

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

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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.	<code>akde(..., debias = FALSE)</code>
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.	<code>akde(..., debias = TRUE)</code>
pHREML	Small (absolute and effective) sample sizes	Improves upon ML and REML autocorrelation estimation, mitigating small sample size biases.	<code>ctmm.select(..., method = "pHREML")</code>
wAKDEc	Irregular sampling schedules or missing data	Upweights observations that occur during under-sampled times, while downweighting those occurring during over-sampled times.	<code>akde(..., weights = TRUE)</code>
Parametric bootstrap	Extremely small effective sample size	Calculates and corrects for autocorrelation estimation biases, by simulating from an approximate sampling distribution.	<code>ctmm.boot(...)</code>

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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

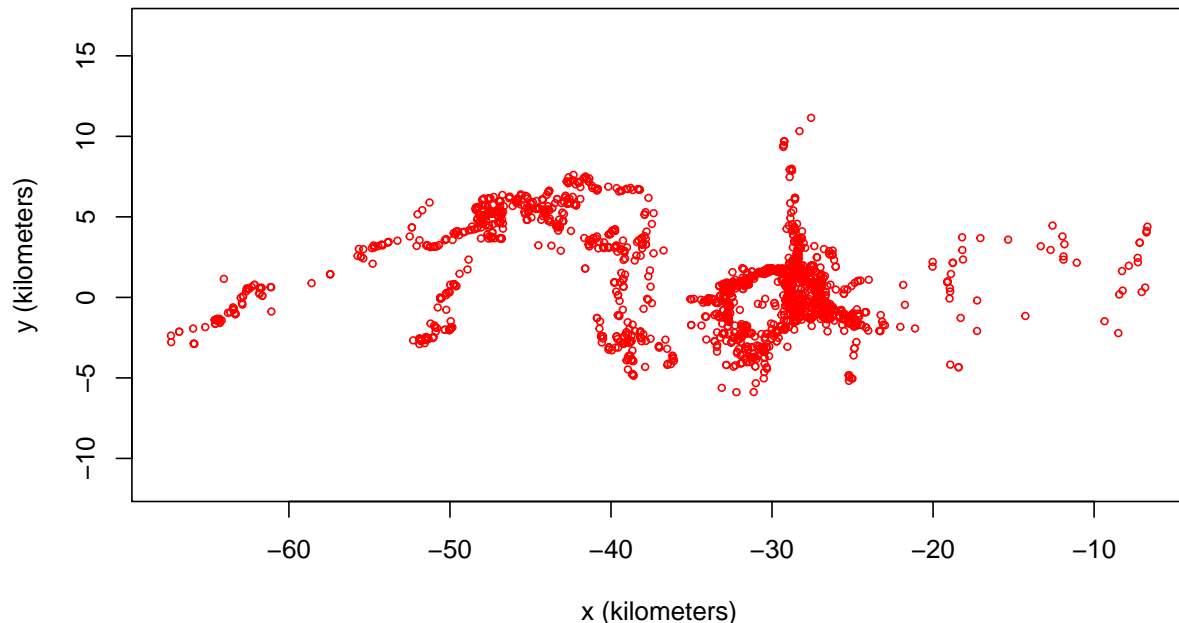
You can import data into R through the `read.table()` or `read.csv()` functions; make sure to navigate to the appropriate folder or working directory. For this tutorial, our data is already prepared into a list of `telemetry` objects which we can load into R:

1.1. Buffalo tracking data

```
data("buffalo")
animal1_buffalo <- buffalo[[4]] # select individual number 4
head(animal1_buffalo)
```

```
##           timestamp longitude latitude      t      x      y
## 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
## 17521 2006-04-25 10:09:00  31.74086 -24.20365 1145959740 -51029.45 -2474.771
## 17522 2006-04-25 11:09:00  31.74098 -24.20370 1145963340 -51022.23 -2463.655
```

```
# Plotting locations:
plot(animal1_buffalo)
```



