hmmTMB user guide

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2020-11-16

1 Introduction

The package hmmTMB implements hidden Markov models (HMMs) with flexible covariate dependence in all model parameters. In this document, we briefly describe the structure of the package, and illustrate its use with several examples. For general background on HMMs, see Zucchini, MacDonald, and Langrock (2016) and, for a presentation of HMMs in the context of ecological studies, see McClintock et al. (2020).

HMMs are time series models involving two processes: an unobserved state process (S_t) specified as a Markov chain, and an observation process (Z_t) . At each time t = 1, 2, ..., the distribution of the observation Z_t depends on the value S_t of the state process. There are therefore two sets of parameters:

- the state process is formulated in terms of a transition probability matrix (and an initial distribution);
- the observation process is formulated in terms of the state-dependent parameters of the observation distributions.

In hmmTMB, the parameters of the state process and of the observation process can depend on covariates, including linear fixed effects, smooth covariate effects using splines, and i.i.d. normal random effects.

2 Package structure

The package hmmTMB uses the R6 framework for object-oriented programming (Chang (2019)), and it is based on three main classes:

- MarkovChain: model for state process, including formulas for the transition probabilities
- Observation: model for observation process, including state-dependent distributions, and formulas for the state-dependent observation parameters
- HMM: contains a state process model (i.e., a MarkovChain object) and an observation model (i.e., an Observation object).

Model specification, model fitting and model visualisation can all be performed by manipulating objects from those three classes. Following the R6 syntax, an object is created with something like

```
hidden <- MarkovChain$new(n_states = 2)
```

This line of code defines a hidden state model with 2 states and no covariate dependence. Once an object is created, various methods (i.e., functions) can be called to manipulate it, using the following syntax:

```
hidden $formulas()
```

```
$`S1>S2`
~1
<environment: 0x559e75651b00>
$`S2>S1`
```

<environment: 0x559e75654c68>

3 Example 1: Elk movement analysis

We illustrate the use of the package on a data set of elk movement (from Morales et al. (2004)), accessible through the moveHMM package. In this example, we showcase the use of smoothing splines to model the relationship between model parameters and covariates.

```
# Load packages
library(hmmTMB)
library(moveHMM)
```

In analyses of animal movement, the step lengths and turning angles are modelled to capture features of the speed and directionality of the movement. We derive those variables from the location using the function prepData in moveHMM.

```
tracks <- prepData(elk_data, type = "UTM", coordNames = c("Easting", "Northing"))
head(tracks)</pre>
```

```
ID
                step
                           angle
                                      х
                                               y dist_water
1 elk-115
           5518.4434
                              NA 769928 4992847
                                                     200.00
2 elk-115
           1416.5663
                      0.1262112 766875 4997444
                                                     600.52
3 elk-115
                      2.3832412 765949 4998516
                                                     561.81
            239.7525
4 elk-115
            432.7600
                      0.9385238 765938 4998276
                                                     550.00
5 elk-115
            103.7545
                      1.1375066 766275 4998005
                                                     302.08
6 elk-115 12416.4659 -0.9687435 766368 4998051
                                                     213.60
```

The data set now has columns for the coordinates (x and y), the step lengths and turning angles, the ID (track identifier), and a covariate (distance to water). hmmTMB doesn't allow for zero inflation for observation distributions with positive support (e.g. gamma), so we jitter step lengths with value zero.

```
indz <- which(tracks$step == 0)
tracks$step[indz] <- runif(length(indz), 0, 1)</pre>
```

The data frame passed to hmmTMB should include a column for ID if necessary, as well as for any response variables and covariates included in the model.

The class Observation encapsulates the observation process model of the hidden Markov model. To create an object of that class, we need to specify distributions for the response variables, as well as initial parameter values. We could also include covariate dependence in the parameters of the observation distributions. We use gamma distributions for the step lengths, and von Mises distributions for the turning angles.

Similarly, we need to create an object of the class MarkovChain for the hidden process model, encapsulating the formulas and parameter values. Here, we formulate the transition probabilities as functions of the distance to water covariate, with smoothing splines specified using the syntax from the package mgcv (Wood (2017)).

