

Step generation for fitting water buffalo step selection models

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Here we are generating random steps for fitting step selection models, and plotting the outputs of the step generation. We are using a Gamma distribution for the step lengths and a von Mises distribution for the turning angles.

Setup packages

```
library(tidyverse)
packages <- c("amt", "lubridate", "terra", "beeswarm", "tictoc")
walk(packages, require, character.only = T)
```

Import data and clean

```
## # A tibble: 6 x 4
##   id     time             lon   lat
##   <chr> <dttm>          <dbl> <dbl>
## 1 2005  2018-07-25 10:04:02 134. -12.3
## 2 2005  2018-07-25 11:04:23 134. -12.3
## 3 2005  2018-07-25 12:04:39 134. -12.3
## 4 2005  2018-07-25 13:04:17 134. -12.3
## 5 2005  2018-07-25 14:04:39 134. -12.3
## 6 2005  2018-07-25 15:04:27 134. -12.3
## [1] "Australia/Queensland"
```

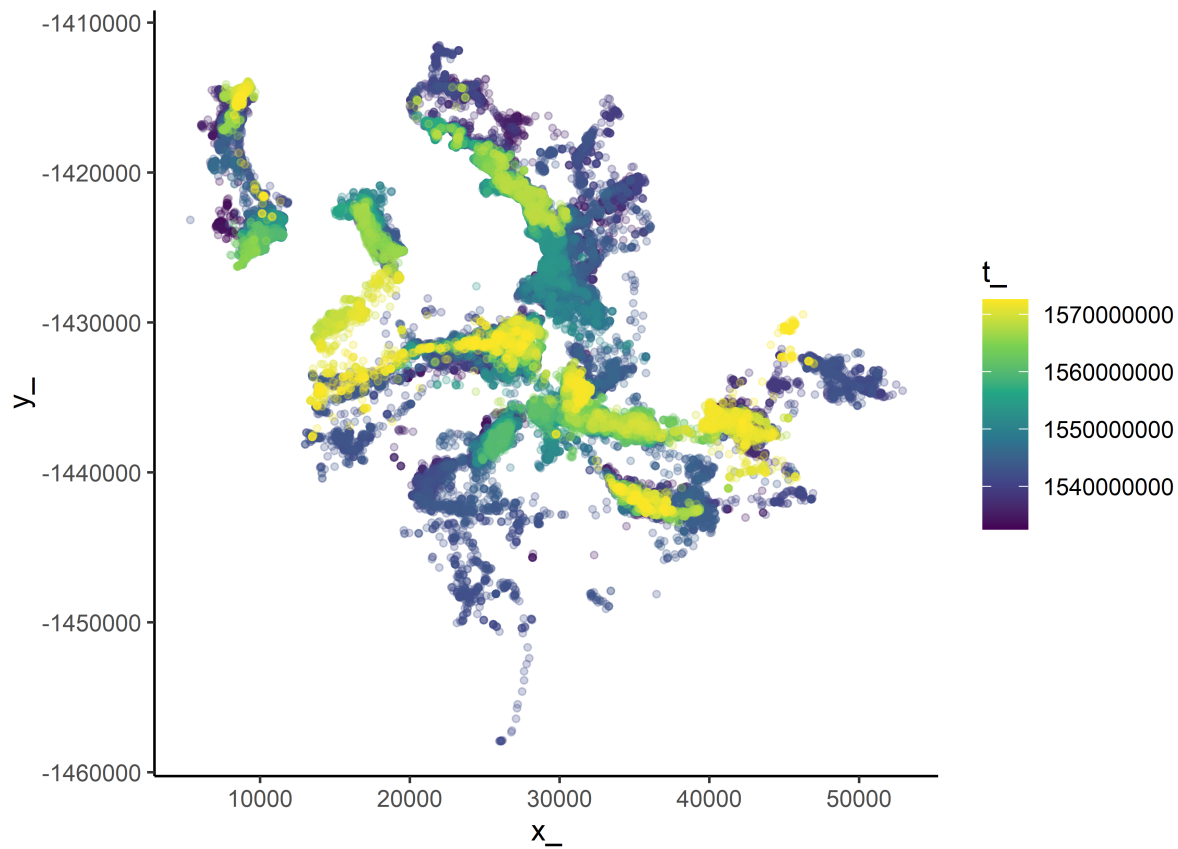
Setup trajectory

Use the `amt` package to create a trajectory object from the cleaned data.

```
buffalo_all <- buffalo_clean %>% mk_track(id = id,
                                              lon,
                                              lat,
                                              time,
                                              all_cols = T,
                                              crs = 4326) %>%
  transform_coords(crs_to = 3112, crs_from = 4326) # Transformation to GDA94 /
# Geoscience Australia Lambert (https://epsg.io/3112)

# plot the data spatially coloured by time
buffalo_all %>%
  ggplot(aes(x = x_, y = y_, colour = t_)) +
  geom_point(alpha = 0.25, size = 1) + # colour = "black",
  coord_fixed() +
```

```
scale_colour_viridis_c() +  
theme_classic()
```



