TRUE)
```

### Modal states sequence

```r
seqplot(biofam.seq, type="ms")
```

### Representative plots

```r
seqmsplot(biofam.seq)
```

### Computing a distance matrix

```r
costs <- seqcost(biofam.seq, method="INDELSLOG")
biofam.om <- seqdist(biofam.seq, method="OM", sm=costs$sm, indel=costs$indel)
```

### Plot of the representative sets grouped by sex

```r
seqrplot(biofam.seq, group=biofam$sex, diss=biofam.om)
```

### Plot of the representative sets grouped by sex

```r
seqrplot(biofam.seq, group=biofam$sex, criterion="dist", diss=biofam.om)
```

### Sequence index plots

### First ten sequences

```r
seqiplot(biofam.seq)
```

### All sequences sorted by age in 2000

```r
seqiplot(actcal.seq, group=actcal$sex, idxs=0, border=NA, space=0, sortv=actcal$age00)
```

### State distribution plot
seqpm

Find substring patterns in sequences

Description
Search for a pattern (substring) into sequences.

Usage
seqpm(seqdata, pattern, sep="")

Arguments
seqdata a sequence object as defined by the seqdef function.
pattern a character string representing the pattern (substring) to search for.
sep state separator used in the pattern definition.

Details
This function searches a pattern (a character string) into a set of sequences and returns the results as a list with two elements: 'Nbmatch' the number of occurrences of the pattern and 'MatchesIndex' the vector of indexes (row numbers) of the sequences that match the pattern (see examples below).

Value
a list with two elements (see details).
seqrecode

Author(s)
Alexis Gabadinho

Examples

data(actcal)
actcal.seq <- seqdef(actcal, 13:24)

## search for pattern "DAAD"
## (no work-full time work-full time work-no work)
## results are stored in the 'daad' object
daad <- seqpm(actcal.seq, "DAAD")

## Looking at the sequences
## containing the pattern
actcal.seq[daad$index,]

## search for pattern "AD"
## (full time work-no work)
seqpm(actcal.seq, "AD")

seqrecode

Recoding state sequence objects and factors

Description
Utilities for recoding factors or state sequence objects created with seqdef.

Usage

seqrecode(seqdata, recodes, otherwise = NULL,
  labels = NULL, cpal = NULL)
recodef(x, recodes, otherwise = NULL, na = NULL)

Arguments

seqdata The state sequence object to be recoded (created with seqdef).
recodes A list specifying the recoding operations where each element is in the form
  newcode=oldcode or newcode=c(oldcode1, oldcode2,...). The rules are
  treated in the same order as they appear, hence subsequent rules may modify the
  first ones.
otherwise NULL or Character. Level given to cases uncovered by the recodes list. If NULL,
  old states remain unchanged.
labels optional state labels used for the color legend of TraMineR’s graphics. If NULL
  (default), the state names in the alphabet are also used as state labels (see seqdef).
cpal

an optional color palette for representing the newly defined alphabet in graphics. If NULL (default), a color palette is created from the colors in seqdata by assigning to newcode the color of the first old state listed as oldcode and by leaving the colors of the other states unchanged.

x

A factor to be recoded.

na

Character vector. If not NULL, the list of states that should be recoded as NA (missing values).

Value

The recoded factor or state sequence object.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

See Also

seqdef to create a state sequence object.

Examples

```r
## Recoding a state sequence object with seqrecode
data(actcal)
## Creating a state sequence object
actcal.seq <- seqdef(actcal,13:24, labels=c("> 37 hours", "19-36 hours", "1-18 hours", "no work"))
## Regrouping states B and C and setting the whole alphabet to A BC D
actcal.new <- seqrecode(actcal.seq,
    recodes = list("A"="A", "BC"=c("B", "C"), "D"="D"))
## Crosstabulate the first column of the recoded and original state sequence objects
table(actcal.new[,1], actcal.seq[,1])

## Same as before but using automatically original codes for unspecified states.
actcal.new2 <- seqrecode(actcal.seq,
    recodes = list("BC"=c("B", "C")))
table(actcal.new2[,1], actcal.seq[,1])

## Same as before but using otherwise
actcal.new3 <- seqrecode(actcal.seq, recodes = list("A"="A", "D"="D"), otherwise="BC")
table(actcal.new3[,1], actcal.seq[,1])

## Recoding factors
## Recoding the marital status to oppose married to all other case
maritalstatus <- recode(actcal$civsta00,
    recodes=list("Married"="married"), otherwise="Single")
summary(maritalstatus)
table(maritalstatus, actcal$civsta00)
```
## Recoding the number of kids in the household
## -2 is a missing value

```r
nbkids <- recode(actcal$nbkid00, 
    recodes=list("None"=0, "One"=1, "Two or more"=2:10), na=-2)
table(nbkids, actcal$nbkid00, useNA="always")
```

### seqrep

Extracting sets of representative sequences

**Description**

Returns either an as small as possible set of non redundant representatives covering (having in their neighborhood) a desired percentage of all sequences, or a given number of patterns with highest coverage. Special cases are single representatives such as the medoid or the sequence pattern with densest neighborhood. See `plot.stslist.rep` for the plot method and `seqplot` for other plot options.

**Usage**

