Supplementary Material for A Guide to Pre-processing High-Frequency Animal Tracking Data

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Introduction

This is a rendering of the Supplementary Material for *A Guide to Pre-processing High-Frequency Animal Tracking Data*, and contains two fully worked out examples that could provide a useful template for structuring pre-processing pipelines for your own high-throughput tracking data.

The code and data used here are available on Github at github.com/pratikunterwegs/atlas-best-practices, and on Zendo at zenodo/atlas-best-practices.
1 Validating the Residence Patch Method with Calibration Data

Here we show how the residence patch method (Barraquand and Benhamou 2008; Bijleveld et al. 2016; Oudman et al. 2018) accurately estimates the duration of known stops in a track collected as part of a calibration exercise in the Wadden Sea. These data can be accessed from the data folder at this link: https://doi.org/10.5281/zenodo.4287462. These data are more fully reported in (Beardsworth et al. 2021).

1.1 Outline of Cleaning Steps

We begin by preparing the libraries we need, and installing atlastools from Github. After installing atlastools, we visualise the data to check for location errors, and find a single outlier position approx. 15km away from the study area (Fig. 1.1, 1.2). This outlier is removed by filtering data by the X coordinate bounds using the function atl_filter_bounds; X coordinate bounds \( \leq 645,000 \) in the UTM 31N coordinate reference system were removed (n = 1; remaining positions = 50,815; Fig. 1.2). We then calculate the incoming and outgoing speed, as well as the turning angle at each position using the functions atl_get_speed and atl_turning_angle respectively, as a precursor to targeting large-scale location errors in the form of point outliers. We use the function atl_filter_covariates to remove positions with incoming and outgoing speeds \( \geq \) the speed threshold of 15 m/s (n = 13,491, 26.5%; remaining positions = 37,324, 73.5%; Fig. 1.3; main text Fig. 7.b). This speed threshold is chosen as the fastest boat speed during the experiment, 15 m/s. Finally, we target small-scale location errors by applying a median smoother with a moving window size \( K = 5 \) using the function atl_median_smooth (Fig. 1.4; main text Fig. 7.c). Smoothing does not reduce the number of positions. We thin the data to a 30 second interval leaving 1,803 positions (4.8% positions of the smoothed track).

1.2 Install atlastools from Github

atlastools is available from Github and is archived on Zenodo (Gupte 2020). It can be installed using remotes or devtools. Here we use the remotes function.

```r
if (!require(remotes)) {
  install.packages("remotes", repos = "http://cran.us.r-project.org")
}

# installation using remotes
if (!require(atlastools)) {
  remotes::install_github("pratikunterwegs/atlastools", upgrade = FALSE)
}
```
A Note on :=

The atlastools package is based on data.table, to be fast and efficient (Dowle and Srinivasan 2020). A key feature is modification in place, where data is changed without making a copy. This is already implemented in R and will be familiar to many users as data_frame$column_name <- values.

The data.table way of writing this assignment would be data_frame[, column_name := values]. We use this syntax throughout, as it provides many useful shortcuts, such as multiple assignment:

data_frame[, c("col_a", "col_b") := list(values_a, values_b)]

Users can use this special syntax, and will find it convenient with practice, but there are no cases where users must use the data.table syntax, and can simply treat the data as a regular data.frame. However, users are advised to convert their data.frame to a data.table using the function data.table::setDT().

1.3 Prepare libraries

First we prepare the libraries we need. Libraries can be installed from CRAN if necessary.

# for data handling
library(data.table)
library(atlastools)
library(stringi)

# for recursion analysis
library(recurse)

# for plotting
library(ggplot2)
library(patchwork)

# making a colour palette
pal <- RColorBrewer::brewer.pal(5, "Set1")
pal[3] <- "seagreen"

1.4 Access data and preliminary visualisation

First we access the data from a local file using the data.table package (Dowle and Srinivasan 2020).

In all, we aim to keep three versions of the data: (1) data_raw, the entirely unprocessed data, (2) data, the working version, and (3) data_unproc, data that has been partially processed, but which is one step behind data. This allows us to better illustrate the pre-processing steps, and prevents us from irreversibly modifying our data — at best, we would have to re-run many pre-processing steps, and at worst, we might overwrite the original data on disk.

We look at the first few rows, using head(). We then visualise the raw data.
# read and plot example data

data <- fread("data/atlas1060_allTrials_annotated.csv")
data_raw <- copy(data)

# see raw data

head(data_raw)

Here we show how data can be easily visualised using the popular plotting package ggplot2. Note that we plot both the points (geom_point) and the inferred path between them (geom_path), and specify a geospatial coordinate system in metres, suitable for the Dutch Wadden Sea (UTM 31N; ESPG code:32631; coord_sf). We save the output to file for future reference.

Since plot code can become very lengthy and complicated, we omit showing further plot code in versions of this document rendered as PDF or HTML; it can however be seen in the online .Rmd version.

# plot data

fig_data_raw <-
  ggplot(data) +
  geom_path(aes(x, y),
            col = "grey", alpha = 1, size = 0.2
          ) +
  geom_point(aes(x, y),
             col = "grey", alpha = 0.2, size = 0.2
          ) +
  ggthemes::theme_few() +
  theme(
    axis.title = element_blank(),
    axis.text = element_blank()
  ) +
  coord_sf(crs = 32631)

# save figure

ggsave(fig_data_raw,
       filename = "supplement/figures/fig_calibration_raw.png",
       width = 185 / 25
)
1.5 Filter by bounding box

We first save a copy of the data, so that we can plot the unprocessed data with the cleaned data plotted over it for comparison. Here, data_unproc, data, and data_raw are still the same, since no pre-processing steps have been applied yet.

```r
# make a copy using the data.table copy function
data_unproc <- copy(data)
```

We then filter by a bounding box in order to remove the point outlier to the far south east of the main track. We use the `atl_filter_bounds` functions using the `x_range` argument, to which we pass the limit in the UTM 31N coordinate reference system. This limit is used to exclude all points with an X coordinate < 645,000.

