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:

```r
> y1 <- c(1,1,1,1,2,1,1,NA,1,1,2,1,1,1,1,1,2,2,2,1,2,2,2,1,1,1,1,1,1,2,2,2,1,1)
```

Calling `smooth.discrete()` on these data gives

```r
> obj <- smooth.discrete(y1)
```

A `smoothDiscrete` object

List of 4

$s : num [1:45] 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

The $s slot of the object contains the smoothed values. We illustrate the results in Figure 1.

```r
> 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(1,1,1,2,2,1,1,2,1,1,1,2,2,2,NA,1,1,2,2,1,2,2,2)
> predict(obj,x=y2)

$s
[1] 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

$x
[1] 1 1 1 1 2 2 2 1 1 2 1 1 1 2 1 1 1 1 1 2 2 2 2 NA 1 1
[26] 1 2 2 1 2 2 2

$N
[1] 32

$loglik

attr(,"class")
[1] "hsmm.data"

Here the smoothed values are in the $s$ 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

  ```
  > table(y1)/sum(table(y1))
  
  1 2
  0.6363636 0.3636364
  ```

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

  ```
  > ttt<-table(y1[-length(y1)],y1[-1])
  > ttt
  
  1 2
  1 19 7
  2 7 9
  
  > sweep(ttt, 1, rowSums(ttt), "/")
  
  1 2
  1 0.7307692 0.2692308
  2 0.4375000 0.5625000
  ```

  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.