The Demonstration of myTraMineR functions

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```
# install.packages("myTraMineR", repos = NULL, type="source")
library(myTraMineR)
library(TraMineR)
data(mvad)
mvad.alphab <- c("employment", "FE", "HE", "joblessness", "school", "training")</pre>
mvad.seq <- seqdef(mvad, 17:86, xtstep = 6, alphabet = mvad.alphab)</pre>
seqfplot(mvad.seq, idxs = 1:20)
     27.1%
   Cum. % freq. (n=712)
        0%
           Sep.93
                                                    Sep.96
                         Sep.94
                                       Sep.95
                                                                  Sep.97
                                                                               Sep.98
```

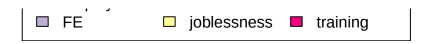


Figure 1: Frequence plot

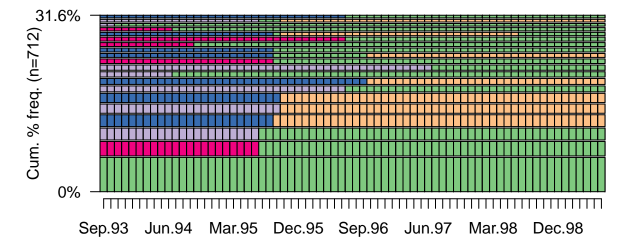
Figure 1 shows that the first 20 most frequent sequences represent 27.1% of the total. Now, let's consider a situation: if a sequence differs by only one state from the most common sequence, which consists solely of the "employment" state, this sequence won't be shown in the plot. The difference may be caused by repeating error—noise, so we are interested in plotting this sequence in the frequency plot as well. In other words, we would like to replace sequences with their representatives and then draw a frequency plot.

Note that it will be important to examine whether small differences among sequences are true or not in the exploratory data analysis process. If users observe a significant improvement in frequency when considering small differences as not true, then in the later analysis, users should consider using the new dataset obtained by replacing these sequences with their representatives.

Function 1: The Extension of seqfplot()

Figure 2 shows that if we consider two sequences different by one Hamming distance as the same, the first 20 most frequent sequences represent 31.6% of the total. This number increased by 16.6% compared to Figure 1.

```
seqdistance <- seqdist(mvad.seq, method="HAM")
clusters <- seqcluster(seqdistance, h=1.5, cmethod='complete')
alphabet <- c("employment", "FE", "HE", "joblessness", "school", "training")
mvad.new.seq <- seqrep_replace(mvad, clusters, var=17:86, alphabet, idxs=1:20, xtsetp=6)
seqfplot(mvad.new.seq, idxs=1:20)</pre>
```



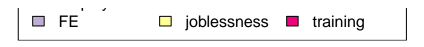


Figure 2: Frequency plot for mvad.new.seq dataset.

We cut the hierarchical tree to enforce the maximal distance within clusters to separate the 20 most frequent sequences. Next, we identified the representative—specifically, the mode—for each cluster and used these representatives to replace every element in the clusters. This new dataset, named mvad.new.seq, is also a stslist object. Finally, we generated a frequency plot for mvad.new.seq.

- 1. seqdist() is used to calculate distance matrix using Hamming distance. This is already in TraMineR.
- 2. seqcluster() uses complete linkage clustering and returns a data frame containing cluster information. There are three columns in the returned data frame—index, cluster, and Freq. It shows which cluster a data point belongs to and the cluster size.
- 3. seqrep_replace() is similar as seqdef(). It returns a stslist object but with elements replaced by their cluster representatives.

If significant improvement on the frequency is observed, then the new stslist object, mvad.new.seq, should be considered in the later analysis.

Use help() to check the usage of two functions—seqcluster() and seqrep_replace() that we created in myTraMineR package.

seqcluster {myTraMineR}

R Documentation

Cluster sequences

Description

This function clusters sequences using hierarchical cluster and returns group memberships of sequences.

Usage

