Package ‘spatsoc’

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Title  Group Animal Relocation Data by Spatial and Temporal Relationship

Version  0.2.2

Description  Detects spatial and temporal groups in GPS relocations (Robitaille et al. (2019) <doi:10.1111/2041-210X.13215>)
It can be used to convert GPS relocations to gambit-of-the-group format to build proximity-based social networks In addition, the randomizations function provides data-stream randomization methods suitable for GPS data.

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URL  https://docs.ropensci.org/spatsoc/, https://github.com/ropensci/spatsoc

BugReports  https://github.com/ropensci/spatsoc/issues

Depends  R (>= 3.4)

Imports  adehabitatHR (>= 0.4.21), data.table (>= 1.10.5), igraph, sf, stats, units

Suggests  asnipe, knitr, markdown, rmarkdown, testthat (>= 2.1.0)

VignetteBuilder  knitr

Encoding  UTF-8

RoxygenNote  7.2.3

SystemRequirements  GDAL (>= 2.0.1), GEOS (>= 3.4.0), PROJ (>= 4.8.0), sqlite3

NeedsCompilation  no

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build_lines generates a simple feature collection with LINESTRINGs from a data.table. The function accepts a data.table with relocation data, individual identifiers, a sorting column and a projection. The relocation data is transformed into LINESTRINGs for each individual and, optionally, combination of columns listed in splitBy. Relocation data should be in two columns representing the X and Y coordinates.

Usage

build_lines(
  DT = NULL,
  projection = NULL,
  id = NULL,
  coords = NULL,
  sortBy = NULL,
  splitBy = NULL
)

Arguments

DT input data.table
projection numeric or character defining the coordinate reference system to be passed to sf::st_crs. For example, either projection = "EPSG:32736" or projection = 32736.
id Character string of ID column name
coords Character vector of X coordinate and Y coordinate column names
build_lines

sortBy   Character string of date time column(s) to sort rows by. Must be a POSIXct.
splitBy  (optional) character string or vector of grouping column name(s) upon which the grouping will be calculated

Details

R-spatial evolution:
Please note, spatsoc has followed updates from R spatial, GDAL and PROJ for handling projections, see more at https://r-spatial.org/r/2020/03/17/wkt.html.
In addition, build_lines previously used sp::SpatialLines but has been updated to use sf::st_as_sf and sf::st_linestring according to the R-spatial evolution, see more at https://r-spatial.org/r/2022/04/12/evolution.html.

Notes on arguments:
The projection argument expects a numeric or character defining the coordinate reference system. For example, for UTM zone 36N (EPSG 32736), the projection argument is either projection = 'EPSG:32736' or projection = 32736. See details in sf::st_crs() and https://spatialreference.org for a list of EPSG codes.
The sortBy argument is used to order the input DT when creating sf LINESTRINGs. It must a column in the input DT of type POSIXct to ensure the rows are sorted by date time.
The splitBy argument offers further control building LINESTRINGs. If in your input DT, you have multiple temporal groups (e.g.: years) for example, you can provide the name of the column which identifies them and build LINESTRINGs for each individual in each year.

build_lines is used by group_lines for grouping overlapping lines generated from relocations.

Value

build_lines returns an sf LINESTRING object with a line for each individual (and optionally splitBy combination).
Individuals (or combinations of individuals and splitBy) with less than two relocations are dropped since it requires at least two relocations to build a line.

See Also

group_lines
Other Build functions: build_polys()

Examples

# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = 'UTC')]
# EPSG code for example data
utm <- 32736

# Build lines for each individual
lines <- build_lines(DT, projection = utm, id = 'ID', coords = c('X', 'Y'),
                      sortBy = 'datetime')

# Build lines for each individual by year
DT[, yr := year(datetime)]
lines <- build_lines(DT, projection = utm, id = 'ID', coords = c('X', 'Y'),
                      sortBy = 'datetime', splitBy = 'yr')

---

**build_polys**  
*Build Polygons*

**Description**

*build_polys* generates a simple feature collection with POLYGONs from a *data.table*. The function accepts a *data.table* with relocation data, individual identifiers, a projection, home range type and parameters. The relocation data is transformed into POLYGONs using either *adehabitatHR::mcp* or *adehabitatHR::kernelUD* for each individual and, optionally, combination of columns listed in *splitBy*. Relocation data should be in two columns representing the X and Y coordinates.

**Usage**

```r
build_polys(
  DT = NULL,
  projection = NULL,
  hrType = NULL,
  hrParams = NULL,
  id = NULL,
  coords = NULL,
  splitBy = NULL,
  spPts = NULL)
```

**Arguments**

- **DT**
  - input *data.table*

- **projection**
  - numeric or character defining the coordinate reference system to be passed to *sf::st_crs*. For example, either `projection = "EPSG:32736"` or `projection = 32736`.

