Supplementary file 2: Autocorrelation-informed home range estimation with the ctmm R package

Inês Silva* Christen H. Fleming Michael J. Noonan Jesse Alston Cody Folta William F. Fagan Justin Calabrese[†]

Introduction

Home range estimation is a key output from tracking datasets, but the inherent properties of animal movement can lead traditional methods to under- or overestimated their size. Autocorrelated Kernel Density Estimation (AKDE) methods were designed to be statistically efficient while explicitly dealing with the complexities and biases of modern movement data, such as autocorrelation, small sample sizes, and missing or irregularly sampled data.

This tutorial is a companion piece to our manuscript "Autocorrelation-informed home range estimation: a review and practical guide". For any definitions, check the main manuscript or the Glossary. The AKDE family of home range estimators will be run using R software (https://www.r-project.org/) and the ctmm package (Calabrese, Fleming, and Gurarie 2016). The techniques and mitigation measures available within this package include:

Method	When to run?	What does it do?	R function
AKDE	Tracking data is autocorrelated	Estimates range distributions from autocorrelated data, by conditioning on an autocorrelation model.	akde(, debias = FALSE)
AKDEc	If using GRF-based KDEs (such as AKDE)	Removes the tendency of Gaussian reference function (GRF) methods to overestimate the area of home ranges.	akde(, debias = TRUE)
pHREML	Small (absolute and effective) sample sizes	Improves upon ML and REML autocorrelation estimation, mitigating small sample size biases.	<pre>ctmm.select(, method = "pHREML")</pre>
wAKDEc	Irregular sampling schedules or missing data	Upweights observations that occur during under-sampled times, while downweighting those occurring during over-sampled times.	<pre>akde(, weights = TRUE)</pre>
Parametric bootstrap	Extremely small effective sample size	Calculates and corrects for autocorrelation estimation biases, by simulating from an approximate sampling distribution.	ctmm.boot()

^{*}i.simoes-silva@hzdr.de

 $^{^{\}dagger}$ j.calabrese@hzdr.de

AKDE_c and pHREML are default arguments within the akde() and ctmm.select() functions, respectively: both will run automatically if arguments debias and method are left unspecified. For most situations, we recommend keeping both of these arguments as the default.

```
# Installing & loading package:
install.packages("ctmm")
library(ctmm)
```

We provide a guide to **home range estimation** using the following workflow:

- Step 1. Formatting and loading an animal tracking dataset;
- Step 2. Checking for the range residency assumption;
- Step 3. Selecting the best-fit movement model through model selection;
- **Step 4.** Feeding a movement model into the *home range estimator*;
- Step 5. Evaluating additional biases, applying mitigation measures.

Data Preparation

We will use two datasets, both available within the ctmm package: African buffalos (Syncerus caffer), and Mongolian gazelles (Procapra gutturosa). Information on the data collection protocol is available in Cross et al. (2009) and Fleming et al. (2014). The ctmm package requires data to conform to Movebank naming conventions (https://www.movebank.org/node/2381). We recommend uploading your data to Movebank (http://www.movebank.org/) as this will facilitate data preparation, and ensure that your data are correctly formatted for ctmm. If needed, Movebank allows you to keep your data private.

We will focus on the simplest data structure:

- animal ID An individual identifier for each animal tracked;
- timestamp or t The date and time corresponding to a sensor measurement;
 - **Example**: 2021-01-01 18:31:00.000
 - Format: yyyy-MM-dd HH:mm:ss.SSS
- longitude or x The geographic longitude of the location as estimated by the sensor.
 - Example: -121.1761111
 - Units: decimal degrees, WGS84 reference system.
- latitude or y The geographic latitude of the location as estimated by the sensor;
 - **Example**: -41.0982423
 - Units: decimal degrees, WGS84 reference system.