```r
seqrep(seqdata, criterion = "density", score = NULL, decreasing = TRUE, 
    coverage = 0.25, nrep = NULL, pradius = 0.10, dmax = NULL, diss = NULL, 
    weighted = TRUE, trep, tsim, dist.matrix, ...)
```

**Arguments**

- **seqdata**: a state sequence object as defined by the `seqdef` function.
- **criterion**: the representativeness criterion for sorting the candidate list. One of "freq" (sequence frequency), "density" (neighborhood density), "mscore" (mean state frequency), "dist" (centrality) and "prob" (sequence likelihood). See details.
- **score**: an optional vector of representativeness scores for sorting the sequences in the candidate list. The length of the vector must be equal to the number of sequences in the sequence object.
- **decreasing**: if a score vector is provided, indicates whether the objects in the candidate list must be sorted in ascending or descending order of this score. Default is TRUE, i.e. descending. The first object in the candidate list is then supposed to be the most representative.
- **coverage**: coverage threshold, i.e., minimum proportion of sequences that should have a representative in their neighborhood (neighborhood radius is defined by `pradius`).
- **nrep**: number of representative sequences. If NULL (default), the size of the representative set is controlled by `coverage`.
- **pradius**: neighborhood radius as a percentage of the maximum (theoretical) distance `dmax`. Defaults to 0.1 (10%). Sequence `y` is redundant to sequence `x` when it is in the neighborhood of `x`, i.e., within a distance `pradius*dmax` from `x`. 

seqrep

dmax  maximum theoretical distance. Used to derive the neighborhood radius as \(pradius\times dmax\). If NULL, the value of \(dmax\) is derived from the dissimilarity matrix.

diss  matrix of pairwise dissimilarities between sequences in seqdata. If NULL, the matrix is computed by calling the seqdist function. In that case, optional arguments to be passed to the seqdist function (see ... hereafter) should also be provided.

weighted logical: Should weights assigned to the state sequence object be accounted for? (See seqdef.) Set as FALSE to ignore the weights.

trep  Deprecated. Use coverage instead.

tsim  Deprecated. Use pradius instead.

distNmatrix  Deprecated. Use diss instead.

... optional arguments to be passed to the seqdist function, mainly dist.method specifying the metric for computing the distance matrix, norm for normalizing the distances, indel and sm for indel and substitution costs when Optimal Matching metric is chosen. See seqdist manual page for details.

Details

The representative set is obtained by an heuristic. Representatives are selected by successively extracting from the sequences sorted by their representativeness score those which are not redundant with already retained representatives. The selection stops when either the desired coverage or the wanted number of representatives is reached. Sequences are sorted either by the values provided as score argument or by specifying one of the following as criterion argument: "freq" (sequence frequency), "density" (neighborhood density), "mscore" (mean state frequency), "dist" (centrality) and "dist" (sequence likelihood).

With the sequence frequency criterion, the more frequent a sequence the more representative it is supposed to be. Therefore, sequences are sorted in decreasing frequency order.

The neighborhood density is the number—density—of sequences in the neighborhood of the sequence. This requires to set the neighborhood radius pradius. Sequences are sorted in decreasing density order.

The mean state frequency criterion is the mean value of the transversal frequencies of the successive states. Let \(s = s_1 s_2 \cdots s_\ell\) be a sequence of length \(\ell\) and \((f_{s_1}, f_{s_2}, \ldots, f_{s_\ell})\) the frequencies of the states at (time-)position \((t_1, t_2, \ldots, t_\ell)\). The mean state frequency is the sum of the state frequencies divided by the sequence length

\[
MSF(s) = \frac{1}{\ell} \sum_{i=1}^{\ell} f_{s_i}
\]

The lower and upper boundaries of \(MSF\) are 0 and 1. \(MSF\) is equal to 1 when all the sequences in the set are identical, i.e. when there is a single sequence pattern. The most representative sequence is the one with the highest score.

The centrality criterion is the sum of distances to all other sequences. The smallest the sum, the most representative the sequence.

The sequence likelihood \(P(s)\) is defined as the product of the probability with which each of its observed successive state is supposed to occur at its position. Let \(s = s_1 s_2 \cdots s_\ell\) be a sequence of length \(\ell\). Then

\[
P(s) = P(s_1, 1) \cdot P(s_2, 2) \cdots P(s_\ell, \ell)
\]
with \( P(s_t, t) \) the probability to observe state \( s_t \) at position \( t \).

The question is how to determinate the state probabilities \( P(s_t, t) \). One commonly used method for computing them is to postulate a Markov Chain model, which can be of various order. The implemented criterion considers the probabilities derived from the first order Markov model, that is each \( P(s_t, t) \), \( t > 1 \) is set to the transition rate \( p(s_t|s_{t-1}) \) estimated across sequences from the observations at positions \( t \) and \( t-1 \). For \( t = 1 \), we set \( P(s_1, 1) \) to the observed frequency of the state \( s_1 \) at position 1.

The likelihood \( P(s) \) being generally very small, we use \(-\log P(s)\) as sorting criterion. The latter quantity reaches its minimum for \( P(s) \) equal to 1, which leads to sort the sequences in ascending order of their score.

Use criterion="dist" and nrep=1 to get the medoid and criterion="density" and nrep=1 to get the densest sequence pattern.

For more details, see Gabadinho & Ritschard, 2013.

### Value

An object of class \texttt{stslistNrep}. This is actually a state sequence object (containing a list of state sequences) with the following additional attributes:

- **Scores**
  - a vector with the representative score of each sequence in the original set given the chosen criterion.

- **Distances**
  - a matrix with the distance of each sequence to its nearest representative.

- **Statistics**
  - a data frame with quality measures for each representative sequence: number \( na \) of sequences attributed to the representative, number \( nb \) of sequences in the representative’s neighborhood, mean distance \( MD \) to the representative and a few other indexes.

- **Quality**
  - overall quality measure.

Print, plot and summary methods are available. More elaborated plots are produced by the \texttt{seqplot} function using the \texttt{type="r"} argument, or the \texttt{seqrplot} alias.

### Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

### References


### See Also

\texttt{seqplot}, \texttt{plot.stslistNrep}, \texttt{dissrep}, \texttt{disscenter}
Examples

```r
## Defining a sequence object with the data in columns 10 to 25
## (family status from age 15 to 30) in the biofam data set
data(biofam)
biofam.lab <- c("Parent", "Left", "Married", "Left+Marr",
"Child", "Left+Child", "Left+Marr+Child", "Divorced")
biofam.seq <- seqdef(biofam, 10:25, labels=biofam.lab)

## Computing the distance matrix
costs <- seqsubm(biofam.seq, method="TRATE")
biofam.om <- seqdist(biofam.seq, method="OM", sm=costs)

## Representative set using the neighborhood density criterion
biofam.repl <- seqrep(biofam.seq, diss=biofam.om, criterion="density")
biofam.rep
summary(biofam.rep)
plot(biofam.rep)
```

seqsep

Adds separators to sequences stored as character string.

Description

Adds separators to sequences stored as character string.

Usage