Prepare environmental covariates

To determine required extent of raster to crop environmental variables to (with a buffer to accommodate the random steps)

```
min(buffalo_all$x_)  
  
## [1] 5337.494  
  
max(buffalo_all$x_)  
  
## [1] 52901.66  
  
min(buffalo_all$y_)  
  
## [1] -1457924  
  
max(buffalo_all$y_)  
  
## [1] -1411517  
  
template_raster_cropped <- terra::rast(xmin = 0,  
                                         xmax = 60000,  
                                         ymin = -1463000,  
                                         ymax = -1406000,
```

```

    resolution = 25,
    crs = "epsg:3112")

template_wgs84 <- project(template_raster_cropped, "epsg:4326")
terra::ext(template_wgs84)

## SpatExtent : 134.000008561276, 134.541785217172, -12.5424223864104, -12.0361830606666 (xmin, xmax, ym

```

Reading in the environmental covariates

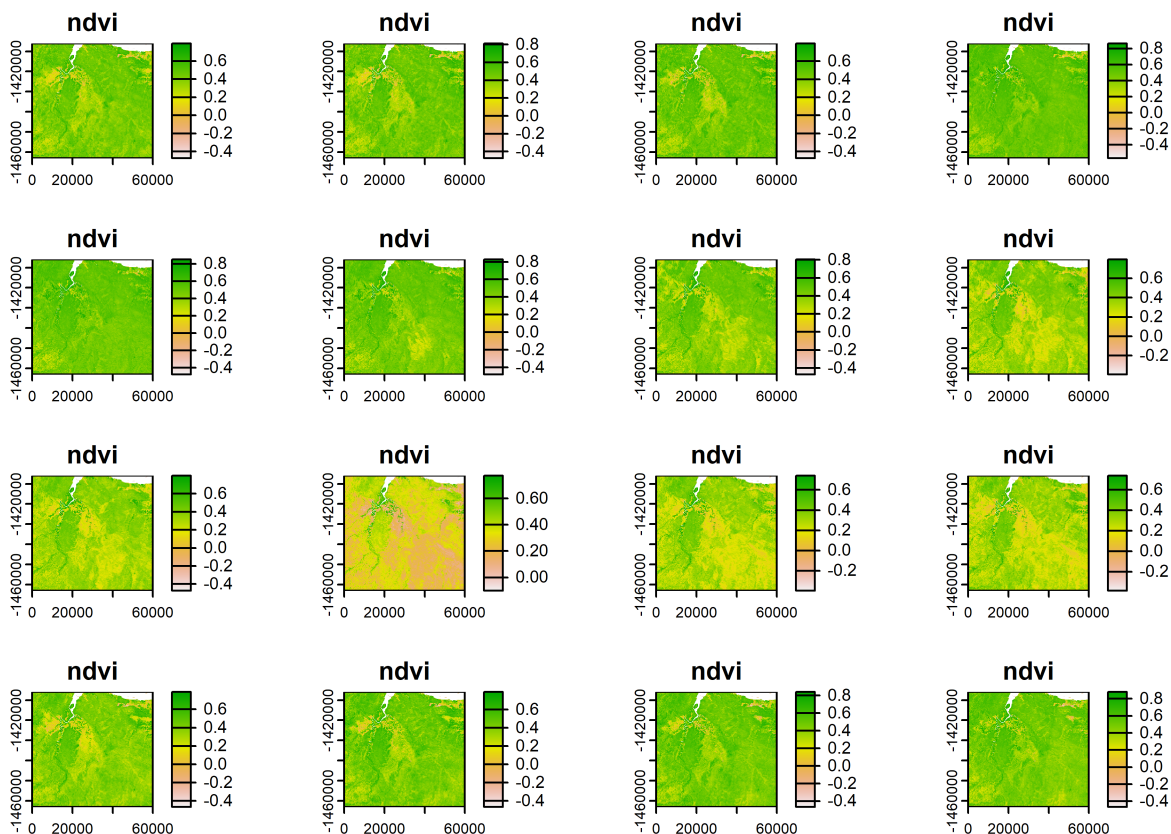
```

ndvi_projected <-
  rast("mapping/cropped rasters/ndvi_GEE_projected_watermask20230207.tif")
slope <- rast("mapping/cropped rasters/slope_raster.tif")
veg_herby <- rast("mapping/cropped rasters/veg_herby.tif")
canopy_cover <- rast("mapping/cropped rasters/canopy_cover.tif")

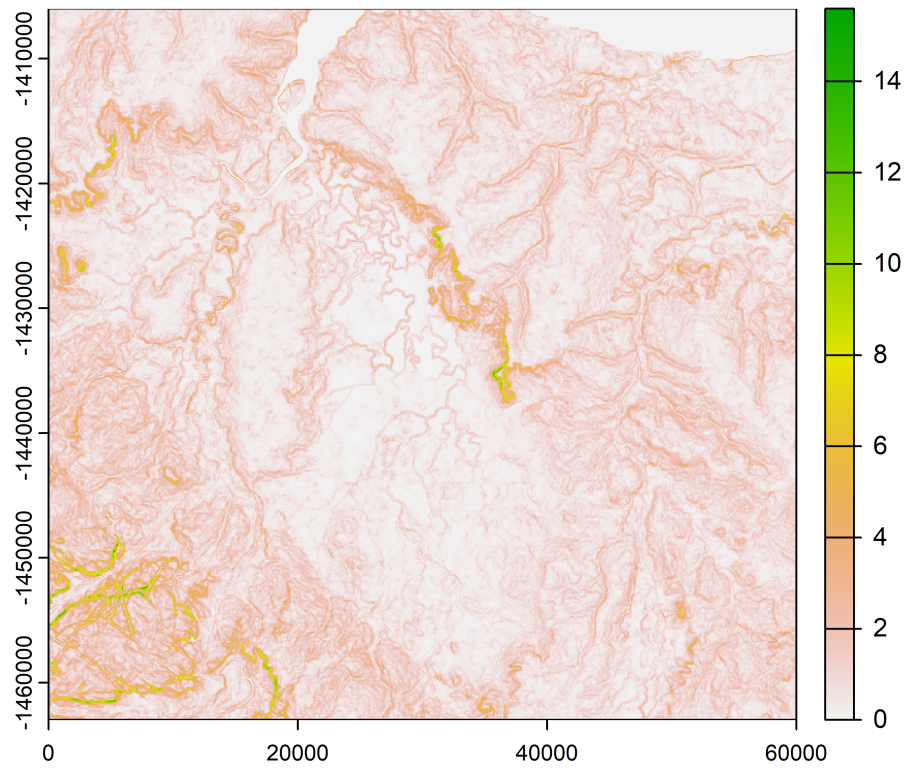
# change the names (these will become the column names when extracting
# covariate values at the used and random steps)
names(ndvi_projected) <- rep("ndvi", terra::nlyr(ndvi_projected))
names(slope) <- "slope"
names(veg_herby) <- "veg_herby"
names(canopy_cover) <- "canopy_cover"

# plot the rasters
plot(ndvi_projected)

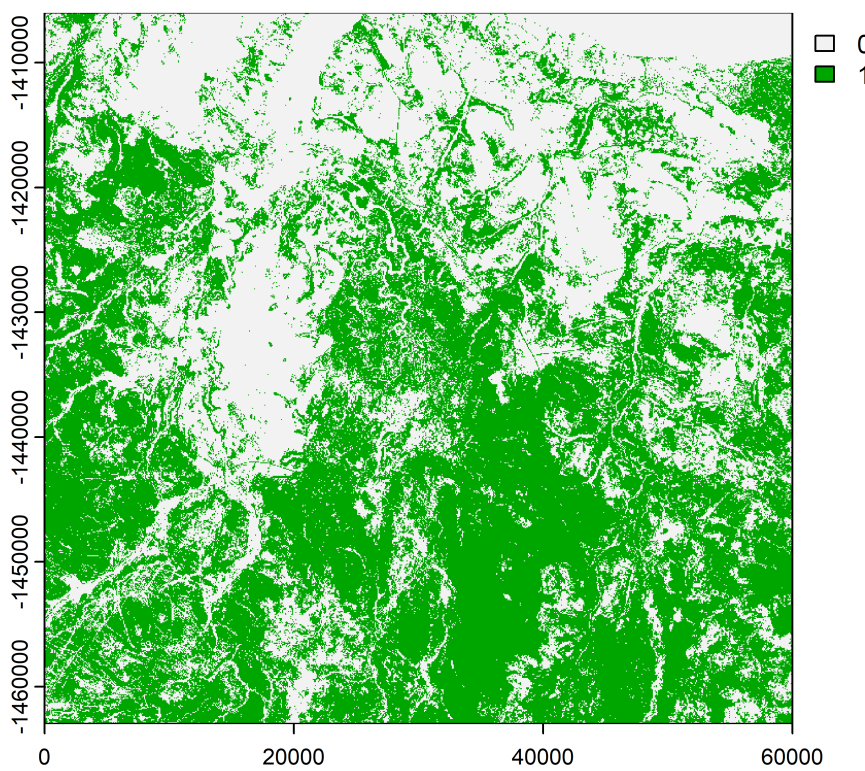
```