Location can also be described as UTM locations instead of latitude/longitude. In this case, you should provide UTM easting, UTM northing, and UTM zone. For all terms and conventions, please see the full

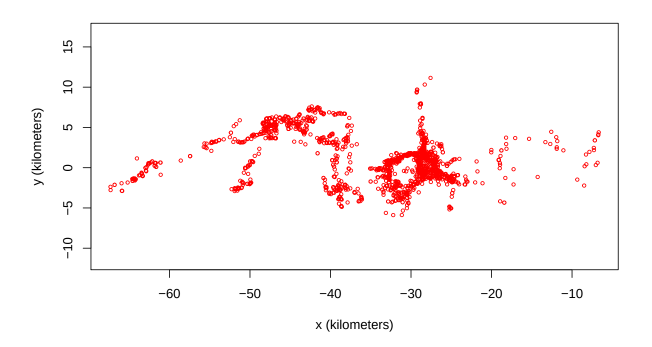
vocabulary list here: http://vocab.nerc.ac.uk/collection/MVB/current/.

Step 1. - Formatting and loading an animal tracking dataset

You can import data into R through the read.table() or read.csv() functions; make sure to navigate to the appropriate folder or working directory. For this tutorial, our data is already prepared into a list of telemetry objects which we can load into R:

1.1. Buffalo tracking data

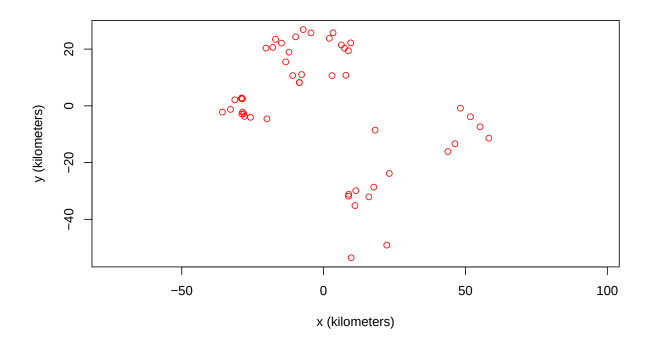
```
data("buffalo")
animal1_buffalo <- buffalo[[4]] # select individual number 4</pre>
head(animal1_buffalo)
##
                   timestamp longitude latitude
## 17517 2006-04-25 05:09:00 31.73749 -24.19705 1145941740 -51803.35 -2715.663
## 17518 2006-04-25 06:09:00
                              31.73653 -24.19929 1145945340 -51569.29 -2845.660
## 17519 2006-04-25 07:09:00
                              31.73946 -24.20100 1145948940 -51340.72 -2576.353
## 17520 2006-04-25 08:09:00
                              31.73987 -24.20092 1145952540 -51344.11 -2533.788
                              31.74086 -24.20365 1145959740 -51029.45 -2474.771
## 17521 2006-04-25 10:09:00
## 17522 2006-04-25 11:09:00
                              31.74098 -24.20370 1145963340 -51022.23 -2463.655
# Plotting locations:
plot(animal1_buffalo)
```



This dataset showcases an irregular sampling schedule: the buffalo nicknamed "Pepper" had a sampling rate shift from one fix every hour to one fix every two hours. We will use this dataset to highlight data irregularity and the **wAKDE** mitigation measure.

1.2. Gazelle tracking data

```
data("gazelle")
animal2_gazelle <- gazelle[[11]] # selecting individual number 11</pre>
head(animal2_gazelle)
##
                X
                                    t
         18152.70
## 2742
                   -8539.799
                                    0
         15931.16 -32069.788 1306800
  2743
## 2744
         17678.84 -28632.329 1396800
         23135.50 -23820.789 1486800
  2745
  2746 -20310.78
                   20348.792 2419200
## 2747 -17920.31
                   20598.668 2509200
# Plotting locations:
plot(animal2_gazelle)
```



Mongolian gazelles have a **home range crossing time** of a few months, and with a maximum longevity around 10 years, it is impossible to get a considerable **effective sample size** no matter the study duration (Christen H. Fleming et al. 2019). We will use this dataset to highlight how to check **effective sample size**