```r
seqsep(seqdata, sl=1, sep="-")
```

Arguments

- `seqdata`: a dataframe or matrix containing sequence data, as vectors of states or events.
- `sl`: the length of the states (the number of characters used to represent them). Default is 1.
- `sep`: the character used as separator. Set by default as "-".

See Also

seqdecomp.

Examples

```r
seqsep("ABAAAAAAD")
```
**Description**

Computes Elzinga’s turbulence for each sequence in a sequence data set.

**Usage**

```r
dseqST(seqdata, norm=FALSE, silent=TRUE)
```

**Arguments**

- `seqdata`: a state sequence object as returned by the `seqdef` function.
- `norm`: logical: should the turbulence index be normalized?
- `silent`: logical: should messages about running operations (extracting dss and durations, computing turbulence) be displayed?

**Details**

Sequence turbulence is a measure proposed by Elzinga & Liefbroer (2007). It is based on the number $\phi(x)$ of distinct subsequences that can be extracted from the distinct successive state (DSS) sequence and the variance of the consecutive times $t_i$ spent in the distinct states. For a sequence $x$, the formula is

$$T(x) = \log_2(\phi(x) \frac{s_{t,\text{max}}^2(x) + 1}{s_t^2(x) + 1})$$

where $s_t^2(x)$ is the variance of the successive state durations in sequence $x$ and $s_{t,\text{max}}^2(x)$ is the maximum value that this variance can take given the total duration of the sequence. This maximum is computed as

$$s_{t,\text{max}}^2 = (d-1)(1-\bar{t})^2$$

where $\bar{t}$ is the mean consecutive time spent in the distinct states, i.e. the sequence duration divided by the number $d$ of distinct states in the sequence.

The function searches for missing states in the sequences and if found, adds the missing state to the alphabet for the computation of the turbulence. In this case the `seqdss` and `seqdur` functions for extracting the distinct successive state sequences and the associated durations are called with the `{with.missing=TRUE}` argument. Thus, a missing state in a sequence is considered as the occurrence of an additional symbol of the alphabet and two or more consecutive missing states are considered as two or more occurrences of this additional state. E.g. the DSS of `A-A++-++-B-B-C-C-D` is `A--B-C-D` and the associated durations are `2-2-2-2-1`.

The normalized value is obtained by subtracting 1 to the index and dividing by the turbulence value of a sequence made by the successive repetition of the alphabet up to the maximal length in `seqdata`. 
Value

A single-column matrix of length equal to the number of sequences in seqdata containing the turbulence value of each sequence. Normalized values are returned when norm=TRUE.

Author(s)

Alexis Gabadinho and Gilbert Ritschard

References


See Also

seqdss, seqdur. For other measures of sequence complexity see seqient and seqici.

Examples

```r
## Loading the 'actcal' example data set
data(actcal)
## Here we consider only the first 10 sequences
actcal <- actcal[1:10,]

## Defining a sequence object with data in columns 13 to 24
## (activity status from January to December 2000)
actcal.seq <- seqdef(actcal[,13:24], informat='STS')

## Computing the sequences turbulence
turb <- seqST(actcal.seq)

## Histogram for the turbulence
hist(turb)

## Normalized turbulence
turb.norm <- seqST(actcal.seq, norm=TRUE)
```

seqstatd

Sequence of transversal state distributions and their entropies

Description

Returns the state frequencies, the number of valid states and the entropy of the state distribution at each position in the sequence.
seqstatd

Usage

`seqstatd(seqdata, weighted=TRUE, with.missing=FALSE, norm=TRUE)`

Arguments

- **seqdata**: a state sequence object as defined by the `seqdef` function.
- **weighted**: if TRUE, distributions account for the weights assigned to the state sequence object (see `seqdef`). Set as FALSE if you want ignore the weights.
- **with.missing**: If FALSE (default value), returned distributions ignore missing values.
- **norm**: if TRUE (default value), entropy is normalized, ie divided by the entropy of the alphabet. Set as FALSE if you want the entropy without normalization.

Details

In addition to the state distribution at each position in the sequence, the `seqstatd` function provides also for each time point the number of valid states and the Shannon entropy of the observed state distribution. Letting $p_i$ denote the proportion of cases in state $i$ at the considered time point, the entropy is

$$h(p_1, \ldots, p_s) = - \sum_{i=1}^{s} p_i \log(p_i)$$

where $s$ is the size of the alphabet. The log is here the natural (base e) logarithm. The entropy is 0 when all cases are in the same state and is maximal when the same proportion of cases are in each state. The entropy can be seen as a measure of the diversity of states observed at the considered time point. An application of such a measure (but with aggregated transversal data) can be seen in Billari (2001) and Fussell (2005).

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

References


See Also

- `plot.stslist.statd` the plot method for objects of class `stslist.statd`
- `seqdplot` for higher level plot of transversal distributions and
- `seqHtplot` for plotting the transversal entropy over sequence positions.
**Examples**

```r
data(biofam)
biofam.seq <- seqdef(biofam, 10:25)
sd <- seqstatd(biofam.seq)
## Plotting the state distribution
plot(sd, type="d")

## Plotting the entropy indexes
plot(sd, type="Ht")

## ================
## example with weights
## ================
data(ex1)
ex1.seq <- seqdef(ex1, 1:13, weights=ex1$weights)

## Unweighted
seqstatd(ex1.seq, weighted=FALSE)
seqstatd(ex1.seq, weighted=TRUE)
```

---

**seqstatf**  
*State frequencies in the whole sequence data set*

**Description**

Overall frequency of each state of the alphabet in the state sequence object.

**Usage**