- **hrType**
  - type of HR estimation, either 'mcp' or 'kernel'

- **hrParams**
  - a named list of parameters for *adehabitatHR* functions

- **id**
  - Character string of ID column name
```r
build_polys

coords Character vector of X coordinate and Y coordinate column names
splitBy (optional) character string or vector of grouping column name(s) upon which the grouping will be calculated
spPts alternatively, provide solely a SpatialPointsDataFrame with one column representing the ID of each point, as specified by adehabitatHR::mcp or adehabitatHR::kernelUD

Details

group_polys uses build_polys for grouping overlapping polygons created from relocations.

R-spatial evolution:
Please note, spatsoc has followed updates from R spatial, GDAL and PROJ for handling projections, see more below and details at https://r-spatial.org/r/2020/03/17/wkt.html.
In addition, build_polys previously used sp::SpatialPoints but has been updated to use sf::st_as_sf according to the R-spatial evolution, see more at https://r-spatial.org/r/2022/04/12/evolution.html.

Notes on arguments:
The DT must be a data.table. If your data is a data.frame, you can convert it by reference using data.table::setDT.
The id, coords (and optional splitBy) arguments expect the names of respective columns in DT which correspond to the individual identifier, X and Y coordinates, and additional grouping columns.
The projection argument expects a character string or numeric defining the coordinate reference system to be passed to sf::st_crs. For example, for UTM zone 36S (EPSG 32736), the projection argument is projection = "EPSG:32736" or projection = 32736. See https://spatialreference.org for a list of EPSG codes.
The hrType must be either one of "kernel" or "mcp". The hrParams must be a named list of arguments matching those of adehabitatHR::kernelUD and adehabitatHR::getverticeshr or adehabitatHR::mcp.
The splitBy argument offers further control building POLYGONs. If in your DT, you have multiple temporal groups (e.g.: years) for example, you can provide the name of the column which identifies them and build POLYGONs for each individual in each year.

Value

build_polys returns a simple feature collection with POLYGONs for each individual (and optionally splitBy combination).

An error is returned when hrParams do not match the arguments of the respective hrType adehabitatHR function.

See Also

group_polys
Other Build functions: build_lines()
```
Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = "UTC")]

# EPSG code for example data
utm <- 32736

# Build polygons for each individual using kernelUD and getverticeshr
build_polys(DT, projection = utm, hrType = "kernel",
            hrParams = list(grid = 60, percent = 95),
            id = "ID", coords = c("X", "Y"))

# Build polygons for each individual by year
DT[, yr := year(datetime)]
build_polys(DT, projection = utm, hrType = "mcp",
            hrParams = list(percent = 95),
            id = "ID", coords = c("X", "Y"), splitBy = "yr")
```

DT Movement of 10 "Newfoundland Bog Cows"

Description

A dataset containing the GPS relocations of 10 individuals in winter 2016-2017.

Format

A data.table with 14297 rows and 5 variables:

- **ID** individual identifier
- **X** X coordinate of the relocation (UTM 36N)
- **Y** Y coordinate of the relocation (UTM 36N)
- **datetime** character string representing the date time
- **population** sub population within the individuals

Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))
```
## dyad_id

**Dyad ID**

### Description

Generate a dyad ID for edge list generated by `edge_nn` or `edge_dist`.

### Usage

```r
dyad_id(DT = NULL, id1 = NULL, id2 = NULL)
```

### Arguments

- **DT**: input `data.table` with columns `id1` and `id2`, as generated by `edge_dist` or `edge_nn`
- **id1**: ID1 column name generated by `edge_dist` or `edge_nn`
- **id2**: ID2 column name generated by `edge_dist` or `edge_nn`

### Details

An undirected edge identifier between, for example individuals A and B will be A-B (and reverse B and A will be A-B). Internally sorts and pastes id columns.

More details in the edge and dyad vignette (in progress).