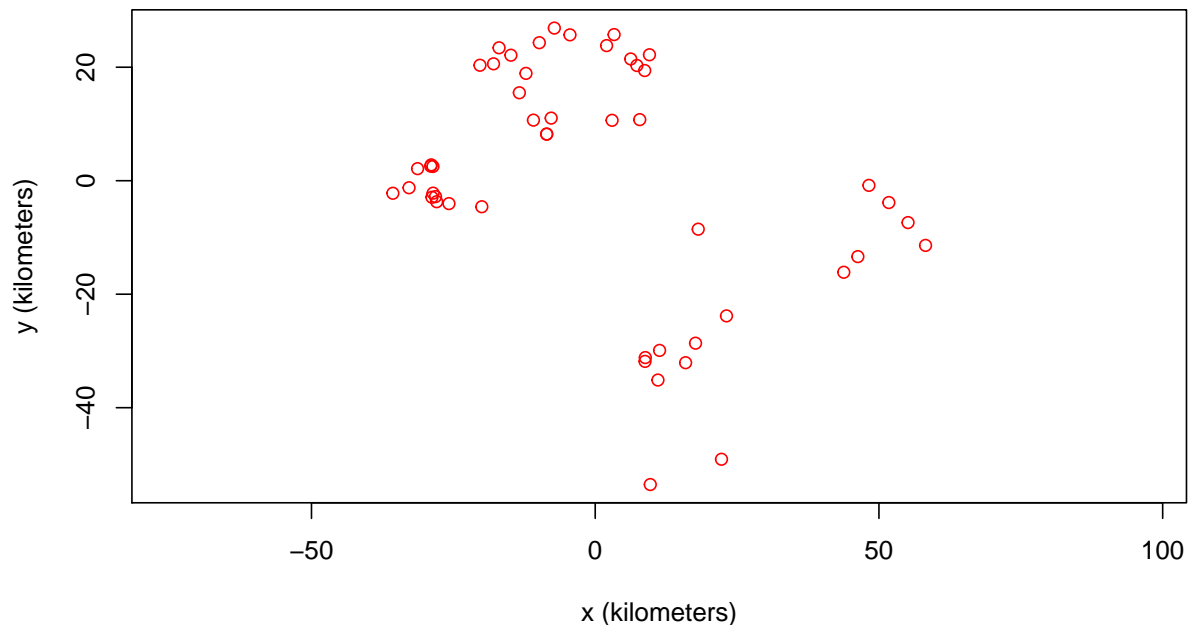
This dataset showcases an irregular sampling schedule: the buffalo nicknamed “*Pepper*” had a sampling rate shift from one fix every hour to one fix every two hours. We will use this dataset to highlight data irregularity and the **wAKDE** mitigation measure.

1.2. Gazelle tracking data

```
data("gazelle")
animal2_gazelle <- gazelle[[11]] # selecting individual number 11
head(animal2_gazelle)
```

```
##           x           y           t
## 2742 18152.70 -8539.799           0
## 2743 15931.16 -32069.788 1306800
## 2744 17678.84 -28632.329 1396800
## 2745 23135.50 -23820.789 1486800
## 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**

and apply the **parametric bootstrap** mitigation.

Data Analysis

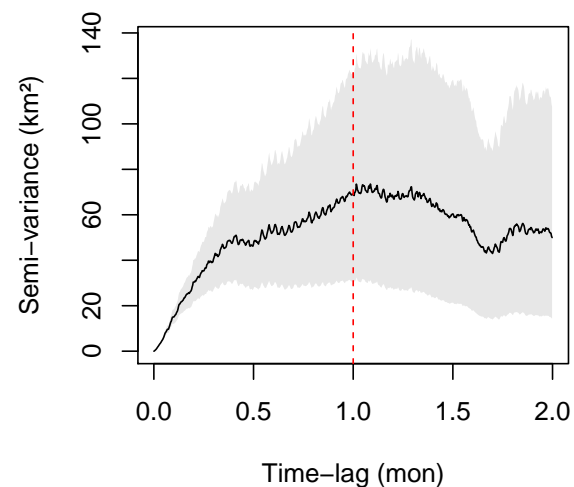
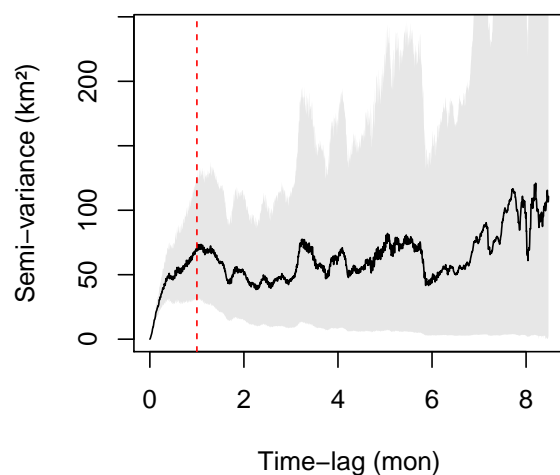
Step 2. – Checking for the *range residency* assumption

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)
```



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(animals1_buffalo, interactive = FALSE)

# Automated model selection, starting from GUESS:
FIT1_ML <- ctmm.select(animals1_buffalo, GUESS1, method = 'ML')
FIT1_pHREML <- ctmm.select(animals1_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      high
## 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 <- ctmselect(animal1_buffalo, GUESS1, verbose = TRUE)
summary(FIT1_pHREML_verbose)
```

	ΔAICc	$\Delta\text{RMSPE (km)}$	$\text{DOF}[\text{area}]$
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)
```

```
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
```