```
seqcluster(seqdistance, h = 1.5, cmethod = "complete")
```

Arguments

segdistance A distance matrix or a distance array returned by TraMineR::segdist()

function.

h Numeric scalar or vector with heights where the tree should be cut. The

default is 1.5.

cmethod The agglomeration method to be used. This should be (an unambiguous

abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC). The default clustering method is "complete", complete-linkage

clustering.

Value

seqcluster returns a data frame with three columns—the index of original data, group memberships, and group size.

Examples

```
data(mvad)
mvad.alphab <- c("employment", "FE", "HE", "joblessness", "school", "tra
mvad.seq <- seqdef(mvad, 17:86, xtstep = 6, alphabet = mvad.alphab)
seqdistance <- seqdist(mvad.seq, method="HAM")
clusters <- seqcluster(seqdistance, h=1.5, cmethod='complete')</pre>
```

[Package myTraMineR version 0.0.0.9000 Index]

Replace sequences with their representatives

Description

Similar as seqdef(). It returns a stslist object but with elements replaced by their cluster representatives.

Usage

segrep replace(data, clusters, var = NULL, alphabet, idxs = 1:10, ...)

Arguments

data A data frame, matrix, or character string vector containing sequence data (tibble

will be converted with as.data.frame).

clusters A data frame, containing three columns—the index of original data, group

memberships, and group size. Or the cluster data frame returned by

seqcluster function.

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.

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.

idxs A integer or an array of integers. The Default is 1:10, meaning replacing

sequences in the 10 largest clusters with their representatives. If idxs=0, then all

clusters will be replaces.

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

format.

Value

An object of class stslist.

Function2: Get the cut height

getheight() returns the cut height for cutting a hierarchical clustering tree. The returned cut height h will satisfy that the first a few most frequent groups, specified by idxs, will represent at least fq of the total.

```
## $h
## [1] 5.5
##
## $frequency
## [1] 0.3300562
getheight(seqdistance, fq=0.3, idxs=1:100)
## There is no need to cluster.
## $h
## [1] 0
##
## $frequency
## [1] 0.4522472
```

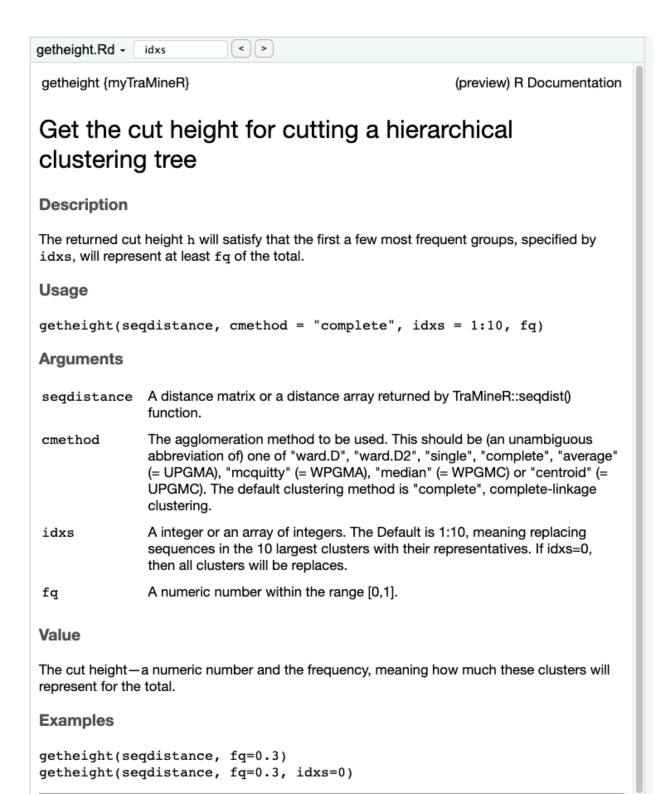


Figure 3: getheight