### Value

`dyad_id` returns the input `data.table` with appended "dyadID" column

### Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = 'UTC')]}

# Temporal grouping
group_times(DT, datetime = 'datetime', threshold = '20 minutes')

# Edge list generation
edges <- edge_dist(
    DT,
    threshold = 100,
    id = 'ID',
    coords = c('X', 'Y'),
```
edge_dist

```r

timegroup = 'timegroup',
returnDist = TRUE,
fillNA = TRUE
)

# Generate dyad IDs
dyad_id(edges, 'ID1', 'ID2')
```

---

**Description**

`edge_dist` returns edge lists defined by a spatial distance within the user defined threshold. The function accepts a `data.table` with relocation data, individual identifiers and a threshold argument. The threshold argument is used to specify the criteria for distance between points which defines a group. Relocation data should be in two columns representing the X and Y coordinates.

**Usage**

```r
edge_dist(
  DT = NULL,
  threshold,
  id = NULL,
  coords = NULL,
  timegroup,
  splitBy = NULL,
  returnDist = FALSE,
  fillNA = TRUE
)
```

**Arguments**

- **DT**
  - input data.table
- **threshold**
  - distance for grouping points, in the units of the coordinates
- **id**
  - Character string of ID column name
- **coords**
  - Character vector of X coordinate and Y coordinate column names
- **timegroup**
  - timegroup field in the DT within which the grouping will be calculated
- **splitBy**
  - (optional) character string or vector of grouping column name(s) upon which the grouping will be calculated
- **returnDist**
  - boolean indicating if the distance between individuals should be returned. If FALSE (default), only ID1, ID2 columns (and timegroup, splitBy columns if provided) are returned. If TRUE, another column "distance" is returned indicating the distance between ID1 and ID2.
- **fillNA**
  - boolean indicating if NAs should be returned for individuals that were not within the threshold distance of any other. If TRUE, NAs are returned. If FALSE, only edges between individuals within the threshold distance are returned.
Details

The DT must be a data.table. If your data is a data.frame, you can convert it by reference using `data.table::setDT`.

The id, coords timegroup (and optional splitBy) arguments expect the names of a column in DT which correspond to the individual identifier, X and Y coordinates, timegroup (generated by group_times) and additional grouping columns.

If provided, the threshold must be provided in the units of the coordinates and must be larger than 0. If the threshold is NULL, the distance to all other individuals will be returned. The coordinates must be planar coordinates (e.g.: UTM). In the case of UTM, a threshold = 50 would indicate a 50m distance threshold.

The timegroup argument is required to define the temporal groups within which edges are calculated. The intended framework is to group rows temporally with `group_times` then spatially with `edge_dist`. If you have already calculated temporal groups without `group_times`, you can pass this column to the timegroup argument. Note that the expectation is that each individual will be observed only once per timegroup. Caution that accidentally including huge numbers of rows within timegroups can overload your machine since all pairwise distances are calculated within each timegroup.

The splitBy argument offers further control over grouping. If within your DT, you have multiple populations, subgroups or other distinct parts, you can provide the name of the column which identifies them to splitBy. `edge_dist` will only consider rows within each splitBy subgroup.

Value

`edge_dist` returns a data.table with columns ID1, ID2, timegroup (if supplied) and any columns provided in splitBy. If 'returnDist' is TRUE, column 'distance' is returned indicating the distance between ID1 and ID2.

The ID1 and ID2 columns represent the edges defined by the spatial (and temporal with `group_times`) thresholds.

See Also

Other Edge-list generation: `edge_nn()`

Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = 'UTC')]

# Temporal grouping
group_times(DT, datetime = 'datetime', threshold = '20 minutes')
```
# Edge list generation

```r
dist <- edge_dist(
  DT,  
  threshold = 100, 
  id = 'ID', 
  coords = c('X', 'Y'), 
  timegroup = 'timegroup', 
  returnDist = TRUE, 
  fillNA = TRUE
)
```

## edge_nn

**Nearest neighbour based edge lists**

**Description**

`edge_nn` returns edge lists defined by the nearest neighbour. The function accepts a `data.table` with relocation data, individual identifiers and a threshold argument. The threshold argument is used to specify the criteria for distance between points which defines a group. Relocation data should be in two columns representing the X and Y coordinates.

**Usage**

```r
dist <- edge_nn(
  DT = NULL, 
  id = NULL, 
  coords = NULL, 
  timegroup, 
  splitBy = NULL, 
  threshold = NULL, 
  returnDist = FALSE
)
```

**Arguments**

- **DT** 
  input `data.table`
- **id** 
  Character string of ID column name
- **coords** 
  Character vector of X coordinate and Y coordinate column names
- **timegroup** 
  timegroup field in the `DT` within which the grouping will be calculated
- **splitBy** 
  (optional) character string or vector of grouping column name(s) upon which the grouping will be calculated
- **threshold** 
  (optional) spatial distance threshold to set maximum distance between an individual and their neighbour.
- **returnDist** 
  boolean indicating if the distance between individuals should be returned. If FALSE (default), only ID, NN columns (and timegroup, splitBy columns if provided) are returned. If TRUE, another column "distance" is returned indicating the distance between ID and NN.
Details

The DT must be a data.table. If your data is a data.frame, you can convert it by reference using `data.table::setDT`.