Data Analysis

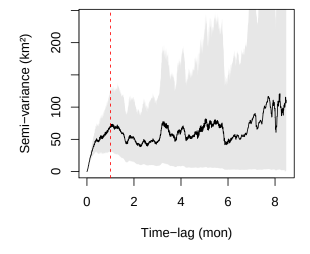
Step 2. – Checking for the range residency assumption

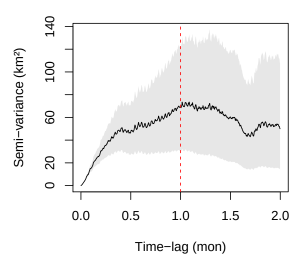
First, we want to check if our first tracking dataset (animal1_buffalo) can be used for home range estimation by checking for range residency. To achieve this, we calculate the semi-variance function (SVF), and visualize it through the variogram() function.

Variograms are an unbiased way to visualize *autocorrelation* structure, representing the average square displacement (y-axis) over a specific time lag (x-axis). To facilitate interpretation, we have the SVF of animal1_buffalo zoomed out (right) to showcase all time lags and (left) zoomed in to showcase time lags up to two months:

```
level <- 0.95 # we want to display 95% confidence intervals
xlim <- c(0,2 %#% "month") # to create a window of 2 months

SVF <- variogram(animal1_buffalo)
par(mfrow = c(1,2))
plot(SVF, fraction = 1, level = level)
abline(v = 1, col = "red", lty = 2) # adding a line at 1 month
plot(SVF, xlim = xlim, level = level)
abline(v = 1, col = "red", lty = 2)</pre>
```





We can see that the variogram flattens (*i.e.*, reaches an asymptote) after approximately **1 month** (red line). This also indicates at how coarse the timeseries needs to be to assume independence (no autocorrelation), and corresponds to when traditional methods—such as **minimum convex polygons** (MCPs) and **Kernel Density Estimators** (KDEs)— could be applied without violating their assumptions.

Step 3. – Selecting the best-fit movement model through model selection

It is necessary to choose a home range estimator that accounts for the autocorrelated structure of the data, now that we see that it is **not** independently and identically distributed (non-IID). We need to test what movement model may explain the autocorrelated structure of our tracking data. We can run different movement processes with **maximum likelihood (ML)** or other parameter estimators, such as **perturbative Hybrid REML (pHREML)**. To facilitate further comparisons, we will run both ML and pHREML with the ctmm.select function.

```
# Calculate an automated model guesstimate:
GUESS1 <- ctmm.guess(animal1_buffalo, interactive = FALSE)</pre>
# Automated model selection, starting from GUESS:
FIT1_ML <- ctmm.select(animal1_buffalo, GUESS1, method = 'ML')</pre>
FIT1_pHREML <- ctmm.select(animal1_buffalo, GUESS1, method = 'pHREML')
## reminder: it will default to pHREML if no method is specified.
summary(FIT1_ML)
## $name
## [1] "OUF anisotropic"
##
## $DOF
##
        mean
                  area
                            speed
##
    11.24296 20.01326 747.72143
##
## $CI
##
                                    low
                                              est
                                                         high
## area (square kilometers) 431.446013 706.20245 1047.55828
## t[position] (days)
                               6.775273 12.18604
                                                    21.91786
## t[velocity] (minutes)
                              31.861346
                                         35.81413
                                                     40.25730
## speed (kilometers/day)
                             15.948352 16.54124
                                                     17.13385
summary(FIT1 pHREML)
## $name
## [1] "OUF anisotropic"
##
## $DOF
##
        mean
                  area
                            speed
##
   10.34454 15.65772 746.09401
##
## $CI
```

```
low
##
                                              est
## area (square kilometers) 439.064822 773.50766 1201.09155
## t[position] (days)
                              6.664052
                                        13.36206
                                                    26.79221
## t[velocity] (minutes)
                             31.807736
                                         35.75689
                                                    40.19636
## speed (kilometers/day)
                             15.949646
                                         16.54326
                                                    17.13658
```

Within these summaries, \$name provides the selected best-fit model, \$DOF provides information on the degrees of freedom (where \$DOF["area"] corresponds to the effective sample size of the home-range area estimate), and \$CI are the parameter outputs (area, position autocorrelation timescale, velocity autocorrelation timescale, and speed).