```r
seqstatf(seqdata, weighted = TRUE)
```

**Arguments**

- `seqdata`  
a sequence object as defined by the `seqdef` function.

- `weighted`  
Logical. Should frequencies account for weights when present in the state sequence object (see `seqdef`). Default is `TRUE`. If no weights were assigned during the creation of the sequence object, `weighted=TRUE` will yield the same result as `weighted=FALSE` since each sequence is allowed a weight of 1.

**Details**

The `seqstatf` function computes the (weighted) count and frequency of each state of the alphabet in `seqdata`, i.e., the (weighted) sum of the occurrences of a state in `seqdata`.

**Value**

A data frame with as many rows as states in the alphabet and two columns, one for the count (Freq) and one for the percentage frequencies (Percent).
seqstatl

List of distinct states or events (alphabet) in a sequence data set.

Description

Returns a list containing distinct states or events found in a data frame or matrix containing sequence data, the alphabet.

Usage

seqstatl(data, var=NULL, format='STS')

Arguments

data a data frame or matrix containing sequence data.
var the list of columns containing the sequences. Default NULL means all columns. Whether the sequences are in the compressed (character strings) or extended format is automatically detected from the number of columns.
format the format of the sequence data set. One of "STS", "SPS", "DSS". Default is "STS". The seqstatl function uses the seqformat function to translate between formats when necessary.
Author(s)
Alexis Gabadinho

References

See Also
seqformat

Examples
```r
data(actcal)
seqstatl(actcal, 13:24)
```

<table>
<thead>
<tr>
<th>seqsubsn</th>
<th>Number of distinct subsequences in a sequence.</th>
</tr>
</thead>
</table>

Description
Computes the number of distinct subsequences in a sequence using Elzinga’s algorithm.

Usage
```r
seqsubsn(seqdata, DSS=TRUE)
```

Arguments
- **seqdata**
  a state sequence object as defined by the `seqdef` function.
- **DSS**
  if TRUE, the sequences of Distinct Successive States (DSS, see `seqdss`) are first extracted (e.g., the DSS contained in 'D>D>D=A>A-A-A-A-A-A-D' is 'D=A-D'), and the number of distinct subsequences in the DSS is computed. If FALSE, the number of distinct subsequences is computed from sequences as they appear in the input sequence object. Hence the number of distinct subsequences is in most cases much higher with the DSS=FALSE option.

Details
The function first searches for missing states in the sequences and if found, adds the missing state to the alphabet for the extraction of the distinct subsequences. A missing state in a sequence is considered as the occurrence of an additional symbol of the alphabet, and two or more consecutive missing states are considered as two or more occurrences of the same state. The `with.missing=TRUE` argument is used for calling the `seqdss` function when DSS=TRUE.
Value

Vector with the number of distinct subsequences for each sequence in the input state sequence object.

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

See Also

seqdss.

Examples

data(actcal)
actcal.seq <- seqdef(actcal,13:24)

### Number of subsequences with DSS=TRUE
seqsubsn(actcal.seq[1:10,])

### Number of subsequences with DSS=FALSE
seqsubsn(actcal.seq[1:10,],DSS=FALSE)

seqtab Frequency table of the sequences

Description

Computes the frequency table of the sequences (count and percent of each sequence).

Usage

seqtab(seqdata, idxs = 1:10, weighted = TRUE, format = "SPS", tlim)

Arguments

seqdata a sequence object as defined by the seqdef function.
idxs returns the table for the sequences at ranks ’idxs’ in the list of distinct sequences sorted in decreasing order of their frequencies. Default is 1:10, i.e. the 10 most frequent sequences. Can be any subset, like 5:10 (fifth to tenth most frequent sequences) or c(2,10) (second and tenth most frequent sequences). Set idxs=0 to get the table for the whole set of distinct sequences.
weighted if TRUE (default), frequencies account for the weights, if any, assigned to the state sequence object (see seqdef). Set to FALSE for ignoring weights.
format format used for displaying the rownames (the sequences) in the output table. Default is SPS format, which yields shorter and more readable sequence representations. Alternatively, "STS" may be specified.
tlim Deprecated. Use idxs instead.
Details

The weighted argument has no effect when no weights were assigned to the state sequence object since weights default in that case to 1.

Value

An object of class stslist.freq. This is actually a state sequence object (containing a list of state sequences) with added attributes, among others the freq attribute containing the frequency table. There are print and plot methods for such objects. More sophisticated plots can be produced with the seqplot function.

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

References


See Also

seqplot, plot.stslist.freq.

Examples

```r
## Creating a sequence object from the actcal data set
data(actcal)
actcal.lab <- c("> 37 hours", "19-36 hours", "1-18 hours", "no work")
actcal.seq <- seqdef(actcal, 13:24, labels=actcal.lab)

## 10 most frequent sequences in the data
seqtab(actcal.seq)

## With idxs=0, we get all distinct sequences in the data set
## sorted in decreasing order of their frequency
stab <- seqtab(actcal.seq, idxs=0)
head(stab)
tail(stab)

## Example with weights
## from biofam data set using weights
data(ex1)
ex1.seq <- seqdef(ex1, 1:13, weights=ex1$weights)

## Unweighted frequencies
seqtab(ex1.seq, weighted=FALSE)

## Weighted frequencies
seqtab(ex1.seq, weighted=TRUE)
```
Description

Computes the number of transitions in each sequence of a sequence object.

Usage

seqtransn(seqdata, with.missing=FALSE, norm=FALSE, pweight=FALSE)

Arguments

seqdata  a state sequence object as defined by the seqdef function.
with.missing  logical. if set as TRUE, missing states (gaps in sequences) are considered as an additional state and included in the DSS sequence. See seqdss.
norm  logical. If set as TRUE, the number of transitions is divided by its theoretical maximum, the length of the sequence minus 1. When length of the sequence is 1, normalized value is set to 0 as in the non-normalized case.
pweight  logical. EXPERIMENTAL! If set as TRUE, when counting transitions each transition does not account for 1 but for its probability (transition rate) as observed in the data.

Details

A transition in a sequence is a state change between time/position \(t\) and \(t + 1\). For example, the sequence "A-A-A-B-A-D-D-D" contains 3 transitions. The maximum number of transitions a sequence can contain is \(t - 1\) where \(t\) is the length of the sequence. The number of transitions is obtained by subtracting 1 to the length of the the Distinct Successive State (DSS) sequence.

Value

a state sequence object containing the number of transitions of each sequence in the object given as argument.

Author(s)

Alexis Gabadinho (with Gilbert Ritschard for the help page)

References


See Also

seqdss.
Examples