The id, coords, timegroup (and optional splitBy) arguments expect the names of a column in DT which correspond to the individual identifier, X and Y coordinates, timegroup (generated by `group_times`) and additional grouping columns.

The threshold must be provided in the units of the coordinates. The threshold must be larger than 0. The coordinates must be planar coordinates (e.g.: UTM). In the case of UTM, a threshold = 50 would indicate a 50m distance threshold.

The `timegroup` argument is required to define the temporal groups within which edge nearest neighbours are calculated. The intended framework is to group rows temporally with `group_times` then spatially with `edge_nn`. If you have already calculated temporal groups without `group_times`, you can pass this column to the `timegroup` argument. Note that the expectation is that each individual will be observed only once per timegroup. Caution that accidentally including huge numbers of rows within timegroups can overload your machine since all pairwise distances are calculated within each timegroup.

The `splitBy` argument offers further control over grouping. If within your DT, you have multiple populations, subgroups or other distinct parts, you can provide the name of the column which identifies them to `splitBy`. `edge_nn` will only consider rows within each `splitBy` subgroup.

Value

`edge_nn` returns a data.table with three columns: `timegroup`, `ID` and `NN`. If `returnDist` is TRUE, column `distance` is returned indicating the distance between ID and NN.

The ID and NN columns represent the edges defined by the nearest neighbours (and temporal thresholds with `group_times`).

If an individual was alone in a timegroup or `splitBy`, or did not have any neighbours within the threshold distance, they are assigned NA for nearest neighbour.

See Also

Other Edge-list generation: `edge_dist()`

Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Select only individuals A, B, C for this example
DT <- DT[ID %in% c('A', 'B', 'C')]

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = 'UTC')]"
## get_gbi

Generate group by individual matrix

### Description

get_gbi generates a group by individual matrix. The function accepts a data.table with individual identifiers and a group column. The group by individual matrix can then be used to build a network using `asnipe::get_network`.

### Usage

```r
get_gbi(DT = NULL, group = "group", id = NULL)
```

### Arguments

- **DT**: input data.table
- **group**: Character string of group column (generated from one of spatsoc's spatial grouping functions)
- **id**: Character string of ID column name

### Details

The DT must be a data.table. If your data is a data.frame, you can convert it by reference using `data.table::setDT`.

The group argument expects the name of a column which corresponds to an integer group identifier (generated by spatsoc’s grouping functions).

The id argument expects the name of a column which corresponds to the individual identifier.
Value

get_gbi returns a group by individual matrix (columns represent individuals and rows represent groups).

Note that get_gbi is identical in function for turning the outputs of spatsoc into social networks as 

`asnipe::get_group_by_individual` but is more efficient thanks to `data.table::dcast`.

See Also

- `group_pts`
- `group_lines`
- `group_polys`

Other Social network tools: `randomizations()`

Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = "UTC")]
DT[, yr := year(datetime)]

# EPSG code for example data
utm <- "EPSG:32736"

group_polys(DT, area = FALSE, hrType = "mcp",
hrParams = list(percent = 95),
projection = utm, id = "ID", coords = c("X", "Y"),
splitBy = "yr")

gbiMtrx <- get_gbi(DT = DT, group = "group", id = "ID")
```

Description

group_lines groups rows into spatial groups by generating LINESTRINGs and grouping based on spatial intersection. The function accepts a data.table with relocation data, individual identifiers and a distance threshold. The relocation data is transformed into sf LINESTRINGs using build_lines and intersecting LINESTRINGs are grouped. The threshold argument is used to specify the distance criteria for grouping. Relocation data should be in two columns representing the X and Y coordinates.
Usage