We then plot the result of filtering, with the excluded point in black, and the points that are retained in green. After this stage, data is filtered and ‘ahead’ of data_raw and data_unproc, which are still the same. This pattern will repeat throughout this material.

```r
# remove inside must be set to FALSE
data <- atl_filter_bounds(
data = data,
x = "x", y = "y",
x_range = c(645000, max(data$x)),
remove_inside = FALSE
)
```
Figure 1.2: Removal of a point outlier using the function `atl_filter_bounds`. The point outlier (black point) is removed based on its X coordinate value, with the data filtered to exclude positions with an X coordinate < 645,000 in the UTM 31N coordinate system. Positions that are retained are shown in green.

1.6 Filter trajectories

1.6.1 Handle time

Time in ATLAS tracks is represented by 64-bit integers (type `long`) that specify time in milliseconds, starting from the beginning of 1970 (the UNIX epoch). This representation of time is called POSIX time and is usually specified in seconds, not milliseconds.

Since about 1.6 billion seconds have passed since the beginning of 1970, current POSIX times in milliseconds cannot be represented by R’s built-in 32-bit integers. A naive conversion results in truncation of out-of-range numbers leading to huge errors (dates many thousands of years in the future).

R does not natively support 64-bit integers. One option is to use the bit64 package, which adds 64-bit integer support to R.

A simpler solution is to convert the times to R’s built-in `double` data type (also called `numeric`), which uses a 64-bit floating point representation. This representation can represent integers with up to 16 digits without error; we only need 13 digits to represent the number of milliseconds since 1970, so the conversion is error free.

We can also perform the conversion and then divide by 1000 so that times are represented in seconds, not milliseconds; this simplifies speed estimation.

If second-resolution is accurate enough (it is for our purposes), the solution that we use is to divide times by 1000 to reduce the resolution from milliseconds to seconds and then to convert the time stamps to R integers. In the spirit of not destroying data, we create a second lower-case column called `time` to store this

```r
# divide by 1000, convert to integer
```
1.6.2 Add speed and turning angle

# add incoming and outgoing speed
data[, 'time' := as.integer(as.numeric(TIME) / 1000)]

# add turning angle
data[, angle := atl_turning_angle(data = data)]

1.6.3 Get 90th percentile of speed and angle

# use sapply
speed_angle_thresholds <- sapply(data[, list(speed_in, speed_out, angle)], quantile, probs = 0.9, na.rm = T)

1.6.4 Filter on speed

Here we use a speed threshold of 15 m/s, the fastest known boat speed. We then plot the data with the extreme speeds shown in grey, and the positions retained shown in green.

Here, data_unproc moves ‘ahead’ of data_raw, and holds the data filtered by a bounding box — data is also moving ahead, and will be filtered on speed.

# make a copy
data_unproc <- copy(data)

# remove speed outliers
data <- atl_filter_covariates(
  data = data,
  filters = c("(speed_in < 15 & speed_out < 15)"
)
)

# recalculate speed and angle
data[, 'time' := as.integer(as.numeric(TIME) / 1000)]
y = "y",
    time = "time"
),
speed_out = atl_get_speed(data, type = "out")
]

# add turning angle
data[, angle := atl_turning_angle(data = data)]

Figure 1.3: Improving data quality by filtering out positions that would require unrealistic movement. We removed positions with speeds \( \geq 15 \text{ m/s} \), which is the fastest possible speed in this calibration data, part of which was collected in a moving boat around Griend. Grey positions are removed, while green positions are retained. Rectangles indicate areas expanded for visualisation in following figures.

1.7 Smoothing the trajectory

We then apply a median smooth over a moving window \((K = 5)\). This function modifies in place, and does not need to be assigned to a new variable. We create a
copy of the data before applying the smooth so that we can compare the data before and after smoothing.

```r
# apply a 5 point median smooth, first make a copy
data_unproc <- copy(data)

# now apply the smooth
atl_median_smooth(
  data = data,
  x = "x", y = "y", time = "time",
  moving_window = 5
)
```

Figure 1.4: Reducing small-scale location error using a median smooth with a moving window \( K = 5 \). Median smoothed positions are shown in green, while raw, unfiltered data is shown in grey. Median smoothing successfully recovers the likely path of the track without a loss of data. The area shown is the upper rectangle from Fig. 1.3.