The typical pool of candidate models includes isotropic (when diffusion is the same in every direction; symmetrical) and anisotropic (when diffusion varies with direction; asymmetrical) variants. The automated model selection shows that *OUF anisotropic* (anisotropic Ornstein-Uhlenbeck foraging process) is our best-fit model. This movement process features a home range, correlated positions, and correlated velocities. To check the full model selection table, we can run the following command:

FIT1_pHREML_verbose <- ctmm.select(animal1_buffalo, GUESS1, verbose = TRUE)
summary(FIT1_pHREML_verbose)</pre>

	$\Delta { m AICc}$	Δ RMSPE (km)	DOF[area]
OUF anisotropic	0.0000	2.224045	15.657669
OU anisotropic	295.0898	2.634793	9.638283
OUF isotropic	330.2236	3.564231	11.204035
OUf anisotropic	1890.8878	0.000000	333.620712

By adding the argument verbose = TRUE we have access to the model selection table. By default, model selection is based on Akaike's Information Criterion adjusted for small sample sizes (AICc). The ctmm package also offers BIC, LOOCV, and HSCV. LOOCV seems to work slightly better for very small datasets, but we recommend AICc for the majority of datasets.

Step 4. – Feeding a movement model into the home range estimator

Now we can fit this movement process into the akde() function, and estimate the home range of animal1_buffalo. This function currently defaults to the area-corrected AKDE, or AKDEc (Fleming & Calabrese 2017):

```
# Run an area-corrected AKDE:
UD1_ML <- akde(animal1_buffalo, FIT1_ML)
UD1_pHREML <- akde(animal1_buffalo, FIT1_pHREML)</pre>
```

```
summary(UD1_pHREML)$CI # home range area estimation

## low est high
## area (square kilometers) 429.6573 756.9343 1175.357
```

We have calculated our home range for animal1_buffalo, resulting in an estimation of 757 km² (with 95% confidence intervals: 430–1,175 km²).

Step 5. – Evaluating additional biases, applying mitigation measures

5.1. Buffalo tracking data

```
summary(UD1_pHREML)$DOF["area"] # effective sample size of animal1

## area
## 15.65772

nrow(animal1_buffalo) # absolute sample size

## [1] 1725
```

Our output here also reveals more information regarding our dataset: the **effective sample size** (N) and the **absolute sample size** (n). We can return this measure with the summary function: in our case, the N for animal1_buffalo is 15.7. Comparatively, our **absolute sample size** is easy to output, as it is the total number of observations within our dataset (n = 1,725).

As mentioned earlier, animal1_buffalo had a device malfunction that led GPS fixes to shift from one fix per hour, to one fix every two hours. As such, this individual is particularly suited for a weighted AKDEc (or wAKDEc), so we can re-run the function with weights set to TRUE:

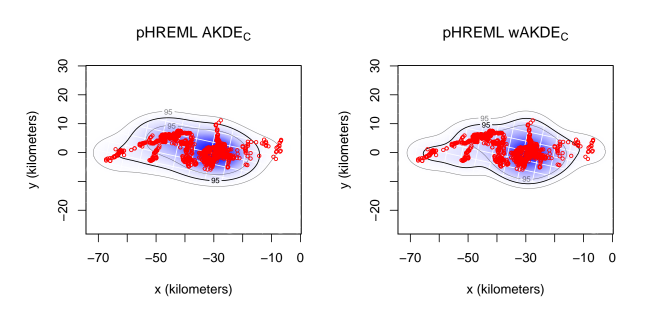
```
UD1w_pHREML <- akde(animal1_buffalo, FIT1_pHREML, weights = TRUE)
summary(UD1w_pHREML)$CI # home range area estimation (weighted)
## low est high
## area (square kilometers) 432.2234 761.4551 1182.376</pre>
```