```r
group_lines(
  DT = NULL,
  threshold = NULL,
  projection = NULL,
  id = NULL,
  coords = NULL,
  timegroup = NULL,
  sortBy = NULL,
  splitBy = NULL,
  sfLines = NULL
)
```

Arguments

- **DT**: input data.table
- **threshold**: The width of the buffer around the lines in the units of the projection. Use `threshold = 0` to compare intersection without buffering.
- **projection**: numeric or character defining the coordinate reference system to be passed to `sf::st_crs`. For example, either `projection = "EPSG:32736"` or `projection = 32736`.
- **id**: Character string of ID column name
- **coords**: Character vector of X coordinate and Y coordinate column names
- **timegroup**: timegroup field in the DT within which the grouping will be calculated
- **sortBy**: Character string of date time column(s) to sort rows by. Must be a POSIXct.
- **splitBy**: (optional) character string or vector of grouping column name(s) upon which the grouping will be calculated
- **sfLines**: Alternatively to providing a DT, provide a simple feature LINESTRING object generated with the sf package. The id argument is required to provide the identifier matching each LINESTRING. If an sfLines object is provided, groups cannot be calculated by timegroup or splitBy.

Details

**R-spatial evolution:**

Please note, spatsoc has followed updates from R spatial, GDAL and PROJ for handling projections, see more at [https://r-spatial.org/r/2020/03/17/wkt.html](https://r-spatial.org/r/2020/03/17/wkt.html).

In addition, `group_lines` (and `build_lines`) previously used `sp::SpatialLines`, `rgeos::gIntersects`, `rgeos::gBuffer` but have been updated to use `sf::st_as_sf`, `sf::st_linestring`, `sf::st_intersects`, and `sf::st_buffer` according to the R-spatial evolution, see more at [https://r-spatial.org/r/2022/04/12/evolution.html](https://r-spatial.org/r/2022/04/12/evolution.html).

**Notes on arguments:**

The DT must be a `data.table`. If your data is a `data.frame`, you can convert it by reference using `data.table::setDT`.
The `id`, `coords`, `sortBy` (and optional `timegroup` and `splitBy`) arguments expect the names of respective columns in `DT` which correspond to the individual identifier, X and Y coordinates, sorting, timegroup (generated by `group_times`) and additional grouping columns.

The projection argument expects a numeric or character defining the coordinate reference system. For example, for UTM zone 36N (EPSG 32736), the projection argument is either `projection = 'EPSG:32736'` or `projection = 32736`. See details in `sf::st_crs()` and [https://spatialreference.org](https://spatialreference.org) for a list of EPSG codes.

The `sortBy` argument is used to order the input `DT` when creating sf LINESTRINGS. It must a column in the input `DT` of type POSIXct to ensure the rows are sorted by date time.

The threshold must be provided in the units of the coordinates. The threshold can be equal to 0 if strict overlap is intended, otherwise it should be some value greater than 0. The coordinates must be planar coordinates (e.g.: UTM). In the case of UTM, a threshold = 50 would indicate a 50m distance threshold.

The `timegroup` argument is optional, but recommended to pair with `group_times`. The intended framework is to group rows temporally with `group_times` then spatially with `group_lines` (or `group_pts`, `group_polys`). With `group_lines`, pick a relevant `group_times` threshold such as '1 day' or '7 days' which is informed by your study species, system or question.

The `splitBy` argument offers further control building LINESTRINGS. If in your input `DT`, you have multiple temporal groups (e.g.: years) for example, you can provide the name of the column which identifies them and build LINESTRINGS for each individual in each year. The grouping performed by `group_lines` will only consider rows within each `splitBy` subgroup.

**Value**

`group_lines` returns the input `DT` appended with a "group" column.

This column represents the spatial (and if `timegroup` was provided - spatiotemporal) group calculated by intersecting lines. As with the other grouping functions, the actual value of `group` is arbitrary and represents the identity of a given group where 1 or more individuals are assigned to a group. If the data was reordered, the group may change, but the contents of each group would not.

A message is returned when a column named "group" already exists in the input `DT`, because it will be overwritten.

**See Also**

`build_lines` `group_times`

Other Spatial grouping: `group_polys()`, `group_pts()`

**Examples**

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Subset only individuals A, B, and C
DT <- DT[ID %in% c('A', 'B', 'C')]
```
# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = 'UTC')]

# EPSG code for example data
utm <- 32736

group_lines(DT, threshold = 50, projection = utm, sortBy = 'datetime',
           id = 'ID', coords = c('X', 'Y'))

## Daily movement tracks
# Temporal grouping
group_times(DT, datetime = 'datetime', threshold = '1 day')

# Subset only first 50 days
DT <- DT[timegroup < 25]

# Spatial grouping
group_lines(DT, threshold = 50, projection = utm,
           id = 'ID', coords = c('X', 'Y'),
           timegroup = 'timegroup', sortBy = 'datetime')

## Daily movement tracks by population
group_lines(DT, threshold = 50, projection = utm,
           id = 'ID', coords = c('X', 'Y'),
           timegroup = 'timegroup', sortBy = 'datetime',
           splitBy = 'population')

---

**group_polys**

**Group Polygons**

**Description**

`group_polys` groups rows into spatial groups by overlapping polygons (home ranges). The function accepts a `data.table` with relocation data, individual identifiers and an `area` argument. The relocation data is transformed into home range POLYGONs using `build_polys()` with `adehabitatHR::mcp` or `adehabitatHR::kernelUD`. If the `area` argument is FALSE, `group_polys` returns grouping calculated by spatial overlap. If the `area` argument is TRUE, `group_polys` returns the area area and proportion of overlap. Relocation data should be in two columns representing the X and Y coordinates.