### 1.8 Thinning the data

Next we thin the data by aggregation to demonstrate thinning after median smoothing. Following this, we plot the median smooth and thinning by aggregation.

```r
# save a copy
data_unproc <- copy(data)

# remove columns we don't need
data <- data[, !c("tID", "Timestamp", "id", "TIME", "UTCtime")]

# thin to a 30s interval
data_thin <- `atl_thin_data`(
  data = data,
  interval = 30,
  method = "aggregate",
  id_columns = "TAG"
)

Figure 1.5: Thinning by aggregation over a 30 second interval (down from 1 second) preserves track structure while reducing the data volume for computation. Here, thinned positions are shown as purple squares, with the size of the square indicating the number of positions within the 30 second bin used to obtain the average position. Green points show the median smoothed data from Fig. 1.4, while the raw data are shown in grey. The area shown is the upper rectangle in Fig. 1.3.

1.9 Residence patches

1.9.1 Get waypoint centroids

We subset the annotated calibration data to select the waypoints and the positions around them which are supposed to be the locations of known stops. Since each stop was supposed to be 5 minutes long, there are multiple points in each known stop.

data_res <- `data_unproc`[`stri_detect`(tID, regex = "(WP)")]

From this data, we get the centroid of known stops, and determine the time difference between the first and last point within 50 metres, and within 10 minutes of the waypoint positions’ median time.

Essentially, this means that the maximum duration of a stop can be 20 minutes, and stops above this duration are not expected.
# get centroid
data_res_summary <- data_res[, list(
  nfixes_real = .N,
  x_median = median(x),
  y_median = median(y),
  t_median = median(time)
),
by = "tID"
]

# now get times 10 mins before and after
data_res_summary[, c("t_min", "t_max") := list(
  t_median - (10 * 60),
  t_median + (10 * 60)
)]

# manually get the duration of the stops
wp_data <- mapply(function(l, u, mx, my) {
  # first select all data whose timestamp is between
  # the upper and lower bounds of the stop (l = lower, u = upper)
  tmp_data <- data_unproc[inrange(time, l, u), ]

  # calculate the distance between the positions selected above
  # and the median X and Y coordinates of the stop (centroid)
  tmp_data[, distance := sqrt((mx - x)^2 + (my - y)^2)]

  # keep positions that are within 50m of the centroid
  tmp_data <- tmp_data[distance <= 50, ]

  # get the duration of the stop as the difference between
  # the minimum and maximum times of the positions retained above
  return(diff(range(tmp_data$time)))}, data_res_summary$t_min, data_res_summary$t_max,
data_res_summary$x_median, data_res_summary$y_median,
# this specifies that a vector, rather than a list, is returned
SIMPLIFY = TRUE)

# get waypoint summary --- rounding median coordinates to the nearest 100m
patch_summary_real <- data_res_summary[, list(
  nfixes_real = nfixes_real,
  x_median = round(median(x_median), digits = -2),
  y_median = round(median(y_median), digits = -2)
),
by = "tID"
]

# add real duration
patch_summary_real[, duration_real := wp_data]
# write to file
fwrite(patch_summary_real, "data/data_real_watlas_stops.csv")

## 1.9.2 Prepare data

First, we filter data where we know the animal (or in this case, the human-carried tag) spent some time at or near a position, as this is the first step to identify residence patches. One way of doing this is by filtering out positions with speeds above which the tag (ideally on an animal) is likely to be in transit. Rather than filtering on instantaneous speed estimates, filtering on a median smoothed speed estimate is more reliable.

## 1.9.3 Exclude transit points

Here, we aim to remove locations where the tag is clearly moving, by filtering on smoothed speed, using a one-way median smooth with \( K = 5 \). The speeds between points must be recalculated here because the speed metrics now associated with the data refer to the raw data before median smoothing.

```r
# get 4 column data
data_for_patch <- copy(data_thin)

# recalculate speeds, removing speed out
data_for_patch[, c("speed_in", "speed_out") := list(
  atl_get_speed(data_for_patch),
  NULL
)]

# get smoothed speed
data_for_patch[, speed_smooth := runmed(speed_in, k = 5)]

# save recurse data
fwrite(data_for_patch, file = "data/data_calib_for_patch.csv")
```

## 1.9.4 Run residence patch method

We subset data with a smoothed speed < 2 m/s in order to construct residence patches. From this subset, we construct residence patches using the parameters: \( \text{buffer\_radius} = 5 \text{ metres}, \text{lim\_spat\_indep} = 50 \text{ metres}, \text{lim\_time\_indep} = 5 \text{ minutes}, \) and \( \text{min\_fixes} = 3. \)

```r
# assign id as tag
data_for_patch[, id := as.character(TAG)]

# on known residence points
patch_res_known <- atl_res_patch(
  data = data_for_patch[speed_smooth < 2, ],
  buffer_radius = 5,
  lim_spat_indep = 50,
  lim_time_indep = 5,
)
A note on summary statistics

Users specifying a summary_variable should make sure that the variable for which they want a summary statistic is present in the data. For instance, requesting mean speed by passing summary_variable = "speed" and summary_function = "mean" to atl_res_patch, should make sure that their data includes a column called speed.

1.9.5 Get spatial and summary objects

Having classified slow-moving or stationary behavioural bouts into residence patches, many animal ecologists would most probably wish to know something about the environment at or around these patches — more accurately, around the point locations classified into patches.

How exactly this is done depends on the relative spatial scales of the residence patches and the resolution of the environmental data layer. For instance, a residence patch some 40m – 50m wide or long may be overlaid on an environmental raster layer with a resolution of 250m. In this case, sampling the layer at the centroid of the patch is as good as sampling at all the patch’s points – the mean is unlikely to differ (except at raster pixel boundaries).

On the other hand, a raster with a 10m resolution (e.g. Sentinel 1 and 2 data) may be worthwhile to sample at all the locations comprising a residence patch, so as to calculate the mean and variance of environmental conditions.

Furthermore, many (if not all) animals integrate cues from quite a distance (10m – 100m) when making decisions on when to settle in an area, and when to leave. Thus it can also be useful to sample environmental layers not at point locations, but to extract the mean and variance from an area, or a buffer, around the animal’s point locations.

We have provided a convenient function to get either (1) the points (classified into patches), or (2) a summary output of the residence patches (i.e., the median coordinates and their attributes), or finally (3) a spatial buffer around the points from (1). This function, atl_patch_summary implements these options using the which_data argument, where the options are (1) “points”, (2) “summary”, or (3) “spatial”.

Here, we choose option (3), using a spatial buffer of 20m. The distance of the buffer is passed to the argument buffer_radius.

```r
# for the known and unknown patches
patch_sf_data <- atl_patch_summary(patch_res_known,
                                   which_data = "spatial",
                                   buffer_radius = 20
)

# assign crs
sf::st_crs(patch_sf_data) <- 32631

# get summary data
```
At this stage, users have successfully pre-processed their data from raw positions to residence patches. Residence patches are essentially sf objects and can be visualised using the sf method for plot; for instance plot(patch_sf_data). Further sections reproduce the analyses in the main manuscript.

