Smoothing discrete data (I) - using the **smooth.discrete()** function in the **mhsmm** package

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1 Introduction

The smooth.discrete() function provides a simple smoothing of a time series of discrete values measured at equidistant times. Under the hood of smooth.discrete() is a hidden Markov model.

More details – and an additional example – is provided in the vignette "Smoothing discrete data (II)"

2 Using smooth.discrete()

For example consider the data:

Calling smooth.discrete() on these data gives

```
> obj <- smooth.discrete(y1)
A 'smoothDiscrete' object
List of 4
$ s   : num [1:45] 1 1 1 1 1 1 1 1 1 1 1 ...
$ model :List of 9
   ..- attr(*, "class")= chr "hmm"
$ data :List of 3
$ initial:List of 3
- attr(*, "class")= chr "smoothDiscrete"
NULL
NULL</pre>
```

The \mathbf{s} slot of the object contains the smoothed values. We illustrate the results in Figure 1.

```
> plot(y1, ylim=c(0.8,2))
> addStates(obj$s)
```

The smoothed sequence of states is by default the jointly most likely sequence of states as obtained by the Viterbi algorithm.

A smooth of a new time series is produced as



Figure 1: Observed and smoothed discrete time time series.

```
> y2 <- c(1,1,1,1,2,2,2,1,1,2,1,1,1,2,1,1,1,1,1,2,2,2,NA,1,1,1,1,2,2,1,2,2,2)
> predict(obj,x=y2)
$s
$x
               2 2 1 1 2 1 1 1 2 1 1 1 1 1 2 2 2 NA 1 1
2 2
           1 2
1 2
[1]
    1
      1
        1
[26]
      2
        2
    1
$N
[1] 32
$loglik
[1] -24.53755
attr(,"class")
[1] "hsmm.data"
```

Here the smoothed values are in the \mathbf{s} slot. Again, the sequence is by default the jointly most likely sequence of states.

The estimated parameters are:

```
> summary(obj)
init:
    1 0
transition:
    [,1] [,2]
[1,] 0.920 0.080
[2,] 0.136 0.864
emission:
$pmf
    [,1] [,2]
[1,] 0.7925235 0.2147958
[2,] 0.3418576 0.6443388
```

3 The arguments to smooth.discrete()

The arguments of smooth.discrete() are

```
> args(smooth.discrete)
```

```
function (y, init = NULL, trans = NULL, parms.emission = 0.5,
    method = "viterbi", details = 0, ...)
NULL
```

• init is a vector of initial probabilities for the Markov chain. If init=NULL then the initial distribution is taken to be the relative frequencies in data, that is

• trans is the transition matrix for the Markov chain. If trans=NULL then the transition matrix is derived from data as:

If **trans** is a vector (of numbers smaller than 1) then these are taken to be the diagonal of the transition matrix and the off-diagonal elements are then, within each row, taken to be identical so that the rows sum to 1. Elements of **trans** are recycled so as to make the dimensions match. Under the hood, the matrix is created as, for example:

```
> createTransition(c(0.8,0.9),2)
```

```
[,1] [,2]
[1,] 0.8 0.2
[2,] 0.1 0.9
```

- parms.emission is a matrix describing the conditional probabilities of the observed states given the latent states. If parms.emission is a vector then the matrix is created following the same scheme as for the transition matrix described above.
- The method argument is either "viterbi" (which produces the jointly most likely sequence of states). The alternative method is smoothed which produces the individually most likely states.
- The dotted arguments are passed on the the hmmfit function. For example, one may specify lock.transition=TRUE in which case the transition matrix is not estimated from data.