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)
```

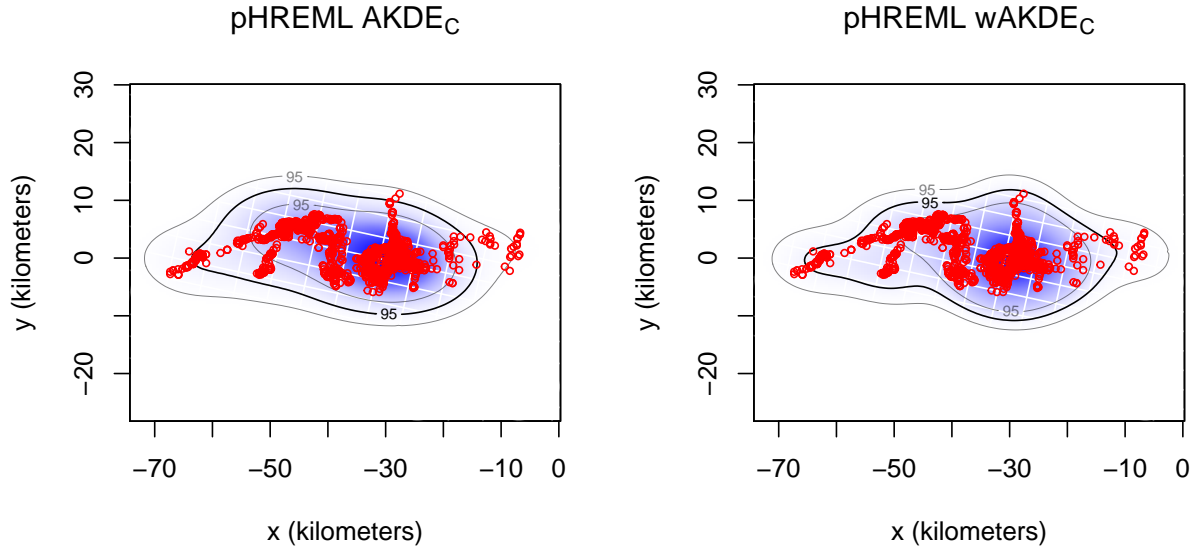
```
# Plotting pHREML (with and without weights) side-by-side:
par(mfrow = c(1,2))
plot(animal1_buffalo, UD = UD1_pHREML, ext = EXT)
```



```

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 <- ctm.guess(animal2_gazelle, interactive = FALSE)

FIT2_ML <- ctm.select(animal2_gazelle, GUESS2, method = 'ML')
FIT2_pHREML <- ctm.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 (~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
```

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
```

```
BOOT <- ctm.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
```

```
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)      0.000000      1.781242      3.758819
```

```
## t[velocity] (hours)       2.248230     10.481973     48.870349
```

```
## speed (kilometers/day)    5.590052      8.072361     10.551247
```

```
1/summary(BOOT)$DOF['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:

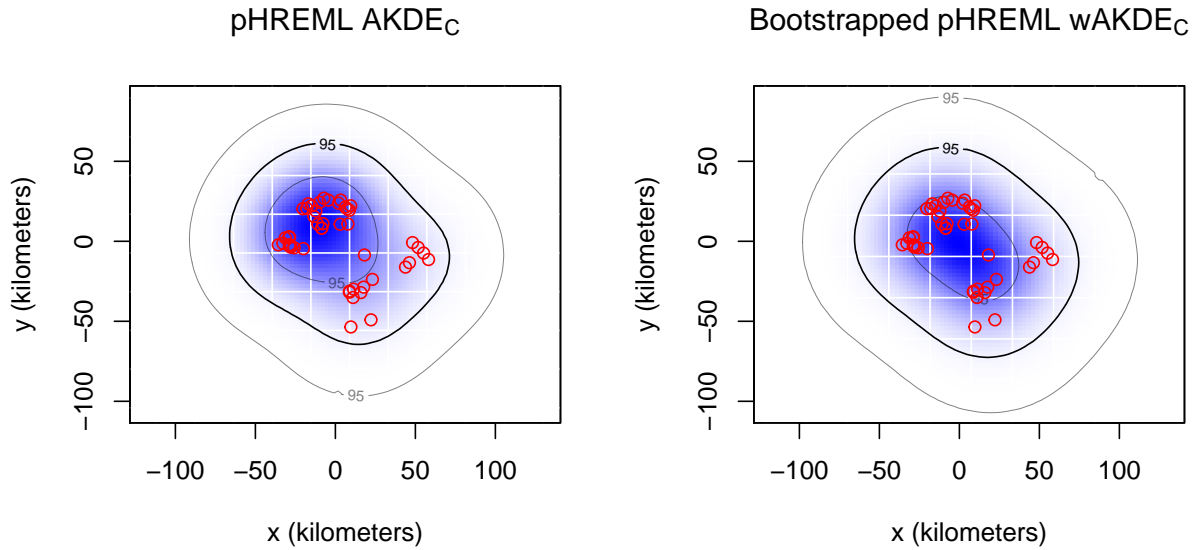
```
UD2_bpHREML <- akde(animal2_gazelle, BOOT, weights = TRUE)
summary(UD2_bpHREML)$CI
```

```
##                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"]))
```



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

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- 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.
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