Combining the MarkovChain and Observation object, we create an object of the HMM class.

```
hmm <- HMM$new(obs = obs, hidden = hid)</pre>
```

We fit the model with the method fit, and print the output of the optimiser with res. It contains the parameters estimates (fixed effects), the value of the objective function at the optimum, and diagnostics from the optimiser.

```
hmm$fit()
############################
## Observation model ##
#########################
+ step ~ gamma(shape, scale)
  * shape.state1 ~ 1
  * shape.state2 ~ 1
  * scale.state1 ~ 1
  * scale.state2 ~ 1
+ angle ~ vm(mu, kappa)
  * mu.state1 ~ 1
  * mu.state2 ~ 1
  * kappa.state1 ~ 1
  * kappa.state2 ~ 1
#############################
## State process model ##
############################
                                   state 1
                                           ~s(dist_water, k = 5, bs = "ts")
state 2 ~s(dist_water, k = 5, bs = "ts")
hmm$out()
$par
  coeff_fe_obs
                  coeff_fe_obs
                                 coeff_fe_obs
                                                 coeff_fe_obs
                                                                 coeff_fe_obs
```

8.66432402

coeff_fe_hid

-2.09231345

6.08410554

coeff_fe_obs

-1.32712147

10.30269244

coeff_fe_hid

-0.38760827

-0.16583071

-0.03581825

coeff_fe_obs

-0.49138014

coeff_fe_obs

-0.55219155

```
log_lambda_hid log_lambda_hid log_delta
3.63477846 -0.14735819 -0.72805839

$value
[1] 6928.693

$counts
function gradient
55 28
```

\$convergence

[1] 0

\$message

NULL

We can get confidence intervals for the estimated (working) parameters with the method CI_coeff By default, it returns 95% confidence intervals, but the argument level can be specified for different confidence levels.

```
hmm$CI_coeff()
```

```
lower
                     estimate
                                                  upper
coeff_fe_obs1
                               -0.29574924 -0.03591218
                -0.1658307069
coeff_fe_obs2
                -0.4913801413
                               -0.82920316 -0.15355713
coeff_fe_obs3
                 6.0841055406
                                5.84834728
                                            6.31986380
coeff_fe_obs4
                 8.6643240218
                                8.37832785
                                           8.95032019
coeff_fe_obs5
                10.3026924394 -33.67595325 54.28133813
coeff_fe_obs6
                -0.0358182545
                               -0.58229227
                                            0.51065576
coeff fe obs7
                -0.5521915451
                               -0.81079847 -0.29358462
coeff_fe_obs8
                -1.3271214746
                               -2.39292525 -0.26131770
coeff_fe_hid1
                -2.0923134514
                               -2.65244837 -1.53217853
coeff_fe_hid2
                -0.3876082707
                               -1.51837376 0.74315722
log_lambda_hid1
                               -2.24415055
                                           9.51370746
                3.6347784575
log lambda hid2 -0.1473581888
                               -3.25566433
                                            2.96094795
log_delta
                               -3.60570677
                -0.7280583914
                                            2.14958998
coeff_re_hid1
                               -0.06345239
                -0.0031987515
                                            0.05705488
coeff_re_hid2
                               -0.08056088
                -0.0005351417
                                            0.07949059
coeff_re_hid3
                -0.0024258152
                               -0.17970435
                                            0.17485272
coeff_re_hid4
                -0.1824971891
                               -0.79497196
                                            0.42997758
coeff_re_hid5
                -0.0026214640
                               -0.38470540
                                            0.37946247
coeff_re_hid6
                 0.0142890913
                               -0.51526323
                                            0.54384141
coeff_re_hid7
                 0.0522455547
                               -1.11801076
                                            1.22250187
coeff_re_hid8
                 1.5866552767
                                0.18576142
                                            2.98754913
```

The methods predict_obspar and predict_tpm can also be used to obtain estimates and confidence intervals of the model parameters directly. For example, say that we want to predict the model parameters for dist_water = 1000,

```
new_data <- data.frame(dist_water = 1000)
hmm$predict_obspar(new_data = new_data, CI = TRUE)</pre>
```

\$estimate

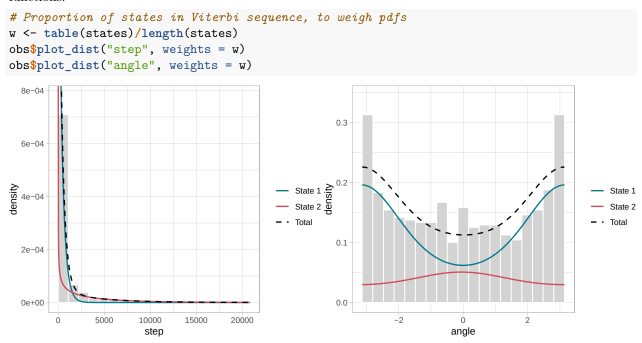
, , 1

```
state 1 state 2 step.shape 0.8471896 0.61178147
```