**Usage**

group_polys(
    DT = NULL,
    area = NULL,
    hrType = NULL,
    hrParams = NULL,
    projection = NULL,)
Arguments

DT          input data.table
area        boolean indicating either overlap group (when FALSE) or area and proportion of
            overlap (when TRUE)
hrType      type of HR estimation, either 'mcp' or 'kernel'
hrParams    a named list of parameters for adehabitatHR functions
projection   numeric or character defining the coordinate reference system to be passed to
              sf::st_crs. For example, either projection = "EPSG:32736" or projection = 32736.
id          Character string of ID column name
coords      Character vector of X coordinate and Y coordinate column names
splitBy     (optional) character string or vector of grouping column name(s) upon which
            the grouping will be calculated
sfPolys     Alternatively, provide solely a simple features object with POLYGONs or MULTIPOLYGONs. If sfPolys are provided, id is required and splitBy cannot be used.

Details

R-spatial evolution:

Please note, spatsoc has followed updates from R spatial, GDAL and PROJ for handling projections, see more below and details at https://r-spatial.org/r/2020/03/17/wkt.html.

In addition, group_polys previously used rgeos::gIntersection, rgeos::gIntersects and rgeos::gArea but has been updated to use sf::st_intersects, sf::st_intersection and sf::st_area according to the R-spatial evolution, see more at https://r-spatial.org/r/2022/04/12/evolution.html.

Notes on arguments:

The DT must be a data.table. If your data is a data.frame, you can convert it by reference using data.table::setDT().

The id, coords (and optional splitBy) arguments expect the names of respective columns in DT which correspond to the individual identifier, X and Y coordinates, and additional grouping columns.

The projection argument expects a character string or numeric defining the coordinate reference system to be passed to sf::st_crs. For example, for UTM zone 36S (EPSG 32736), the projection argument is projection = "EPSG:32736" or projection = 32736. See https://spatialreference.org for a list of EPSG codes.

The hrType must be either one of "kernel" or "mcp". The hrParams must be a named list of arguments matching those of adehabitatHR::kernelUD() or adehabitatHR::mcp().
The `splitBy` argument offers further control over grouping. If within your DT, you have multiple populations, subgroups or other distinct parts, you can provide the name of the column which identifies them to `splitBy`. The grouping performed by `group_polys` will only consider rows within each `splitBy` subgroup.

**Value**

When `area` is `FALSE`, `group_polys` returns the input DT appended with a group column. As with the other grouping functions, the actual value of `group` is arbitrary and represents the identity of a given group where 1 or more individuals are assigned to a group. If the data was reordered, the group may change, but the contents of each group would not. When `area` is `TRUE`, `group_polys` returns a proportional area overlap data.table. In this case, `ID` refers to the focal individual of which the total area is compared against the overlapping area of `ID2`.

If `area` is `FALSE`, a message is returned when a column named `group` already exists in the input DT, because it will be overwritten.

Along with changes to follow the R-spatial evolution, `group_polys` also now returns area and proportion of overlap with units explicitly specified through the `units` package.

**See Also**

`build_polys()` `group_times()`

Other Spatial grouping: `group_lines()`, `group_pts()`

**Examples**

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = "UTC")]

# EPSG code for example data
utm <- 32736

group_polys(DT, area = FALSE, hrType = "mcp",
            hrParams = list(percent = 95), projection = utm,
            id = "ID", coords = c("X", "Y"))

areaDT <- group_polys(DT, area = TRUE, hrType = "mcp",
                      hrParams = list(percent = 95), projection = utm,
                      id = "ID", coords = c("X", "Y"))

print(areaDT)
```
**group_pts**  

**Group Points**

**Description**

`group_pts` groups rows into spatial groups. The function accepts a `data.table` with relocation data, individual identifiers and a threshold argument. The threshold argument is used to specify the criteria for distance between points which defines a group. Relocation data should be in two columns representing the X and Y coordinates.

**Usage**