```r
## Creating a sequence object from columns 13 to 24
## in the 'actcal' example data set
data(actcal)
actcal.seq <- seqdef(actcal, 13:24)

## Computing the number of transitions
actcal.trans <- seqtransn(actcal.seq)

## Displaying the DSS for the first 10 sequences
actcal.trans[1:10]

## Example with with.missing argument
data(ex1)
ex1.seq <- seqdef(ex1, 1:13)

seqtransn(ex1.seq)
seqtransn(ex1.seq, with.missing=TRUE)
```

---

**seqtrate**

*Compute transition rates between states*

**Description**

Returns a matrix with transition rates between states, computed from a set of sequences.

**Usage**

```r
seqtrate(seqdata, sel.states = NULL, time.varying = FALSE, weighted = TRUE,
          lag = 1, with.missing = FALSE, count = FALSE, statl)
```

**Arguments**

- `seqdata`  
  a sequence object as defined by the `seqdef` function.

- `sel.states`  
  a list of states or events for which the transition rates will be computed. If omitted (default), transition rates are computed between the distinct states in `seqdata` (obtained with the `alphabet` function).

- `time.varying`  
  Logical. If TRUE, return an array containing a distinct matrix for each time unit. The time is the third dimension (subscript).

- `weighted`  
  Logical. If TRUE, compute transition rates using weights specified in `seqdata`.

- `lag`  
  Integer. Time between the two states considered to compute transition rates (one by default).

- `with.missing`  
  Logical. If FALSE (default value), returned transition rates ignore missing values.

- `count`  
  Logical. Should counts of transition be returned instead of transition probabilities. Default is FALSE.

- `statl`  
  Deprecated. Use `sel.states` instead.
Details

Transition rates are the probabilities of transition from one state to another observed in the sequence data. Substitution costs based on transition rates can be used when computing distances between sequences with the optimal matching method (see seqdist).

Value

a matrix of dimension \(ns \times ns\), where \(ns\) is the number of states in the alphabet of the sequence object.

Author(s)

Matthias Studer, Alexis Gabadinho, and Gilbert Ritschard

References


See Also

seqdist seqsubm alphabet.

Examples

```r
## Loading the 'actcal' example data set
data(actcal)

## Defining a sequence object with data in columns 13 to 24
## (activity status from January to December 2000)
actcal.seq <- seqdef(actcal[,13:24])

## Computing transition rates
seqtrate(actcal.seq)

## Computing transition rates between states "A" and "B" only
seqtrate(actcal.seq, c("A","B"))

## ================
## Example with weights
## ================
data(ex1)
ex1.seq <- seqdef(ex1[,1:13], weights=ex1$weights)

seqtrate(ex1.seq, weighted=FALSE)
seqtrate(ex1.seq, weighted=FALSE, count=TRUE)

## weights are accounted for by default
seqtrate(ex1.seq)
seqtrate(ex1.seq, count=TRUE)
```
**seqtree**

Tree structured analysis of a state sequence object.

**Description**

Facility for growing a regression tree for a state sequence object.

**Usage**

```r
seqtree(formula, data = NULL, weighted = TRUE, min.size = 0.05,
         max.depth = 5, R = 1000, pval = 0.01, weight.permutation = "replicate",
         seqdist.args = list(method = "LCS", norm = "auto"), diss = NULL,
         squared = FALSE, first = NULL, minSize, maxdepth, seqdist_arg)
```

**Arguments**

- `formula`: a formula where the left hand side is a state sequence object (see `seqdef`) and the right hand specifies the candidate variables for partitioning the set of sequences.
- `weighted`: Logical. If `TRUE`, use the weights of the state sequence object.
- `data`: a data frame where variables in the formula will be searched
- `min.size`: minimum number of cases in a node, in percentage if less than 1.
- `max.depth`: maximum depth of the tree.
- `R`: Number of permutations used to assess the significance of the split.
- `pval`: Maximum p-value, in percent.
- `weight.permutation`: Weights permutation method: "diss" (attach weights to the dissimilarity matrix), "replicate" (replicate case according to the `weights` arguments), "rounded-replicate" (replicate case according to the rounded `weights` arguments), "random-sampling" (random assignment of covariate profiles to the objects using distributions defined by the weights.)
- `seqdist.args`: list of arguments directly passed to `seqdist`, only used if `diss=NULL`
- `diss`: An optional dissimilarity matrix. If not provided, a dissimilarity matrix is computed using `seqdist` and `seqdist.args`
- `squared`: Logical. If `TRUE`, the dissimilarity matrix is squared
- `first`: Character. An optional variable name to force the first split.
- `minSize`: Deprecated. Use `min.size` instead.
- `maxdepth`: Deprecated. Use `max.depth` instead.
- `seqdist_arg`: Deprecated. Use `seqdist.args` instead.

**Details**

The function provides a simplified interface for applying `distree` on state sequence objects. The `seqtree` objects can be "plotted" with `seqtreedisplay`. A print method is also available which prints the medoid sequence for each terminal node.
Value

A seqtree object with same attributes as `disstree` objects.

The leaf membership is in the first column of the fitted attribute. For example, the leaf memberships for a tree `dt` are in `dt$fitted[,1]`.

Author(s)

Matthias Studer (with Gilbert Ritschard for the help page)

References


See Also

`seqtreedisplay`, `disttree`

Examples

data(mvad)

```r
## Defining a state sequence object
mvad.seq <- seqdef(mvad[, 17:86])

## Growing a seqtree from Hamming distances:
## Warning: The R=10 used here to save computation time is
## much too small and will generate strongly unstable results.
## We recommend to set R at least as R=1000.
## To comply with this small R value, we set pval = 0.1.
set.seed(123)
seqt <- seqtree(mvad.seq ~ male + Grammar + funemp + gcseSeq + fmpr + livboth, 
  data=mvad, R=10, pval=0.1, seqdist.arg=list(method="HAM", norm="auto"))
print(seqt)