### 1.9.6 Prepare to plot data

We read in the island’s shapefile to plot it as a background for the residence patch figure.

```r
# read griend and hut
griend <- sf::st_read("data/griend_polygon/griend_polygon.shp", quiet = TRUE)
hut <- sf::st_read("data/griend_hut.gpkg", quiet = TRUE)
```

### 1.10 Compare patch metrics

We filter these data to exclude one exceedingly long outlier of about an hour (WP080).

```r
# round median coordinate for inferred patches
patch_summary_inferred <-
  patch_summary_data[
    ,
    c(
      "x_median", "y_median",
      "nfixes", "duration", "patch"
    )
  ][, `:=`
    x_median = round(x_median, digits = -2),
    y_median = round(y_median, digits = -2)
  ]
```

We add data from the known patches, matching by X and Y median.

```r
# join with respatch summary
patch_summary_compare <-
  merge(patch_summary_real,
        patch_summary_inferred,
        on = c("x_median", "y_median"),
        all.x = TRUE, all.y = TRUE)
```

# drop nas
patch_summary_compare <- na.omit(patch_summary_compare)

# drop patch around WP080
patch_summary_compare <- patch_summary_compare[tID != "WP080", ]
Figure 1.6: Classifying thinned data into residence patches yields robust estimates of the duration of known stops. The island of Griend (53.25°N, 5.25°E) is shown in beige. Residence patches (green polygons; function parameters in text) correspond well to the locations of known stops (purple triangles). However, the algorithm identified all areas with prolonged residence, including those which were not intended stops (n = 12; green polygons without triangles). The field station on Griend (red triangle) was not intended to be a stop, but the tags were stored here before the trial, and the method correctly picked up this prolonged stationary data as a residence patch. The algorithm failed to find two stops of 6 and 15 seconds duration, since these were lost in the data thinning step (purple triangle without green polygon shows one of these). The area shown is the lower rectangle in Fig. 1.3.
7 patches are identified where there are no waypoints, while 2 waypoints are not identified as patches. These waypoints consisted of 6 and 15 (WP098 and WP092) positions respectively, and were lost when the data were aggregated to 30 second intervals.

1.10.1 Linear model durations

We run a simple linear model.

```r
# get linear model
model_duration <- lm(duration_real ~ duration,
  data = patch_summary_compare)

# get R2
summary(model_duration)
```

```
Call:
  lm(formula = duration_real ~ duration, data = patch_summary_compare)

Residuals:
  Min     1Q    Median     3Q    Max
-105.07 -16.51    -4.26    9.54   91.66

Coefficients:
  Estimate Std. Error t value Pr(>|t|)
(Intercept) 102.4395  47.4097  2.16  0.046 *
duration   1.0225   0.0786 13.02  6.3e-10 ***
---
Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Residual standard error: 50.2 on 16 degrees of freedom
Multiple R-squared: 0.914, Adjusted R-squared: 0.908
F-statistic: 169 on 1 and 16 DF, p-value: 6.29e-10
```

# write to file
writeLines(
  text = capture.output(
    summary(model_duration)
  ),
  con = "data/model_output_residence_patch.txt"
)

1.10.2 Linear model summary

```r
cat(
  readLines(
    con = "data/model_output_residence_patch.txt",
    encoding = "UTF-8"
  ),
  sep = "\n"
)
Figure 1.7: The inferred duration of residence patches corresponds very closely to the real duration (grey circles, red line shows linear model fit), with an underestimation of the true duration of around 2%. The dashed black line represents $y = x$ for reference.

```r
#>
#> Call:  
#> lm(formula = duration_real ~ duration, data = patch_summary_compare)  
#>  
#> Residuals:  
#>    Min 1Q Median 3Q Max  
#> -105.07 -16.51 -4.26 9.54 91.66  
#>  
#> Coefficients:  
#>             Estimate Std. Error t value Pr(>|t|)  
#> (Intercept) 102.4395   47.4097  2.16  0.046 *  
#> duration   1.0225    0.0786 13.02 6.3e-10 ***  
#>  
#> Residual standard error: 50.2 on 16 degrees of freedom  
#> Multiple R-squared: 0.914, Adjusted R-squared: 0.908  
#> F-statistic: 169 on 1 and 16 DF, p-value: 6.29e-10
```

1.11 Main text Figure 7

See main text for Figure 7. Plotting code is not shown in PDF and HTML form, see the `.Rmd` file.
2 Processing Egyptian Fruit Bat Tracks

We show the pre-processing pipeline at work on the tracks of three Egyptian fruit bats (*Rousettus aegyptiacus*), and construct residence patches.

2.1 Prepare libraries

Install the required R libraries that are required from CRAN if not already installed.

```r
# libs for data
library(data.table)
library(RSQLite)
library(atlastools)

# libs for plotting
library(ggplot2)
library(patchwork)

# recursion analysis
library(recurse)

# prepare a palette
pal <- pals::kovesi.rainbow_bgyr_35_85_c73(3)
if (!require(remotes)) {
    install.packages("remotes", repos = "http://cran.us.r-project.org")
}

# installation using remotes
if (!require(atlastools)) {
    remotes::install_github("pratikunterwegs/atlastools", upgrade = FALSE)
}
```

2.2 Read bat data

Read the bat data from an SQLite database local file and convert to a plain text csv file. This data can be found in the “data” folder.

```r
# prepare the connection
con <- dbConnect(
    drv = SQLite(),
    dbname = "data/Three_example_bats.sql"
)

# list the tables
table_name <- dbListTables(con)

# prepare to query all tables
query <- sprintf('select * from %s', table_name)

# query the database
data <- dbGetQuery(conn = con, statement = query)

# disconnect from database
dbDisconnect(con)

Convert data to csv, and save a local copy in the folder “data”.