```r
group_pts(
  DT = NULL,
  threshold = NULL,
  id = NULL,
  coords = NULL,
  timegroup,
  splitBy = NULL
)
```

**Arguments**

- `DT` input `data.table`
- `threshold` distance for grouping points, in the units of the coordinates
- `id` Character string of ID column name
- `coords` Character vector of X coordinate and Y coordinate column names
- `timegroup` timegroup field in the `DT` within which the grouping will be calculated
- `splitBy` (optional) character string or vector of grouping column name(s) upon which the grouping will be calculated

**Details**

The `DT` must be a `data.table`. If your data is a `data.frame`, you can convert it by reference using `data.table::setDT`.

The `id`, `coords`, `timegroup` (and optional `splitBy`) arguments expect the names of a column in `DT` which correspond to the individual identifier, X and Y coordinates, timegroup (typically generated by `group_times`) and additional grouping columns.

The `threshold` must be provided in the units of the coordinates. The `threshold` must be larger than 0. The coordinates must be planar coordinates (e.g.: UTM). In the case of UTM, a `threshold` = 50 would indicate a 50m distance threshold.

The `timegroup` argument is required to define the temporal groups within which spatial groups are calculated. The intended framework is to group rows temporally with `group_times` then spatially with `group_pts` (or `group_lines`, `group_polys`). If you have already calculated temporal groups...
without `group_times`, you can pass this column to the `timegroup` argument. Note that the expectation is that each individual will be observed only once per `timegroup`. Caution that accidentally including huge numbers of rows within `timegroups` can overload your machine since all pairwise distances are calculated within each `timegroup`.

The `splitBy` argument offers further control over grouping. If within your `DT`, you have multiple populations, subgroups or other distinct parts, you can provide the name of the column which identifies them to `splitBy`. The grouping performed by `group_pts` will only consider rows within each `splitBy` subgroup.

### Value

`group_pts` returns the input `DT` appended with a group column.

This column represents the spatial-temporal group. As with the other grouping functions, the actual value of `group` is arbitrary and represents the identity of a given group where 1 or more individuals are assigned to a group. If the data was reordered, the `group` may change, but the contents of each group would not.

A message is returned when a column named `group` already exists in the input `DT`, because it will be overwritten.

### See Also

- `group_times`
- Other Spatial grouping: `group_lines()`, `group_polys()`

### Examples

```r
# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Select only individuals A, B, C for this example
DT <- DT[ID %in% c('A', 'B', 'C')]

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = 'UTC')]

# Temporal grouping
group_times(DT, datetime = 'datetime', threshold = '20 minutes')

# Spatial grouping with timegroup
group_pts(DT, threshold = 5, id = 'ID',
         coords = c('X', 'Y'), timegroup = 'timegroup')

# Spatial grouping with timegroup and splitBy on population
group_pts(DT, threshold = 5, id = 'ID', coords = c('X', 'Y'),
         timegroup = 'timegroup', splitBy = 'population')
```
**group_times**

**Group Times**

**Description**

`group_times` groups rows into time groups. The function accepts date time formatted data and a threshold argument. The threshold argument is used to specify a time window within which rows are grouped.

**Usage**