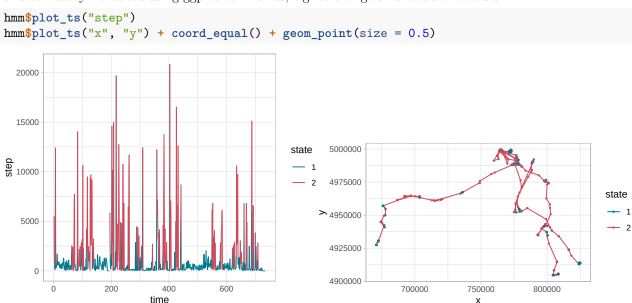
```
step.scale 438.8271240 5792.52760426
angle.mu
              3.1413819
                           -0.05625717
angle.kappa
              0.5756868
                            0.26523966
$low
, , 1
                state 1
                               state 2
step.shape
              0.7507598
                            0.43713635
step.scale 345.1287339 4368.26456543
angle.mu
                           -0.87875092
             -3.1415927
angle.kappa
              0.4522470
                            0.09462405
$upp
, , 1
                state 1
                              state 2
step.shape
                            0.8461125
              0.9560138
step.scale 555.3290994 7687.1515889
angle.mu
              3.1415927
                            0.8256917
                            0.7736108
angle.kappa
              0.7393386
hmm$predict_tpm(new_data = new_data, CI = TRUE)
$estimate
, , 1
                     [,2]
          [,1]
[1,] 0.8950868 0.1049132
[2,] 0.5187398 0.4812602
$low
, , 1
          [,1]
                      [,2]
[1,] 0.8374103 0.06600354
[2,] 0.2181967 0.22503597
$upp
, , 1
          [,1]
                     [,2]
[1,] 0.9339965 0.1625897
[2,] 0.7749640 0.7818033
The method viter implements the Viter ialgorithm, to estimate the most likely state sequence.
states <- hmm$viterbi()</pre>
head(states)
```

[1] 2 2 1 1 1 2

Various plotting functions are provided to visualise a fitted model. Using the method Observation\$plot_dist, we can plot histograms of the observed data, overlaid with the estimated state-dependent probability density functions.

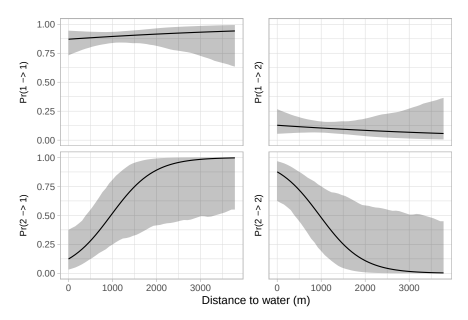


We can plot the data coloured by the Viterbi-decoded states. All plotting functions return ggplot objects, and can easily be edited using ggplot commands, e.g. to change axis labels or colours.

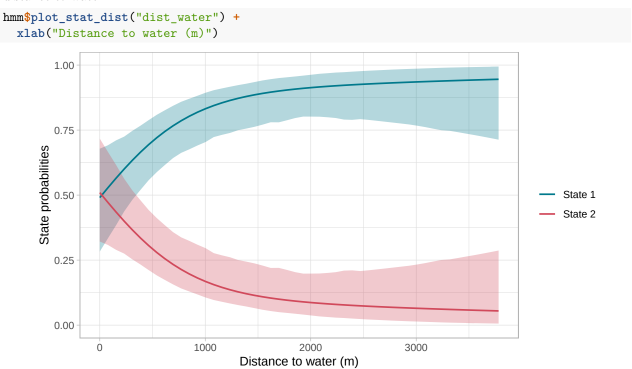


The function plot_tpm creates graphs of the transition probabilities as functions of a covariate, here distance to water.

```
hmm$plot_tpm("dist_water") + xlab("Distance to water (m)")
```



With the function plot_stat_dist, we can visualise the stationary state probabilities as functions of the distance to water.