# convert data to datatable
setDT(data)

# write data for QGIS
fwrite(data, file = "data/bat_data.csv")

2.3 Exploratory Data Analysis Panels: Main Text Figure 1

Here, we make some basic figures for exploratory data analysis shown in Figure 1 of the main text.

Plot the bat data as a sanity check, and inspect it visually for errors. The plot code is hidden in the rendered copy (PDF) of this supplementary material, but is available in the Rmarkdown file “supplement/06_bat_data.Rmd”.

2.3.1 Heatmap of Locations

Here we demonstrate a basic heatmap of locations, aggregating over all individuals. In this one instance, the plotting code is also shown as a guide for readers, but in general, plotting code is hidden throughout this document.

data_heatmap <- copy(data)
data_heatmap[, c("xround", "yround") := list(
    plyr::round_any(X, 250),
    plyr::round_any(Y, 250)
)]
data_heatmap <- data_heatmap[, .N, by = c("xround", "yround")]

fig_heatmap <-
ggplot(data_heatmap) +
  geom_tile(
    aes(xround, yround, fill = N),
    size = 0.1,
    show.legend = F
  ) +
  scale_fill_viridis_c(
    option = "A",
    ..., 21
direction = -1,
    trans = "log10"
  ) +
  theme_void() +
  coord_sf(crs = 2039)

  ggsave(fig_heatmap,
    filename = "supplement/figures/fig_bat_heatmap_raw.png",
    dpi = 300,
    width = 6, height = 4
  )

2.3.2 Sampling Intervals

Here, we create the histogram of sampling intervals shown in Figure 1 of the main text. The plotting code is hidden in the PDF version, but available in the source code.

2.3.3 Localisation Error Measured by Systems

Here, we create the histogram of location error (variance in X) (Weiser et al. 2016) shown in Figure 1 of the main text. The plotting code is hidden in the PDF version, but available in the source code.

2.3.4 Plot paths from raw tracking data

Here, we plot the paths of individual bats from the raw tracking data to visually inspect them for errors.

  # this figure for the panel in main text figure 1
  fig_bat_focus_bad_speed <-
    fig_bat_raw +
    coord_sf(
      crs = 2039,
      xlim = c(253000, NA),
      ylim = c(772000, NA)
    )

  # save to supplement figures
  ggsave(fig_bat_focus_bad_speed,
    filename = "supplement/figures/fig_bat_focus_bad_speed.png",
    dpi = 300,
    width = 4, height = 6
  )

2.4 Prepare data for filtering

Here we apply a series of simple filters. It is always safer to deal with one individual at a time, so we split the data.table into a list of data.tables to avoid mixups among individuals.
Figure 2.1: Movement data from three Egyptian fruit bats tracked using the ATLAS system (*Rousettus aegyptiacus*; (Toledo et al. 2020; Shohami and Nathan 2020)). The bats were tracked in the Hula Valley, Israel (33.1°N, 35.6°E), and we use three nights of tracking (5th, 6th, and 7th May, 2018), for our demonstration, with an average of 13,370 positions (SD = 2,173; range = 11,195 – 15,542; interval = 8 seconds) per individual. After first plotting the individual tracks, we notice severe distortions, making pre-processing necessary.
This is a very rudimentary demonstration of the principle behind *batch processing* — splitting data into smaller, independent subsets, and applying the same steps to each subset.

### 2.4.1 Prepare data per individual

```r
# split bat data by tag
# first make a copy using the data.table function copy
# this prevents the original data from being modified by atlastools
# functions which DO MODIFY BY REFERENCE!
data_split <- copy(data)

# now split
data_split <- split(data_split, by = "TAG")
```

### 2.5 Filter by system-generated error attributes

No natural bounds suggest themselves, so instead we proceed to filter by system-generated attributes of error, since point outliers are obviously visible.

We use filter out positions with $SD > 20$ and positions calculated using only 3 base stations, using the function `atl_filter_covariates`.

First we calculate the variable $SD$, which for ATLAS systems is calculated as:

$$SD = \sqrt{VARX + VARY + 2 \times COVXY}$$

```r
# get SD.
# since the data are data.tables, no assignment is necessary
invisible(
  lapply(data_split, function(dt) {
    dt[, SD := sqrt(VARX + VARY + (2 * COVXY))]
  })
)
```

Then we pass the filters to `atl_filter_covariates`. We apply the filter to each individual’s data using `lapply` – this separates the data from each individual into a separate data frame, lessening the chances of inter-individual mix-ups.

This is another basic example of the principles behind batch-processing, and could be parallelised using the R package *furrr* (see https://CRAN.R-project.org/package=furrr).

```r
# filter for SD <= 20
# here, reassignment is necessary as rows are being removed
data_split <- lapply(data_split, function(dt) {
  dt <- atl_filter_covariates(
    data = dt,
    filters = c("SD <= 20",
               "NBS > 3")
  )
})
```
2.5.1 Sanity check: Plot filtered data

We plot the data to check whether the filtering has improved the data (Fig. 2.2). The plot code is once again hidden in this rendering, but is available in the source code file.

2.6 Filter by speed

Some point outliers remain, and could be removed using a speed filter.