```r
group_times(DT = NULL, datetime = NULL, threshold = NULL)
```

**Arguments**

- **DT**: input data.table
- **datetime**: name of date time column(s). Either 1 POSIXct or 2 IDate and ITime. e.g.: 'datetime' or c('idate', 'itime')
- **threshold**: threshold for grouping times. e.g.: '2 hours', '10 minutes', etc. If not provided, times will be matched exactly. Note that provided threshold must be in the expected format: '## unit'

**Details**

The `DT` must be a data.table. If your data is a data.frame, you can convert it by reference using `data.table::setDT`.

The `datetime` argument expects the name of a column in `DT` which is of type POSIXct or the name of two columns in `DT` which are of type IDate and ITime.

The `threshold` must be provided in units of minutes, hours or days. The character string should start with an integer followed by a unit, separated by a space. It is interpreted in terms of 24 hours which poses the following limitations:

- minutes, hours and days cannot be fractional
- minutes must divide evenly into 60
- minutes must not exceed 60
- minutes, hours which are nearer to the next day, are grouped as such
- hours must divide evenly into 24
- multi-day blocks should divide into the range of days, else the blocks may not be the same length

In addition, the `threshold` is considered a fixed window throughout the time series and the rows are grouped to the nearest interval.

If `threshold` is NULL, rows are grouped using the `datetime` column directly.
Value

group_times returns the input DT appended with a timegroup column and additional temporal grouping columns to help investigate, troubleshoot and interpret the timegroup.

The actual value of timegroup is arbitrary and represents the identity of a given timegroup which 1 or more individuals are assigned to. If the data was reordered, the group may change, but the contents of each group would not.

The temporal grouping columns added depend on the threshold provided:

- threshold with unit minutes: "minutes" column added identifying the nearest minute group for each row.
- threshold with unit hours: "hours" column added identifying the nearest hour group for each row.
- threshold with unit days: "block" columns added identifying the multiday block for each row.

A message is returned when any of these columns already exist in the input DT, because they will be overwritten.

See Also

group_pts group_lines group_polys

Examples

# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Cast the character column to POSIXct
DT[, datetime := as.POSIXct(datetime, tz = "UTC")]

group_times(DT, datetime = "datetime", threshold = "5 minutes")
group_times(DT, datetime = "datetime", threshold = "2 hours")
group_times(DT, datetime = "datetime", threshold = "10 days")
randomizations

Description

randomizations performs data-stream social network randomization. The function accepts a data.table with relocation data, individual identifiers and a randomization type. The data.table is randomized either using step or daily between-individual methods, or within-individual daily trajectory method described by Spiegel et al. (2016).

Usage

randomizations(
  DT = NULL,
  type = NULL,
  id = NULL,
  group = NULL,
  coords = NULL,
  datetime = NULL,
  splitBy = NULL,
  iterations = NULL
)

Arguments

DT input data.table

 type one of 'daily', 'step' or 'trajectory' - see details
 id Character string of ID column name
 group generated from spatial grouping functions - see details
 coords Character vector of X coordinate and Y coordinate column names
 datetime field used for providing date time or time group - see details
 splitBy List of fields in DT to split the randomization process by
 iterations The number of iterations to randomize

Details

The DT must be a data.table. If your data is a data.frame, you can convert it by reference using data.table::setDT.

Three randomization types are provided:

1. step - randomizes identities of relocations between individuals within each time step.
2. daily - randomizes identities of relocations between individuals within each day.
3. trajectory - randomizes daily trajectories within individuals (Spiegel et al. 2016).

Depending on the type, the datetime must be a certain format:

- step - datetime is integer group created by group_times
- daily - datetime is POSIXct format
- trajectory - datetime is POSIXct format
The id, datetime, (and optional splitBy) arguments expect the names of respective columns in DT which correspond to the individual identifier, date time, and additional grouping columns. The coords argument is only required when the type is "trajectory", since the coordinates are required for recalculating spatial groups with group_pts, group_lines or group_polys.

Please note that if the data extends over multiple years, a column indicating the year should be provided to the splitBy argument. This will ensure randomizations only occur within each year.

The group argument is expected only when type is 'step' or 'daily'.

For example, using data.table::year:

```r
DT[, yr := year(datetime)] randomizations(DT, type = 'step', id = 'ID', datetime = 'timegroup', splitBy = 'yr')
```

iterations is set to 1 if not provided. Take caution with a large value for iterations with large input DT.

Value

randomizations returns the random date time or random id along with the original DT, depending on the randomization type. The length of the returned data.table is the original number of rows multiplied by the number of iterations + 1. For example, 3 iterations will return 4x - one observed and three randomized.

Two columns are always returned:

- observed - if the rows represent the observed (TRUE/FALSE)
- iteration - iteration of rows (where 0 is the observed)

In addition, depending on the randomization type, random ID or random date time columns are returned:

- step - randomID each time step
- daily - randomID for each day and jul indicating julian day
- trajectory - a random date time ("random" prefixed to datetime argument), observed jul and randomJul indicating the random day relocations are swapped to.

References

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See Also

Other Social network tools: get_gbi()
Examples

# Load data.table
library(data.table)

# Read example data
DT <- fread(system.file("extdata", "DT.csv", package = "spatsoc"))

# Select only individuals A, B, C for this example
DT <- DT[ID %in% c('A', 'B', 'C')]

# Date time columns
DT[, datetime := as.POSIXct(datetime)]
DT[, yr := year(datetime)]

# Temporal grouping
group_times(DT, datetime = 'datetime', threshold = '5 minutes')

# Spatial grouping with timegroup
group_pts(DT, threshold = 5, id = 'ID', coords = c('X', 'Y'), timegroup = 'timegroup')

# Randomization: step
randStep <- randomizations(
  DT,
  type = 'step',
  id = 'ID',
  group = 'group',
  datetime = 'timegroup',
  splitBy = 'yr',
  iterations = 2
)

# Randomization: daily
randDaily <- randomizations(
  DT,
  type = 'daily',
  id = 'ID',
  group = 'group',
  datetime = 'datetime',
  splitBy = 'yr',
  iterations = 2
)

# Randomization: trajectory
randTraj <- randomizations(
  DT,
  type = 'trajectory',
  id = 'ID',
  group = NULL,
  coords = c('X', 'Y'),
  datetime = 'datetime',
  splitBy = 'yr',
  iterations = 2
)
randomizations

iterations = 2
)

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