Our new home range area estimation for animal1_buffalo is 761 km² (with 95% confidence intervals: 432–1,182 km²). We can now plot our home range estimate for animal1_buffalo:

```
# Creating an extent that includes both UDs at the 95% CI level:
EXT <- extent(list(UD1_ML, UD1_pHREML, UD1w_pHREML), level = 0.95)

# Plotting pHREML (with and without weights) side-by-side:
par(mfrow = c(1,2))
plot(animal1_buffalo, UD = UD1_pHREML, ext = EXT)</pre>
```

```
title(expression("pHREML AKDE"["C"]))
plot(animal1_buffalo, UD = UD1w_pHREML, ext = EXT)
title(expression("pHREML wAKDE"["C"]))
```



For animal1_buffalo, the difference between model parameter estimators is not substantial; we only have a ~5.7% AKDE area underestimation by ML compared to pHREML. However, the data fits the spatial locations much better.

```
( 1 - summary(UD1_ML)$CI[1,2] / summary(UD1w_pHREML)$CI[1,2] ) * 100
## [1] 5.742588
```

5.2. Gazelle tracking data

We can also check the difference with animal2_gazelle's tracking data, where the small effective sample size issue is clearer:

```
GUESS2 <- ctmm.guess(animal2_gazelle, interactive = FALSE)

FIT2_ML <- ctmm.select(animal2_gazelle, GUESS2, method = 'ML')

FIT2_pHREML <- ctmm.select(animal2_gazelle, GUESS2, method = 'pHREML')

UD2_ML <- akde(animal2_gazelle, FIT2_ML)

UD2_pHREML <- akde(animal2_gazelle, FIT2_pHREML)
```

With animal2_gazelle, we have a more substantial area underestimation by ML compared to pHREML (\sim 15.2%). We can also see that our effective sample size is only 4.5, with an absolute sample size of 49 (N

```
\ll n).
```

```
(1 - summary(UD2_ML)$CI[1,2] / summary(UD2_pHREML)$CI[1,2] ) * 100
## [1] 15.19245
summary(UD2_pHREML)$DOF["area"] # effective sample size
       area
## 4.528926
nrow(animal2_gazelle) # absolute sample size
```

[1] 49

speed (kilometers/day)

At this point, we have selected a movement process, fed it into a home range area estimation with different model parameter estimators, and corrected for irregular sampling rates. With small effective sample sizes, it is important to see if **parametric bootstrapping** may be worth it to further reduce our estimation error. In order to do so, we can check the expected order of bias from pHREML:

```
# Expected order of pHREML bias:
1/summary(FIT2_pHREML)$DOF['area']^2
##
         area
## 0.04875392
```

The bias is currently $\mathcal{O}(5\%)$ ("in the order of" 5%). As such, we will run parametric bootstrapping for animal2_gazelle. The relative error target is 1% by default (argument error = 0.01), but can be adjusted if necessary.

```
start_time <- Sys.time() # start recording running time</pre>
BOOT <- ctmm.boot(animal2_gazelle, FIT2_pHREML, trace = 2)
## note: this function incurs substantial computational cost, may take hours.
( total_time <- Sys.time() - start_time ) # output running time</pre>
summary(BOOT)
## $name
## [1] "OUF isotropic"
##
## $DOF
##
        mean
                  area
                            speed
   3.196957 3.530735 10.000358
##
##
## $CI
##
                                     low
                                                   est
                                                               high
## area (square kilometers) 3429.704883 14089.021613 32137.455223
## t[position] (months)
                                                           3.758819
                             0.000000
                                             1.781242
                                                          48.870349
## t[velocity] (hours)
                                2.248230
                                            10.481973
```

8.072361

10.551247

5.590052

```
1/summary(B00T)$D0F['area']^3 # expected order of bias

## area
## 0.02271981
```

We can see that the expected order of bias was reduced to 2.3%, which is comparable to the numerical error target of 1%. To reduce the numerical error further, we would need to change the default relative error target of ctmm.boot, but the computational cost would continue to increase, and the comparably large statistical bias (2%) would remain.