First we calculate speeds, using `atl_get_speed`. We must assign the speed output to a new column in the data.table, which has a special syntax which modifies in place, and is shown below. This syntax is a feature of the data.table package, not strictly of atlastools (Dowle and Srinivasan 2020).

```r
# get speeds as with SD, no reassignment required for columns
invisible(
  lapply(data_split, function(dt) {
    # first process time to seconds
    # assign to a new column
    dt[, time := floor(TIME / 1000)]

    dt[, ,:= (~ (speed_in = atl_get_speed(dt,
      x = "X", y = "Y",
      time = "time",
      type = "in" ),
      speed_out = atl_get_speed(dt,
      x = "X", y = "Y",
      time = "time",
      type = "out"
    )]
    )]
  )
)
```

Now filter for speeds > 20 m/s (around 70 km/h), passing the predicate (a statement return TRUE or FALSE) to `atl_filter_covariates`. First, we remove positions which have NA for their `speed_in` (the first position) and their `speed_out` (last position).

```r
# filter speeds
# reassignment is required here
data_split <- lapply(data_split, function(dt) {
  dt <- na.omit(dt, cols = c("speed_in", "speed_out"))

  dt <- atl_filter_covariates(
```
Figure 2.2: Bat data filtered for large location errors, removing observations with standard deviation $> 20$. Grey crosses show data that were removed. Since the number of base stations used in the location process is a good indicator of error (Weiser et al. 2016), we also removed observations calculated using fewer than four base stations. Both steps used the function \texttt{atl_filter_covariates}. This filtering reduced the data to an average of 10,447 positions per individual (78% of the raw data on average). However, some point outliers remain.
data = dt, filters = c(
    "speed_in <= 20",
    "speed_out <= 20"
)

2.6.1 Sanity check: Plot speed filtered data

The speed filtered data is now inspected for errors (Fig. 2.3). The plot code is once again hidden.

2.7 Median smoothing

The quality of the data is relatively high, and a median smooth is not strictly necessary. We demonstrate the application of a 5 point median smooth to the data nonetheless (Fig. 2.4).

Since the median smoothing function `atl_median_smooth` modifies in place, we first make a copy of the data, using `data.table`'s copy function. No reassignment is required, in this case. The `lapply` function allows arguments to `atl_median_smooth` to be passed within `lapply` itself.

In this case, the same moving window $K$ is applied to all individuals, but modifying this code to use the multivariate version `Map` allows different $K$ to be used for different individuals. This is a programming matter, and is not covered here further.

```r
# since the function modifies in place, we shall make a copy
data_smooth <- copy(data_split)

# split the data again
data_smooth <- split(data_smooth, by = "TAG")

# apply the median smooth to each list element
# no reassignment is required as THE FUNCTION MODIFIES IN PLACE!
invisible(

    # the function arguments to atl_median_smooth
    # can be passed directly in lapply

    lapply(
        X = data_smooth, FUN = atl_median_smooth, time = "time", moving_window = 5
    )
)
```
Figure 2.3: Bat data with unrealistic speeds removed. Notice, compared with the previous figure, that spikes of unrealistic movement in all three tracks have been removed. Grey crosses show data that were removed. We calculated the incoming and outgoing speed of each position using `atl_get_speed`, and filtered out positions with speeds > 20 m/s using `atl_filter_covariates`, leaving 10,337 positions per individual on average (98% from the previous step).
Figure 2.4: Bat data after applying a median smooth with a moving window $K = 5$. Grey circles show data prior to smoothing. The smoothing step did not discard any data.


2.7.1 Sanity check: Plot smoothed data

2.8 Making residence patches

2.8.1 Calculating residence time

First, the data is put through the `recurse` package to get residence time (Bracis, Bildstein, and Mueller 2018).

```r
# split the data
data_smooth <- split(data_smooth, data_smooth$TAG)
```

We calculated residence time, but since bats may revisit the same features, we want to prevent confusion between frequent revisits and prolonged residence.

For this, we stop summing residence times within Z metres of a location if the animal exited the area for one hour or more. The value of Z (radius, in `recurse` parameter terms) was chosen as 50m.

This step is relatively complicated and is only required for individuals which frequently return to the same location, or pass over the same areas repeatedly, and for which revisits (cumulative time spent) may be confused for residence time in a single visit.

While a simpler implementation using total residence time divided by the number of revisits is also possible, this does assume that each revisit had the same residence time.

```r
# get residence times
data_residence <- lapply(data_smooth, function(dt) {
  # do basic recurse -- refer to Bracis et al. (2018) Ecography
dt_recurse <- getRecursions(
    x = dt[, c("X", "Y", "time", "TAG")],
    radius = 50,
    timeunits = "mins"
  )

  # get revisit stats column provided as recurse output
dt_recurse <- setDT(
    dt_recurse[["revisitStats"]]
  )

  # count long absences from the each position
dt_recurse[, timeSinceLastVisit :=
    ifelse(is.na(timeSinceLastVisit), -Inf, timeSinceLastVisit)]
dt_recurse[, longAbsenceCounter := cumsum(timeSinceLastVisit > 5),
    by = .(coordIdx)
  ]

  # filter data to exclude revisits after the first long absence of 60 mins
dt_recurse <- dt_recurse[longAbsenceCounter < 1, ]

  # calculate the residence time as the sum of times inside
  # before the first 'long absence'
```

# also calculate the First Passage Time and the number of revisits
dt_recurse <- dt_recurse[, list(
  resTime = sum(timeInside),
  fpt = first(timeInside),
  revisits = max(visitIdx)
),
by = .(coordIdx, x, y)
]

# prepare to merge existing data with recursion data
dt[, coordIdx := seq(nrow(dt))]

dt <- merge(dt,
  dt_recurse[, c("coordIdx", "resTime")],
  by = c("coordIdx")
)

# ensure the data are ordered in ascending order of time
setorderv(dt, "time")

# print message when done
message(sprintf("TAG %s residence times done", unique(dt$TAG)))

# return the dataframe
dt

We bind the data together and assign a human readable timestamp column.