We can simulate from the fitted model with the method simulate, for example to check how well the model captures features of the real data. Here, we simulate step lengths and turning angles from the model.

```
# Simulate from fitted model
sim_data <- hmm$simulate(n = n_sim, data = new_covs)</pre>
```

The output is a data frame with the same columns as the input, plus columns for the simulated variables, and a column for the simulated states.

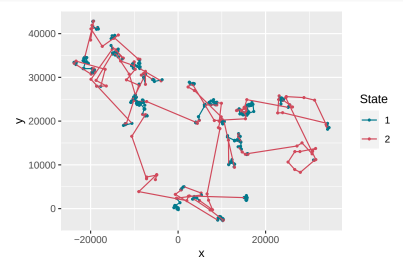
head(sim_data)

```
dist_water ID state
                            step
                                     angle
1
      527.38
              1
                        23.58317 4.881601
2
      206.16
             1
                     1 237.89069 4.895131
3
     1948.24
                     1 296.59420 2.321896
4
                     1 405.08077 2.102369
     3414.13
              1
5
      213.60
              1
                     1 221.83249 1.290461
6
      300.00
                     1 112.87189 3.242383
```

We then derive the corresponding locations to visualise the simulated movement trajectory.

```
# Get locations from simulated step lengths and turning angles
bearings <- cumsum(sim_data$angle)
sim_dxy <- sim_data$step * cbind(cos(bearings), sin(bearings))
sim_data$x <- cumsum(c(0, sim_dxy[-n_sim, 1]))
sim_data$y <- cumsum(c(0, sim_dxy[-n_sim, 2]))

# Plot simulated trajectory
ggplot(sim_data, aes(x, y, col = factor(state), group = NA)) +
    scale_color_manual("State", values = hmmTMB_cols) +
    geom_point(size = 0.7) +
    geom_path() +
    coord_equal()</pre>
```



4 Example 2: wild haggis movement

We further illustrate the functionalities of hmmTMB with the analysis of the wild haggis tracking data set from Michelot, Langrock, and Patterson (2016). The objective of the study is to investigate the effect of terrain slope on the behaviour of wild haggises, accounting for inter-individual differences.

We load the data from Michelot, Langrock, and Patterson (2016), which contains locations for 15 haggis tracks, as well as slope and temperature measurements,

```
ID x y slope temp
1 1 0.000000 0.000000 25.957002 10.344959
2 1 -1.068761 -0.194650 18.606632 8.352531
3 1 -6.152549 2.051343 16.524004 13.529650
4 1 -6.703983 3.338480 9.154917 10.951095
5 1 -6.541667 3.553843 5.547686 11.243328
6 1 -7.160298 1.960377 8.129402 13.187280
```

We use the prepData function from moveHMM to calculate step lengths, and we transform ID into a factor variable,

```
data <- prepData(raw, type = "UTM")
data$ID <- factor(data$ID)
head(data)</pre>
```

```
ID
         step
                   angle
                                                 slope
                                                            temp
 1 1.0863417
                      NA 0.000000 0.000000 25.957002 10.344959
 1 5.5578218 -0.5961622 -1.068761 -0.194650 18.606632 8.352531
 1 1.4002860 -0.7500230 -6.152549
                                    2.051343 16.524004 13.529650
4 1 0.2696813 -1.0506197 -6.703983
                                    3.338480
                                             9.154917 10.951095
5 1 1.7093394 -2.8660552 -6.541667
                                    3.553843 5.547686 11.243328
6 1 1.1529149 2.3676683 -7.160298 1.960377 8.129402 13.187280
```

We define the hidden state process as a 2-state Markov chain, with a smooth effect of "slope" (using thin-plate regression splines), and a normal random intercept for "ID" to allow for differences between haggises. We don't include the effect of temperature because Michelot, Langrock, and Patterson (2016) found no clear effect.