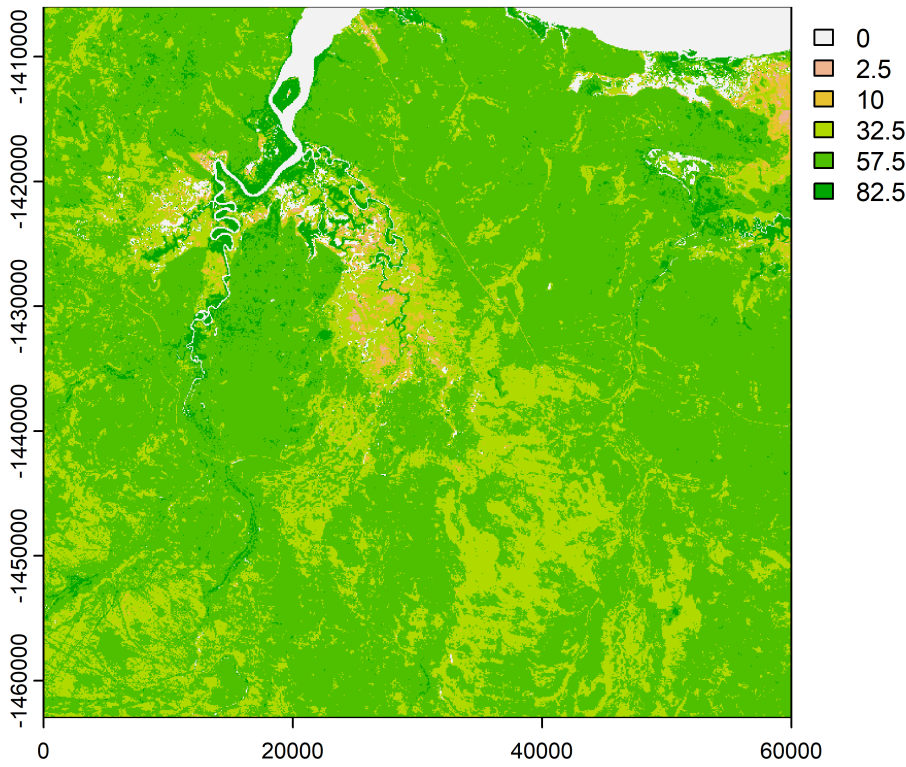
```
plot(slope)
```



```
plot(veg_herby)
```



```
plot(canopy_cover)
```



Creating a step object (by bursts)

```

# nest the data by individual
buffalo_all_nested <- buffalo_all %>% arrange(id) %>% nest(data = -"id")

buffalo_all_nested_steps_by_burst <- buffalo_all_nested %>%
  mutate(steps = map(data, function(x)
    x %>% track_resample(rate = hours(1), tolerance = minutes(10)) %>%
      # to filter out bursts with less than 3 locations
      # amt::filter_min_n_burst(min_n = 3) %>%
      steps_by_burst())))

```

unnest the data after creating 'steps' objects

```

buffalo_all_steps_by_burst <- buffalo_all_nested_steps_by_burst %>%
  amt::select(id, steps) %>%
  amt::unnest(cols = steps)

buffalo_all_steps_by_burst <- buffalo_all_steps_by_burst %>%
  mutate(t2_rounded = round_date(t2_, "hour"), # round the time to the nearest hour
         hour_t2 = ifelse(hour(t2_rounded) == 0, 24, hour(t2_rounded))) # change the 0 hour to 24

head(buffalo_all_steps_by_burst, 10)

```

```

## # A tibble: 10 x 14
##       id     burst_     x1_     x2_     y1_     y2_ sl_ direction_p     ta_
##   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl> <dbl> <dbl> <dbl>
## 1     1       1     100     100    -1435000 -1435000  100     1     100
## 2     1       1     100     100    -1435000 -1435000  100     1     100
## 3     1       1     100     100    -1435000 -1435000  100     1     100
## 4     1       1     100     100    -1435000 -1435000  100     1     100
## 5     1       1     100     100    -1435000 -1435000  100     1     100
## 6     1       1     100     100    -1435000 -1435000  100     1     100
## 7     1       1     100     100    -1435000 -1435000  100     1     100
## 8     1       1     100     100    -1435000 -1435000  100     1     100
## 9     1       1     100     100    -1435000 -1435000  100     1     100
## 10    1       1     100     100    -1435000 -1435000  100     1     100
## # ... with 9 more rows, and 4 more variables:
## #   direction_p <dbl>, sl_ <dbl>, t1_ <dbl>, t2_ <dbl>