# bind the list
data_residence <- rbindlist(data_residence)

# get time as human readable
data_residence[, ts := as.POSIXct(time, origin = "1970-01-01")]

# get hour of day to filter for nighttime
data_residence[, hour := data.table::hour(ts)]

2.8.2 Movements away from the roost

To focus on night-time bat foraging around fruit trees, we shall filter data both on the
timestamps, to select night-time positions, and on the locations, to select positions >
1 km away from the roost-cave at Har Gershom (see main text Fig. 8).

Combining these two filters allows us to exclude bat positions at the roost-cave that
may be due to individual-differences in bats' departure or return times to and from
their foraging areas.

# read in roosts and select the Har Gershom roost
roosts <- fread("data/Roosts.csv")
setnames(roosts, "Species", "roost_name")
roosts <- roosts[roost_name == "Har Gershom"]
# define a simple distance function that is vectorised
get_distance_adhoc <- function(x1, y1, x2, y2) {
  sqrt(((x2 - x1)^2) + ((y2 - y1)^2))
}

# calculate distance to roost cave at Har Gershom
data_residence[, distance_roost] = get_distance_adhoc(
  x1 = roosts$X,
  x2 = data_residence$X,
  y1 = roosts$Y,
  y2 = data_residence$Y
)

Users should plot the data to examine the effect of applying filters — this code is shown, but the figure is hidden for brevity.

# plot for sanity check --- this plot is not shown
fig_roost.exclude <- ggplot(data_residence) +
  geom_point(
    aes(X, Y, col = distance_roost)
  ) +
  geom_point(
    data = roosts,
    aes(X, Y),
    shape = 21, size = 4,
    fill = "blue",
    alpha = 0.5
  ) +
  scale_colour_viridis_c(
    option = "B", direction = -1,
    trans = "log10",
    breaks = c(1000, 2500, 5000),
    labels = function(x) {
      scales::comma(
        x,
        scale = 0.001, suffix = "km"
      )
    },
    limits = c(1000, NA),
    na.value = "lightblue"
  ) +
  ggspatial::annotation_scale(location = "br") +
  theme_test() +
  theme(
    axis.text = element_blank(),
    axis.title = element_blank(),
    legend.position = "top",
    legend.key.height = unit(2, "mm"),
    legend.title = element_text(vjust = 1)
  )
We now filter the data to exclude both day-time data, as well as data that is < 1 km from the roost.

```r
# filter for hour between 8pm and 5am and distance > 1km
data_residence <- atl_filter_covariates(
  data = data_residence,
  filters = c(
    "hour > 20 | hour < 5",
    "distance_roost > 1000"
  )
)
```

### 2.8.3 Split data by night-id

We assign a night-id to each position, i.e., the night-time spanning two calendar days. We then filter for data with a residence time > 5 minutes, as we expect that a bat stopped at a location for more than 5 minutes is likely to be foraging.

```r
# split data into separate nights
data_residence[, night := 1 + c(0, cumsum(diff(hour) > 12)), by = "TAG"]

# filter for residence time > 5 minutes
data_residence <- data_residence[resTime > 5, ]
```

### 2.8.4 Constructing residence patches

Some preparation is required. First, the function requires columns x, y, time, and id, which we assign using the `data.table` syntax. The time column is already present, but the other columns need to be renamed to lower case.

```r
# add an id column
data_residence[, `:=`(id = TAG, 
  x = X, y = Y )]

# get mean residence time per id
data_residence[, list(
  mean_residence = mean(resTime),
  sd_residence = sd(resTime)
), by = "TAG"]
```

```r
#>          TAG mean_residence sd_residence
#> 1: 972001004424        238.4         116.8
#> 2: 972001004449          52.8          33.5
#> 3: 972001004452          53.5          38.5
```
# get mean residence time for all bats pooled

data_residence[, list(
    mean_residence = mean(resTime),
    sd_residence = sd(resTime)
)]

#> mean_residence sd_residence
#> 1: 133 123

# average positions per bat after removing transit points

data_residence[, list(.N), by = "TAG"][, list(mean_positions_ = mean(N))]

#> mean_positions_
#> 1: 5549

# split the data

data_residence <- split(data_residence, by = c("TAG", "night"))

We apply the residence patch method, using the default argument values
(lim_spat_indep = 100 (metres), lim_time_indep = 30 (minutes)). We change the
buffer_radius to 25 metres (twice the buffer radius is used, so points must be
separated by 50m to be independent bouts), and min_fixes = 3.

# segment into residence patches

data_patches <- lapply(data_residence, atl_res_patch,
    buffer_radius = 25,
    min_fixes = 3,
    summary_variable = c("night", "distance_roost"),
    summary_functions = c("mean", "sd")
)

2.8.5 Getting residence patch data

We extract the residence patch data as spatial sf-MULTIPOLYGON objects. These are
returned as a list and must be converted into a single sf object. These objects and
the raw movement data are shown in Fig. 2.5.

# get data spatial

data_spatials <- lapply(data_patches, atl_patch_summary,
    which_data = "spatial",
    buffer_radius = 25
)

# bind list

data_spatials <- rbindlist(data_spatials)

# convert to sf

library(sf)
data_spatials <- st_sf(data_spatials, sf_column_name = "polygons")

# assign a crs

st_crs(data_spatials) <- st_crs(2039)
2.8.6 Write patch spatial representations

```r
st_write(data_spatials,
    dsn = "data/data_bat_residence_patches.gpkg",
    append = FALSE
)
#> Deleting layer `data_bat_residence_patches' using driver `GPKG'
#> Writing layer `data_bat_residence_patches' to data source
#>  `data/data_bat_residence_patches.gpkg' using driver `GPKG'
#> Writing 45 features with 22 fields and geometry type Multi Polygon.
```

Write cleaned bat data.

```r
fwrite(rbindlist(data_smooth),
    file = "data/data_bat_smooth.csv"
)
```