## Growing a seqtree from an existing distance matrix
mvad.dhd <- seqdist(mvad.seq, method="DHD")
seqt <- seqtree(mvad.seq ~ male + Grammar + funemp + gcseSeq + fmpr + livboth, 
  data=mvad, R=10, pval=0.1, diss=mvad.dhd)
print(seqt)

## Following commands only work if GraphViz is properly installed
## Not run:
seqtreedisplay(seqt, type="d", border=NA)
seqtreedisplay(seqt, type="I", sortv=cmdscale(mvad.dhd, k=1))
```

## End(Not run)
seqtreedisplay  

Graphical rendering of a sequence regression tree

Description

Generate a graphical representation of a regression tree of state sequence data.

Usage

```
seqtreedisplay(tree, filename = NULL, seqdata = tree$info$object,
only.leaf = FALSE, sortv = NULL, diss = NULL, cex.main = 3,
with.legend = "auto", cex.legend = cex.main, axes = FALSE,
image.format = "png", with.quality = TRUE, cex.quality = cex.main,
legend.text = NULL, show.tree = TRUE, show.depth = FALSE,
imgLeafOnly, dist.matrix, title.cex, withlegend, legend.fontsize,
imageformat, withquality, quality.fontsize, legendtext, showtree,
showdepth, ...)
```

```
disstreedisplay(tree, filename = NULL, image.data=NULL, image.fun=plot,
only.leaf = FALSE, cex.main = 3, image.format = "png",
with.quality = TRUE, cex.quality = cex.main,
legend.text = NULL, show.tree = TRUE, show.depth = FALSE,
imagedata, imagefunc, imgLeafOnly, title.cex, imageformat,
withquality, quality.fontsize, legendtext, showtree, showdepth, ...)
```

Arguments

- **tree**: A seqtree object (as produced by `seqtree`) for `seqtreedisplay`. A distree object (as produced by `disstree`) for `disstreedisplay`.
- **filename**: The name of a file where to save the plot (overwriting existing file). If NULL, a temporary file is created.
- **seqdata**: The sequence object containing the state sequences plotted in the nodes.
- **only.leaf**: Logical. If TRUE sequences are plotted only in terminal nodes.
- **sortv**: Argument passed to `seqplot`
- **diss**: Argument passed to `seqplot`
- **cex.main**: Node title magnification. See `par`.
- **with.legend**: Logical. Should the color legend be displayed on the plot?
- **cex.legend**: Legend magnification. See `par`. If not specified, use the value of `cex.main`.
- **axes**: Argument passed to `seqplot`
- **image.format**: Image format of the output file (filename)
- **with.quality**: If TRUE, a node displaying fitting measures of the tree is added to the plot.
- **cex.quality**: Fitting measure text magnification. See `par`. If not specified, use the value of `cex.main`. 


seqtreedisplay

```r
legend.text  Character. Optional text information that should be added.
show.tree   Logical. Should the tree be shown on the screen?
show.depth  Logical. If TRUE, the splits are ordered according to their global pseudo-R2.
image.fun   A function to plot the individuals in a node, see details.
image.data  a data.frame that will be passed to image.fun.
imgLeafOnly Deprecated. Use only.leaf instead.
dist.matrix Deprecated. Use diss instead.
title.cex   Deprecated. Use cex.main instead.
withlegend  Deprecated. Use with.legend instead.
legendfontsize Deprecated. Use cex.legend instead.
imageformat Deprecated. Use image.format instead.
withquality  Deprecated. Use with.quality instead.
quality.fontsize Deprecated. Use cex.quality instead.
legendtext  Deprecated. Use legend.text instead.
showtree    Deprecated. Use show.tree instead.
showdepth   Deprecated. Use show.depth instead.
imagedata   Deprecated. Use image.data instead.
imagefunc   Deprecated. Use image.fun instead.
...        additional arguments passed to seqplot
```

**Details**

This function generates a tree image. For each node, it invokes `seqplot` for the selected lines of `seqdata` as argument. You should at least specify the type of the plot to use (type="d" for instance, see `seqplot` for more details).

The plot is actually not generated as an R plot, but with GraphViz (www.graphviz.org). Hence, `seqtreedisplay` only works when GraphViz is correctly installed.

Conversion to image formats other than "jpeg" or "png" is done using ImageMagick (www.imagemagick.org). To use this feature, ImageMagick (www.imagemagick.org) should hence also be installed.

**Value**

None

**Author(s)**

Matthias Studer (with Gilbert Ritschard for the help page)

**See Also**

See `seqtree` and `disstree` for examples, and `disstree2dot` for generating "dot" files.
stlab

Get or set the state labels of a sequence object

Description

This function gets or sets the state labels of a sequence object, that is, the long labels used when displaying the state legend in plotting functions.

Usage

stlab(seqdata)
stlab(seqdata) <- value

Arguments

seqdata a state sequence object as defined by the seqdef function.
value a vector of character strings containing the labels, of length equal to the number of states in the alphabet. Each string is attributed to the corresponding state in the alphabet, the order being the one returned by the alphabet.

Details

The state legend is plotted either automatically by the plot functions provided for visualizing sequence objects or with the seqlegend function. A long label is associated to each state of the alphabet and displayed in the legend. The state labels are defined when creating the sequence object, either automatically using the values found in the data or by specifying a user defined vector of labels. The stlab function can be used to get or set the state labels of a previously defined sequence object.

Value

For 'stlab' a vector containing the labels.
For 'stlab<-’ the updated sequence object.

See Also

seqdef

Examples

## Creating a sequence object with the columns 13 to 24
## in the 'actcal' example data set
## The color palette is automatically set
data(actcal)
actcal.seq <- seqdef(actcal,13:24)

## Retrieving the color palette
stlab(actcal.seq)
seqiplot(actcal.seq)

## Changing the state labels
stlab(actcal.seq) <- c("Full time","Part time (19-36 hours)",
  "Part time (1-18 hours)", "No work")
seqiplot(actcal.seq)

---

**TraMineR.check.depr.args**

*Checking and managing deprecated arguments*

**Description**

Checks the presence of deprecated arguments, assigns value of a deprecated argument to the corresponding new argument name, and issues warning messages.

**Usage**