```

```

## * <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dttm> <dttm>
## 1 2005 1 41940. 41968. -1.44e6 -1.44e6 206. 1.43 NA 2018-07-25 10:04:02 2018-0
## 2 2005 1 41968. 41921. -1.44e6 -1.44e6 50.7 2.80 1.37 2018-07-25 11:04:23 2018-0
## 3 2005 1 41921. 41778. -1.44e6 -1.44e6 152. 2.78 -0.0214 2018-07-25 12:04:39 2018-0
## 4 2005 1 41778. 41840. -1.44e6 -1.44e6 70.7 -0.507 2.99 2018-07-25 13:04:17 2018-0
## 5 2005 1 41840. 41655. -1.44e6 -1.44e6 188. 2.98 -2.80 2018-07-25 14:04:39 2018-0
## 6 2005 1 41655. 41618. -1.44e6 -1.44e6 37.1 -3.02 0.285 2018-07-25 15:04:27 2018-0
## 7 2005 1 41618. 41687. -1.44e6 -1.44e6 522. -1.44 1.58 2018-07-25 16:04:24 2018-0
## 8 2005 1 41687. 41683. -1.44e6 -1.44e6 8.13 2.10 -2.75 2018-07-25 17:04:28 2018-0
## 9 2005 1 41683. 41674. -1.44e6 -1.44e6 9.75 -3.13 1.05 2018-07-25 18:04:31 2018-0
## 10 2005 1 41674. 41424. -1.44e6 -1.44e6 308. 2.52 -0.635 2018-07-25 19:04:33 2018-0
## # i 2 more variables: t2_rounded <dttm>, hour_t2 <dbl>

```

Fitting step length and turning angle distributions

Fitting exponential and von Mises distributions to the steps of ALL individuals (only one Gamma and one von Mises distribution for the whole population). This should be done when fitting a hierarchical model to update the ‘population’ parameters, but also makes it straightforward to update after model fitting to each individual separately.

```

# fitting step length and turning angle distributions to all locations
gamma_dist <- fit_distr(buffalo_all_steps_by_burst$sl_, "gamma")
vonmises_dist <- fit_distr(buffalo_all_steps_by_burst$ta_, "vonmises")

# checking parameters - which can then be saved to update movement parameters
# after fitting the step selection model
gamma_dist$params$shape

## [1] 0.438167
gamma_dist$params$scale

## [1] 534.3507
vonmises_dist$params$kappa

## [1] 0.1848126
vonmises_dist$params$mu

## Circular Data:
## Type = angles
## Units = radians
## Template = none
## Modulo = asis
## Zero = 0
## Rotation = counter
## [1] 0

```

For some reason, the `random_steps` function does not work when using the `bursted_steps_xyt` class. I’m not sure why (the error is `Error in bursts[[i]] : subscript out of bounds`), but it works when that class label is removed, and appears to sample random steps correctly. For taxa that have irregular fixes and many bursts, this may be worth exploring in more detail.

```

class(buffalo_all_steps_by_burst)

## [1] "bursted_steps_xyt" "steps_xyt"           "steps_xy"          "tbl_df"           "tbl"

```

```

## [6] "data.frame"
class(buffalo_all_steps_by_burst) <- class(buffalo_all_steps_by_burst)[-1]
class(buffalo_all_steps_by_burst)

## [1] "steps_xyt"   "steps_xy"    "tbl_df"      "tbl"        "data.frame"

tic()
buffalo_parametric_popn_GvM <- buffalo_all_steps_by_burst %>%
  random_steps(n_control = 10,
                sl_distr = gamma_dist,
                ta_distr = vonmises_dist) %>%
  mutate(y = as.numeric(case_))
toc()

## 58.54 sec elapsed