Now we will calculate the **AKDEc** based on the estimated parameters, and plot the home range of animal2_gazelle. Because of small effective sample size, we set optimal weights to TRUE for improved statistical efficiency:

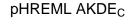
```
UD2_bpHREML <- akde(animal2_gazelle, BOOT, weights = TRUE)
summary(UD2_bpHREML)$CI</pre>
```

```
## low est high
## area (square kilometers) 3231.504 13274.82 30280.25
```

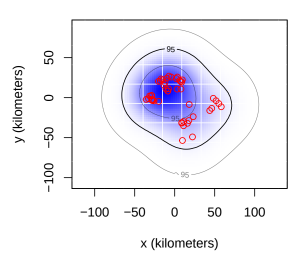
Finally, we have calculated our home range for animal2_gazelle, with an estimated area of 13,274 square kilometers (with 95% confidence intervals: 3,231–30,280 km²). Our uncertainty with animal2_gazelle is substantially higher than with animal1_buffalo, as expected due to the small effective sample size.

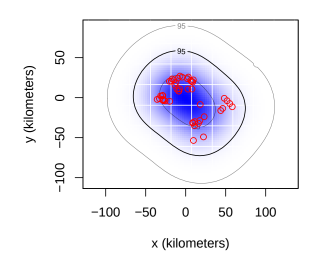
```
# Creating an extent that includes both UDs at the 95% CI level:
EXT <- extent(list(UD2_pHREML, UD2_bpHREML), level = 0.95)

# Plotting pHREML and bootstrapped-pHREML side-by-side:
par(mfrow = c(1,2))
plot(animal2_gazelle, UD = UD2_pHREML, ext = EXT)
title(expression("pHREML AKDE"["C"]))
plot(animal2_gazelle, UD = UD2_bpHREML, ext = EXT)
title(expression("Bootstrapped pHREML wAKDE"["C"]))</pre>
```



Bootstrapped pHREML wAKDE_C





The results presented here were generated with R version 4.0.5, and ctmm version 0.6.0.

Glossary

- *Home range*: the area repeatedly used throughout an animal's lifetime for all its normal behaviors and activities, excluding occasional exploratory moves.
- Range residency: the tendency of an animal to remain within their home range.
- Home range crossing time: the time required for an animal to cross the linear extent of its home range.
- Absolute sample size (n): the observations in a dataset.
- Effective sample size (N): number of range crossings that occurred during the observation period. Can be roughly estimated by dividing the duration of the tracking dataset by the average home range crossing time parameter.

References

Calabrese, Justin M., Chris H. Fleming, and Eliezer Gurarie. 2016. "Ctmm: An r Package for Analyzing Animal Relocation Data as a Continuous-time Stochastic Process." *Methods in Ecology and Evolution* 7

- (9): 1124–32.
- Cross, P. C., D. M. Heisey, J. A. Bowers, C. T. Hay, J. Wolhuter, P. Buss, M. Hofmeyr, A. L. Michel, Roy G. Bengis, and T. L. F. Bird. 2009. "Disease, Predation and Demography: Assessing the Impacts of Bovine Tuberculosis on African Buffalo by Monitoring at Individual and Population Levels." Journal of Applied Ecology 46 (2): 467–75.
- Fleming, Chris H., Justin M. Calabrese, Thomas Mueller, Kirk A. Olson, Peter Leimgruber, and William F. Fagan. 2014. "From Fine-Scale Foraging to Home Ranges: A Semivariance Approach to Identifying Movement Modes Across Spatiotemporal Scales." *The American Naturalist* 183 (5): E154–67.
- Fleming, Christen H., Michael J. Noonan, Emilia Patricia Medici, and Justin M. Calabrese. 2019. "Overcoming the Challenge of Small Effective Sample Sizes in Home-range Estimation." *Methods in Ecology and Evolution* 10 (10): 1679–89.