```r
TraMineR.check.depr.args(arg.pairs)
```

**Arguments**

- `arg.pairs` List of pairs of old and new argument names
  (e.g. `alist(newname1 = oldname1, newname2 = oldname2)`)  

**Details**

To be used inside functions. For developers only.

For each specified pair of new and old argument names, the function checks if the old argument name is specified. If so and the new one is not, a warning message is raised and the argument value is assigned to the new argument name. If one of the names declared in check.depr.args() arguments is not an argument of the parent function or if both the new and old argument names are specified an error is raised.

The function does not detect when the new and the old argument names are specified together and the new argument value is its default value. In this case, the value associated with the old argument name is assigned to the new name and a warning message is raised.

The function works whether the argument names are explicitly declared or not in the call to the checked function.

The only requirement for the function to work is that the deprecated arguments should be listed WITHOUT default values in the definition of the checked function.

**Value**

None.

**Author(s)**

Pierre-Alexandre Fonta, Gilbert Ritschard
TraMineR.checkupdates  Check for TraMineR updates

**Description**

Check if the installed version of TraMineR is up-to-date. This function only prints a message and does not need any argument. It connects to the TraMineR webserver ([http://traminer.unige.ch](http://traminer.unige.ch)).

**Usage**

```r
TraMineR.checkupdates()
```

**Value**

Return your current version number of TraMineR and the latest stable and development version number if more recent versions are available.

**Author(s)**

Nicolas S. Müller

---

TraMineRInternal  Access to TraMineR internal functions

**Description**

Functions allowing other packages to access some TraMineR internal functions. Corresponding functions are respectively `TraMineR.setlayout`, `TraMineR.Legend`, `DTNInit`, `seqeage`, `seqgbar`, `DTNsplt`, and `tmrWeightedInertiaDist`. For experts only.

**Usage**