```

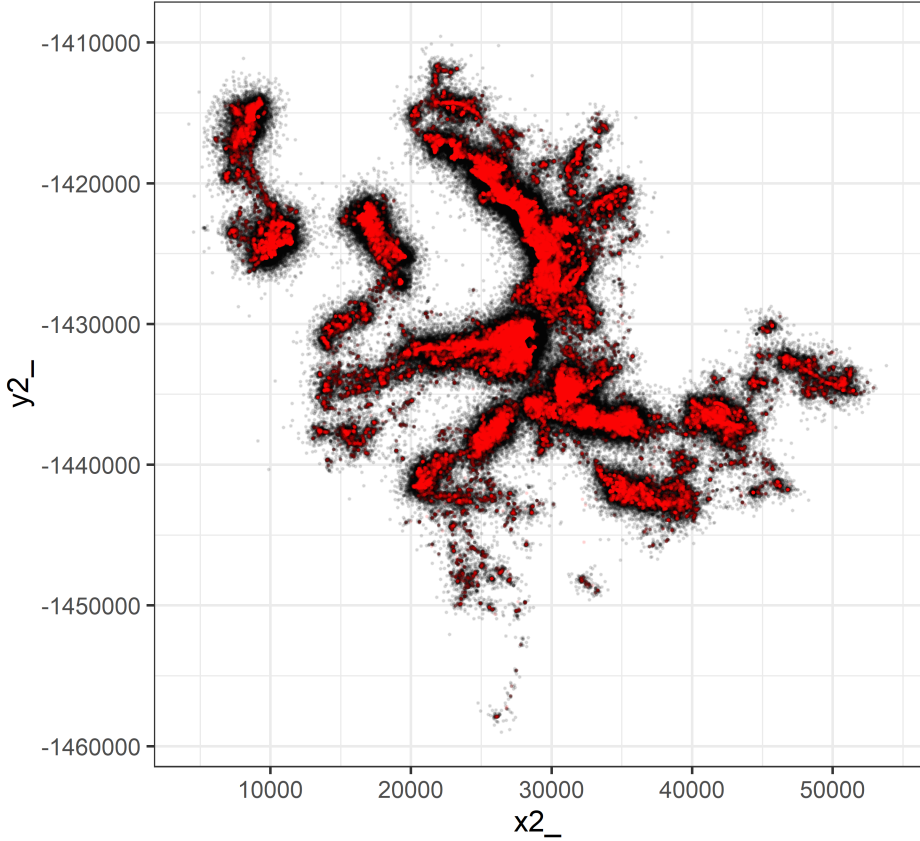
Plotting the random step distributions

Spatially plot the used and random steps, with the used steps in red.

```

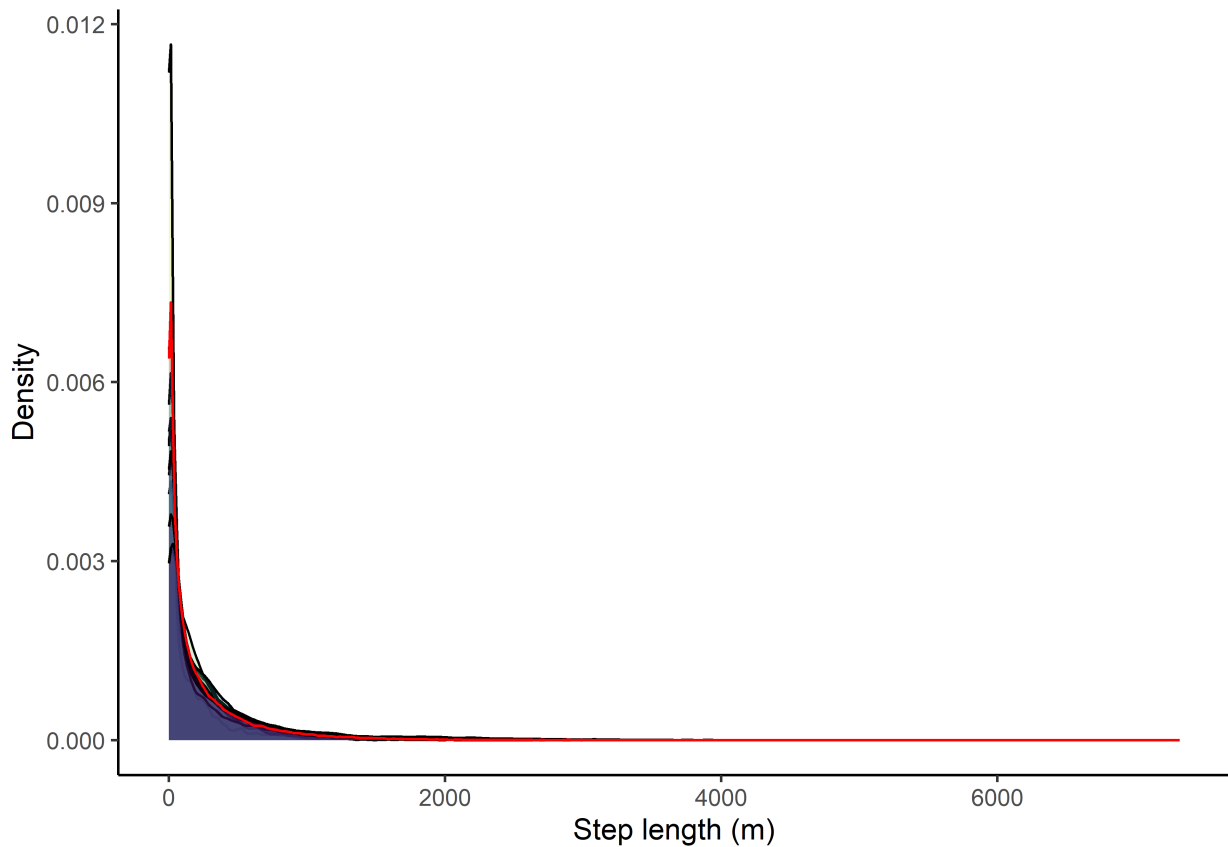
buffalo_parametric_popn_GvM %>% ggplot() +
  geom_point(data = . %>% filter(y == 0), aes(x = x2_, y = y2_),
             colour = "black", size = 0.1, alpha = 0.1) +
  geom_point(data = . %>% filter(y == 1), aes(x = x2_, y = y2_),
             colour = "red", size = 0.1, alpha = 0.1) +
  coord_equal() +
  theme_bw()