To define the observation model, we specify that the step lengths should be modelled with gamma distributions. We choose initial parameter values for the mean and standard deviation based on data visualisations, and derive initial values for the shape and scale based on that,

```
par = par0)
```

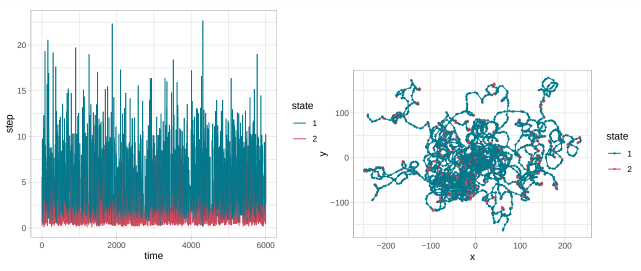
We create the HMM object from the MarkovChain and Observation components, fit it using the fit method, and compute the most likely state sequence using the viter is method,

```
hmm <- HMM$new(hidden = hid, obs = obs)
hmm$fit()</pre>
```

```
data$state <- hmm$viterbi()</pre>
```

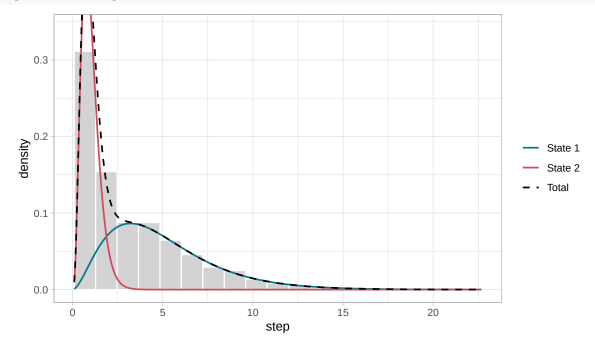
We can plot the time series of step lengths, or the tracks, coloured by the estimated state sequence,

```
hmm$plot_ts("step")
hmm$plot_ts("x", "y") +
    geom_point(size = 0.5) +
    coord_equal()
```



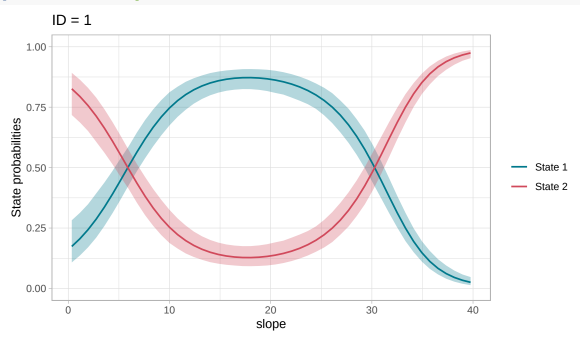
These plots suggest that state 1 captured fast movement behaviour (long step lengths) and state 2 captured slow movement (short step lengths). This can also be seen from the estimated state-dependent distributions of step lengths,





To visualise the effect of slope on the behaviour of wild haggises, we plot the stationary state probabilities as functions of the slope,

hmm\$plot_stat_dist("slope")



When there are several covariates, the plotting functions fix the other covariates to their mean value (for continuous covariates) or to their first level (for factor covariates), as shown at the top of the plot. This can be changed by specifying the argument "covs", a data frame with a single row and one named column for each covariate.

We observe the same pattern found by Michelot, Langrock, and Patterson (2016): wild haggises tended to adopt the fast movement state at intermediate slopes (between 5 and 30 degrees), and the slow movement

state on very flat or very steep terrains. Michelot, Langrock, and Patterson (2016) compared a linear and a quadratic effect of slope; the smoothing splines used here are more flexible, and the implementation automatically estimates the smoothness of the relationship during model fitting.

A measure of inter-individual heterogeneity is given by the standard deviation of the distribution of random intercepts, and can be obtained using the vcomp method,

hid\$vcomp()

```
[,1]
S1>S2.s(ID) 0.12888757
S1>S2.s(slope) 4.47960634
S2>S1.s(ID) 0.01866037
S2>S1.s(slope) 3.74073593
```

References

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Michelot, Théo, Roland Langrock, and Toby A Patterson. 2016. "moveHMM: An R Package for the Statistical Modelling of Animal Movement Data Using Hidden Markov Models." *Methods in Ecology and Evolution* 7 (11): 1308–15.

Morales, Juan Manuel, Daniel T Haydon, Jacqui Frair, Kent E Holsinger, and John M Fryxell. 2004. "Extracting More Out of Relocation Data: Building Movement Models as Mixtures of Random Walks." *Ecology* 85 (9): 2436–45.

Wood, S. N. 2017. Generalized Additive Models: An Introduction with R. 2nd ed. Chapman; Hall/CRC.

Zucchini, Walter, Iain L MacDonald, and Roland Langrock. 2016. Hidden Markov Models for Time Series: An Introduction Using R, Second Edition. CRC press.