Write patch summary.

```r
# get summary
patch_summary <- lapply(data_patches, atl_patch_summary)

# bind summary
patch_summary <- rbindlist(patch_summary)

# write
fwrite(
    patch_summary,
    "data/data_bat_patch_summary.csv"
)
```

2.8.7 Duration at foraging sites

We exclude the first and last patch of each day as being roosting related, and examine how much of the total foraging time (time between the remaining first and last patch) was spent at foraging sites. It follows that the remainder of the time must have been spent in transit, or otherwise not foraging.

```r
# make patch summary a datatable
setDT(patch_summary)

# get mean and sd of duration in patches
patch_summary[, list(
    mean_duration = mean(duration / 60),
    sd_duration = sd(duration / 60)
)]
#> mean_duration sd_duration
#> 1: 57 62.2

# assign night id
patch_summary[, c("hour", "day") := list(
    data.table::hour(as.POSIXct(time_start, origin = "1970-01-01")),
    data.table::mday(as.POSIXct(time_start, origin = "1970-01-01"))
)]
# get total foraging time
foraging_proportion <- patch_summary[, list(
  time_total_forage = (max(time_end) - min(time_start)) / 60,
  time_forage_site = sum(duration / 60)
),
by = c("id", "night_mean")
]

# get proportion of foraging that is at a foraging site
foraging_proportion[, list(
  mean_foraging_prop = mean(time_forage_site / time_total_forage),
  sd_foraging_prop = sd(time_forage_site / time_total_forage)
)]
#> mean_foraging_prop sd_foraging_prop
#> 1: 0.838 0.155

2.9 **Main text Figure 8**

See Fig. 8 in the main text, made with QGIS.
Figure 2.5: A visual examination of plots of the bats’ residence patches and linear approximations of paths between them showed that though all three bats roosted at the same site, they used distinct areas of the study site over the three nights (a). Bats tended to be resident near fruit trees, which are their main food source, travelling repeatedly between previously visited areas (b, c). However, bats also appeared to spend some time at locations where no fruit trees were recorded, prompting questions about their use of other food sources (b, c). When bats did occur close together, their residence patches barely overlapped, and their paths to and from the broad area of co-occurrence were not similar (c). Constructing residence patches for multiple individuals over multiple activity periods suggests interesting dynamics of within- and between-individual overlap (b, c).
3 References

Barraquand, Frédéric, and Simon Benhamou. 2008. “Animal Movements in Heterogeneous Landscapes: Identifying Profitable Places and Homogeneous Movement Bouts.” *Ecology* 89 (12): 3336–48. https://doi.org/10.1890/08-0162.1.

Beardsworth, Christine E., Evy Gobbens, Frank van Maarseveen, Bas Denissen, Anne Dekinga, Ran Nathan, Sivan Toledo, and Allert I. Bijleveld. 2021. “Validating a High-Throughput Tracking System: ATLAS as a Regional-Scale Alternative to GPS.” *bioRxiv*, February. Cold Spring Harbor Laboratory, 2021.02.09.430514. https://doi.org/10.1101/2021.02.09.430514.

Bijleveld, Allert Imre, Robert B MacCurdy, Ying-Chi Chan, Emma Penning, Richard M. Gabrielson, John Cluderay, Erik L. Spaulding, et al. 2016. “Understanding Spatial Distributions: Negative Density-Dependence in Prey Causes Predators to Trade-Off Prey Quantity with Quality.” *Proceedings of the Royal Society B: Biological Sciences* 283 (1828): 20151557. https://doi.org/10.1098/rspb.2015.1557.

Bracis, Chloe, Keith L. Bildstein, and Thomas Mueller. 2018. “Revisitation Analysis Uncovers Spatio-Temporal Patterns in Animal Movement Data.” *Ecography* 41 (11): 1801–11. https://doi.org/10.1111/ecog.03618.

Dowle, Matt, and Arun Srinivasan. 2020. *Data.Table: Extension of ‘data.Frame’*. Manual.

Gupte, Pratik Rajan. 2020. “Atlastools: Pre-Processing Tools for High Frequency Tracking Data.” Zenodo. https://doi.org/10.5281/ZENODO.4033154.

Oudman, Thomas, Theunis Piersma, Mohamed V. Ahmedou Salem, Marieke E. Feis, Anne Dekinga, Sander Holthuijsen, Job ten Horn, Jan A. van Gils, and Allert I. Bijleveld. 2018. “Resource Landscapes Explain Contrasting Patterns of Aggregation and Site Fidelity by Red Knots at Two Wintering Sites.” *Movement Ecology* 6 (1): 24–24. https://doi.org/10.1186/s40462-018-0142-4.

Shohami, David, and Ran Nathan. 2020. “Cognitive Map-Based Navigation in Wild Bats Revealed by a New High-Throughput Tracking System.” Dryad. https://doi.org/10.5061/DRODA.G4F4QRFN2.

Toledo, Sivan, David Shohami, Ingo Schiffner, Emmanuel Lourie, Yotam Orchan, Yoav Bartan, and Ran Nathan. 2020. “Cognitive Map-Based Navigation in Wild Bats Revealed by a New High-Throughput Tracking System.” *Science* 369 (6500). American Association for the Advancement of Science: 188–93. https://doi.org/10.1126/science.aax6904.

Weiser, Adi Weller, Yotam Orchan, Ran Nathan, Motti Charter, Anthony J. Weiss, and Sivan Toledo. 2016. “Characterizing the Accuracy of a Self-Synchronized Reverse-GPS Wildlife Localization System.” In *2016 15th ACM/IEEE International Conference on Information Processing in Sensor Networks (IPSN)*, 1–12. https://doi.org/10.1109/IPSN.2016.7460662.