```



Gamma distribution of the step lengths

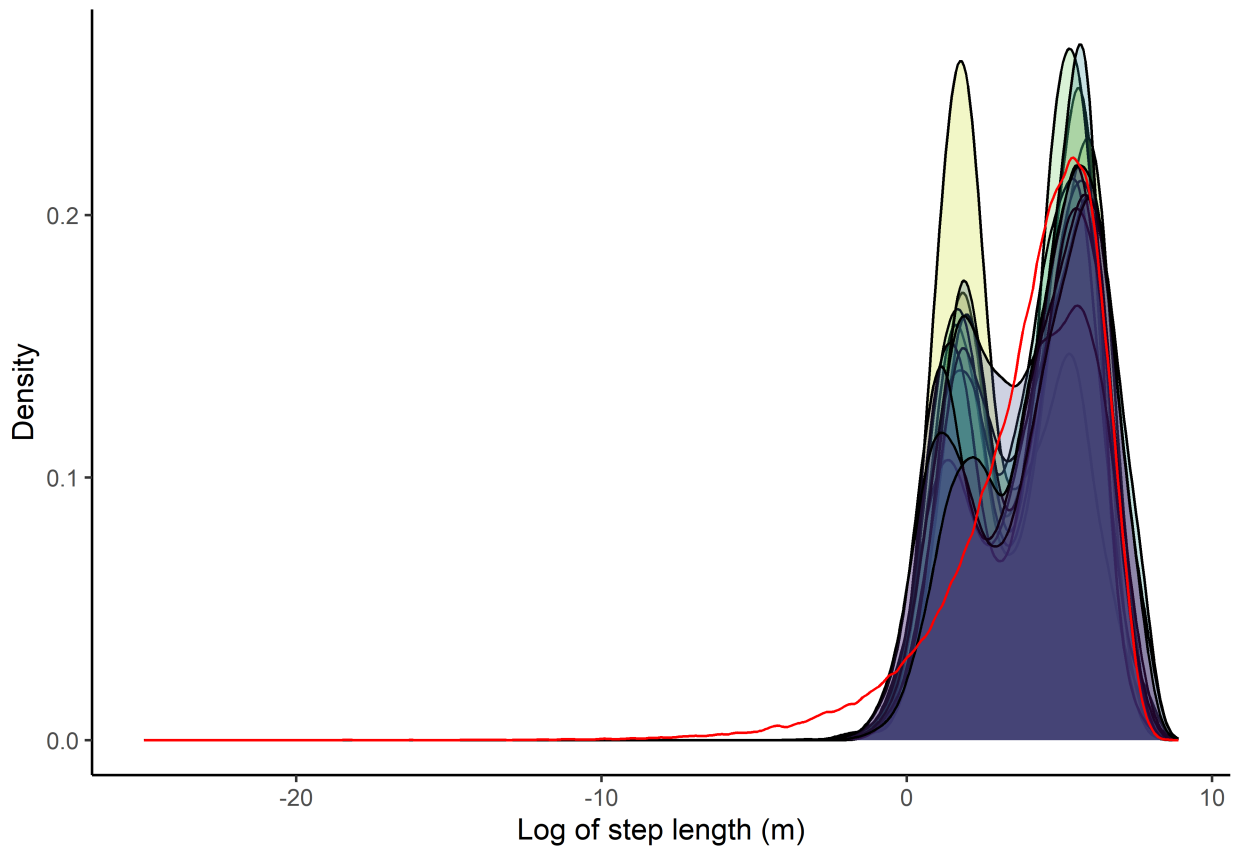
```
buffalo_parametric_popn_GvM %>% ggplot() +
  geom_density(data = . %>% filter(y == 1), aes(x = sl_, fill = factor(id)),
               alpha = 0.25) +
  geom_density(data = . %>% filter(y == 0), aes(x = sl_), colour = "red") +
  scale_y_continuous("Density") +
  scale_x_continuous("Step length (m)") +
  scale_fill_viridis_d(direction = -1) +
  theme_classic() +
  theme(legend.position = "none")
```



```
ggsave(paste0("outputs/plots/manuscript_figs/step_length_density_",
              Sys.Date(), ".png"),
       width=150, height=90, units="mm", dpi = 600)
```

When log-transforming the step length, it is clear that there is a bimodal pattern to the step lengths, which is not captured by the Gamma distribution of the random steps.

```
buffalo_parametric_popn_GvM %>% ggplot() +
  geom_density(data = . %>% filter(y == 1), aes(x = log(sl_), fill = factor(id)),
               alpha = 0.25) +
  geom_density(data = . %>% filter(y == 0), aes(x = log(sl_)), colour = "red") +
  scale_y_continuous("Density") +
  scale_x_continuous("Log of step length (m)") +
  scale_fill_viridis_d(direction = -1) +
  theme_classic() +
  theme(legend.position = "none")
```