```r
TraMineRInternalLayout(...)  
TraMineRInternalLegend(...)  
TraMineRInternalNodeInit(...) 
TraMineRInternalSeqeage(...) 
TraMineRInternalSeqgbar(...) 
TraMineRInternalSplitInit(...) 
TraMineRInternalWeightedInertiaDist(diss, diss.size, is.dist, individuals, sweights, var)
```
Arguments

... Arguments passed to or from other methods.
diss See tmrWeightedInertiaDist().
diss.size See tmrWeightedInertiaDist().
is.dist See tmrWeightedInertiaDist().
individuals See tmrWeightedInertiaDist().
sweights See tmrWeightedInertiaDist().
var See tmrWeightedInertiaDist().
## Index

### Data handling
- read.tda.mdist, 45
- seqcomp, 47
- seqconc, 48
- seqdecomp, 52
- seqdef, 53
- seqecreate, 76
- seqetm, 82
- seqfind, 85
- seqformat, 86
- seagen, 92
- seqnum, 106
- seqrecode, 119
- seqsep, 124
- seqstatl, 129

### Datasets
- actcal, 5
- actcal.tse, 6
- bfspell, 8
- biofam, 9
- ex1, 28
- ex2, 29
- famform, 30
- mvad, 30

### Dissimilarity measures
- seqcost, 49
- seqdist, 60
- seqdistmc, 66
- seqLLCP, 99
- seqLLCS, 100
- seqmpos, 105

### Dissimilarity-based analysis
- dissassoc, 12
- disscenter, 14
- dissmfacw, 16
- seqrep, 18
- distree, 21
- distree2dot, 23
- distreeleaf, 26
- dissvar, 27
- plot.seqdiff, 32
- seqalign, 46
- seqdiff, 57
- seqrep, 121
- seqltree, 136
- seqtree.display, 138

### Event sequences
- plot.subseqelist, 43
- plot.subseqelistchisq, 44
- seqeapplysub, 70
- seqecmpgroup, 72
- seqeconstraint, 73
- seqecontain, 75
- seqecreate, 76
- seqefsub, 78
- seqefid, 81
- seqelength, 81
- seqetm, 82
- seqeweight, 84
- seqpcplot, 107

### Global characteristics
- seqmeant, 102
- seqstatf, 128
- seqtrate, 134

### Longitudinal characteristics
- seqdss, 68
- seqdur, 69
- seqelength, 81
- seqfpos, 91
- seqici, 93
- seqient, 94
- seqistatd, 96
- seqlength, 98
- seqlogp, 101
- seqST, 125
- seqsubsn, 130
- seqtransn, 133

### Plot
- 144
disttree2dot, 23
plot.seqdiff, 32
plot stslist, 33
plot stslist.freq, 35
plot stslist.meant, 37
plot stslist.modst, 38
plot stslist.rep, 39
plot stslist.statd, 39
plot subseqelist, 43
plot subseqelistschisq, 44
seqlegend, 97
seqpcplot, 107
seqplot, 112
seqtreedisplay, 138

*Topic Sequence-object attributes
alphabet, 7
cpal, 11
seqdim, 59
seqeid, 81
seqeweight, 84
stlab, 140

*Topic State sequences
seqdef, 53
seqfind, 85
seqgen, 92
seqici, 93
seqient, 94
seqistatd, 96
seqlopp, 101
seqnum, 106
seqpm, 118
seqstatf, 128

*Topic Transversal characteristics
seqmodst, 104
seqstatd, 126
seqtab, 131

*Topic package
TraMineR-package, 4
TraMineR.checkupdates, 142

actcal, 5, 6
actcal.tse, 6
alphabet, 7, 11, 50, 91, 107, 134, 135, 140
alphabet<- (alphabet), 7
array, 50, 101, 134

barplot, 43, 45
bfspell20 (bfspell), 8
bfspell, 8

bfscheckupdates, 142
biofam, 8, 9
colors, 11, 55
cpal, 11
cpal<- (cpal), 11
daisy, 50
dissassoc, 12, 15, 17, 18, 22, 28, 58
disscenter, 14, 14, 18, 20, 22, 28, 123
disseqfacw, 12–15, 16, 22, 28
dissrep, 18, 123
dissstree, 13, 15, 18, 21, 26–28, 136–139
dissstree2dot, 21, 23, 139
dissstree2dotp (distree2dot), 23
dissstreealisp, 22, 26
dissstreedisplay (seqtreedisplay), 138
dissstreeleaf, 26
dissvar, 13, 15, 18, 22, 27
dist, 12, 14, 18, 27, 61, 66

ex1, 28
ex2, 29

famform, 30

getwd, 26
gower_matrix (disseqfacw), 16

hist.dissassoc (dissassoc), 12
is.seqelist, 80
is.subseqelist (seqefsub), 78

layout, 114
legend, 32, 98, 114
lines, 32

mvad, 30

order, 77

par, 34, 36, 37, 39, 40, 42, 43, 46, 108, 109,
113, 114, 138
pdf, 34, 115
plot.eseq (seqpcplot), 107
plot.seqalign (seqalign), 46
plot.seqdiff, 32
plot.seqelist (seqpcplot), 107
plot.tslist, 33, 56, 114–116
plot.tslist.freq, 35, 114–116, 132
plot.stslist.meant, 37, 103, 115, 116
plot.stslist.modst, 38, 104, 115, 116
plot.stslist.rep, 39, 115, 116, 121, 123
plot.stslist.statd, 41, 115, 116, 127
plot.subseqelist, 43, 80
plot.subseqelistchisq, 44, 73
png, 34, 115
postscript, 34, 115
print.dissassoc (dissassoc), 12
print.dissmultifactor (dissmfacw), 16
print.distree (distree), 21
print.seqalign (seqalign), 46
print.seqdiff (seqdiff), 57
print.sequeconstraint (sequeconstraint), 73
print.stslist (seqdef), 53
print.subseqelist (seqefsub), 78
read.tda.mdist, 45
recode (seqrecode), 119
rownames, 34
runif, 92

seqalign, 46
seqcomp, 47
seqconc, 48, 53, 86
seqcost, 49, 61, 63, 64
seqdecomp, 49, 52, 86, 124
seqdef, 4, 7, 8, 11, 25, 32–34, 37, 46, 47, 49,
  51, 53, 57, 60, 62–64, 66–69, 76, 77,
  83, 85, 88, 89, 91, 93, 95–98, 103,
  104, 106–108, 110, 113, 114,
  118–122, 125, 127, 128, 130, 131,
  133, 134, 136, 140
seqdiff, 32, 33, 57
seqdim, 59
seqdist, 14, 46, 47, 49, 51, 58, 60, 66, 67,
  100, 122, 135, 136
seqdistmc, 64, 66
seqdplot, 127
seqdplot (seqplot), 112
seqdss, 68, 69, 70, 94, 125, 126, 130, 131, 133
seqdur, 68, 69, 125, 126
seqeapplysub, 70, 73, 75, 77, 79, 80
seqecmpgroup, 44, 45, 72, 77, 84
sequeconstraint, 70, 72, 73, 79
sequecontain, 75

seqecreate, 4, 7, 56, 71, 76, 76, 80, 81, 83,
  108, 110
seqefsub, 43, 70, 72, 73, 75–77, 78, 84
seqid, 81
seqlength, 77, 81
seqlength<- (seqlength), 81
seqtm, 76, 77, 82, 86
seqeweight, 77, 79, 84
seqeweight<- (seqeweight), 84
seqfcheck, 54, 86
seqfind, 47, 85
seqformat, 6, 8, 54–56, 77, 83, 86, 129, 130
seqfplot (seqplot), 112
seqfpos, 47, 91
seqgen, 92
seqhtplot, 127
seqhtplot (seqplot), 112
seqic, 93, 96, 126
seqic, 94, 94, 126
seqiplot (seqplot), 112
seqiplot (seqplot), 112
seqistatd, 96, 129
seqlegend, 97, 140
seqlength, 98
seqLLCP, 99, 105
seqLLCS, 100, 105
seqlogp, 101
seqmean, 37, 102, 115
seqmodst, 39, 104, 115
seqpos, 105
seqmsplot (seqplot), 112
seqmtplot, 37, 103
seqmtplot (seqplot), 112
seqnum, 106
seqpcfilter (seqpcplot), 107
seqpcplot, 107, 115, 116
seqplot, 25, 26, 34, 36, 37, 39, 41, 42, 56,
  103, 104, 107, 110, 112, 121, 123,
  132, 138, 139
seqpm, 47, 118
seqrecode, 119
seqrep, 20, 39, 40, 115, 116, 121
seqrplot, 25, 41, 123
seqrplot (seqplot), 112
seqsep, 124
seqST, 94, 96, 125
seqstatd, 41, 42, 94, 96, 114, 115, 126, 129
seqstatf, 128
INDEX

seqstatl, 54, 55, 129
seqsubm, 63, 64, 66, 67, 135
seqsubm (seqcost), 49
seqsubsn, 130
seqtab, 36, 114, 115, 131
seqtransn, 133
seqtree, 50, 51, 134
seqtree, 21, 22, 26, 136, 138, 139
seqtree2dot (disstree2dot), 23
seqtree2display, 22, 26, 136, 137, 138
setwd, 26
stlab, 140
stlab<- (stlab), 140
str.seqelist, 79

title, 24, 42
TraMineR (TraMineR-package), 4
TraMineR-package, 4
TraMineR.check.depr.args, 141
TraMineR.checkupdates, 142
TraMineRInternal, 142
TraMineRInternalLayout
   (TraMineRInternal), 142
TraMineRInternalLegend
   (TraMineRInternal), 142
TraMineRInternalNodeInit
   (TraMineRInternal), 142
TraMineRInternalSeqage
   (TraMineRInternal), 142
TraMineRInternalSeqbar
   (TraMineRInternal), 142
TraMineRInternalSplitInit
   (TraMineRInternal), 142
TraMineRInternalWeightedInertiaDist
   (TraMineRInternal), 142