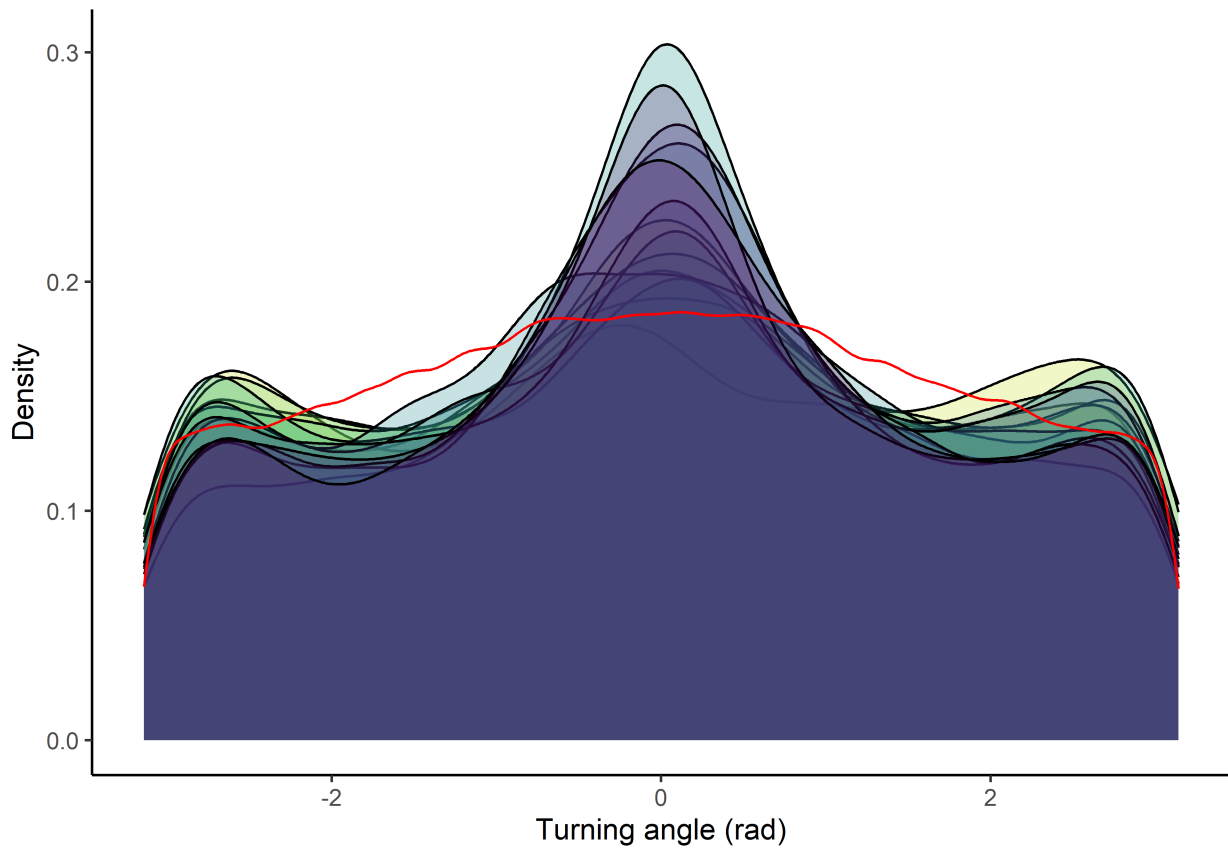


```
ggsave(paste0("outputs/plots/manuscript_figs/step_length_density_log_",
              Sys.Date(), ".png"),
       width=150, height=90, units="mm", dpi = 600)
```

Plot the von Mises distribution of the turning angles

```
buffalo_parametric_popn_GvM %>% ggplot() +
  geom_density(data = . %>% filter(y == 1), aes(x = ta_, fill = factor(id)),
               alpha = 0.25) +
  geom_density(data = . %>% filter(y == 0), aes(x = ta_), colour = "red",
               alpha = 0.01) +
  scale_y_continuous("Density") +
  scale_x_continuous("Turning angle (rad)") +
  scale_fill_viridis_d(direction = -1) +
  theme_classic() +
  theme(legend.position = "none")
```

```
## Warning: Removed 8053 rows containing non-finite values ('stat_density()').
```



```
ggsave(paste0("outputs/plots/manuscript_figs/turning_angle_density_",
              Sys.Date(), ".png"),
       width=150, height=90, units="mm", dpi = 600)
```

```
## Warning: Removed 8053 rows containing non-finite values ('stat_density()').
```

Sample values of the environmental covariates at the end of the steps.

```
buffalo_parametric_popn_covs <- buffalo_parametric_popn_GvM %>%
  extract_covariates_var_time(ndvi_projected,
    where = "end",
    when = "any",
    max_time = days(15),
    name_covar = "ndvi_temporal") %>%
  extract_covariates(veg_herby,
    where = "end") %>%
  extract_covariates(canopy_cover,
    where = "end") %>%
  extract_covariates(slope,
    where = "end") %>%
  mutate(y = as.numeric(case_),
    cos_ta_ = cos(ta_),
    log_sl_ = log(sl_))
```

```
# write the output to a csv file
write_csv(buffalo_parametric_popn_covs,
          paste0("outputs/buffalo_parametric_popn_covs_GvM_10rs_", Sys.Date(), ".csv"))

beep(sound = 2)
```