

Package ‘wpeR’

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Type Package

Title Streamlined Analysis of Wild Pedigree Data

Version 0.1.0

Description Analyzing pedigree data of wild populations. While primarily designed to process outputs from the 'COLONY' (Jones & Wang (2010) <[doi:10.1111/j.1755-0998.2009.02787.x](https://doi.org/10.1111/j.1755-0998.2009.02787.x)>) pedigree reconstruction software, it can also accommodate data from other sources. By linking reconstructed pedigrees with genetic sample metadata, 'wpeR' produces spatial and temporal visualizations as well as tabular summaries that support interpretation of family structures and dynamics. The main goal of the package is to provide a solution for the analysis of complex wild pedigree data and to help the user to gain insights into genetic relationships within wild animal populations.

License GPL (>= 3)

URL <https://gr3602.github.io/wpeR/>

BugReports <https://github.com/GR3602/wpeR/issues>

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anim_timespan	<i>Get dates of individuals first and last sample</i>
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Description

Takes data frame of all samples and returns the dates of individuals first and last sample. Besides that the functions determines if animal is dead based on predefined sample type eg. tissue.

Usage

```
anim_timespan(individual_id, sample_date, sample_type, dead = "Tissue")
```

Arguments

individual_id	Column in the dataframe of all samples containing individual animal identifier code. Defined as dataframe\$column.
sample_date	Column in the dataframe of all samples containing the date of sample collection. Must be in Date format. Defined as dataframe\$column.
sample_type	Column in the dataframe of all samples containing the data on the type (eg. scat, tissue, saliva) of particular sample. Defined as dataframe\$column.
dead	Single value or vector of different lethal sample types. If no lethal samples are included in the sampledata the dead parameter can be set to FALSE (dead = FALSE). Defaults to "Tissue".

Value

A data frame with four columns and one row for each `individual_id`. Returned data frame columns correspond to individual identification key (ID), date of first (`FirstSeen`) and last (`LastSeen`) sample of individual and logical (TRUE/FALSE) value that identifies if the individual is dead (`IsDead`).

Examples

```
anim_timespan(
  individual_id = wolf_samples$AnimalRef,
  sample_date = wolf_samples$Date,
  sample_type = wolf_samples$SType,
  dead = c("Tissue")
)
```

check_sampledata	<i>Check and prepare genetic sample metadata</i>
------------------	--------------------------------------------------

Description

Verifies the consistency of columns in the genetic sample metadata and prepares it for use with other functions in the `wpeR` package. The function ensures that the provided data is properly formatted and conforms to the standards of functions that make up the `wpeR` package.

Usage

```
check_sampledata(
  Sample,
  Date,
  AnimalRef,
  GeneticSex,
  lat,
  lng,
  SType,
  extraCols = NULL
)
```

Arguments

Sample	A vector of sample unique identifier codes.
Date	A vector of sample collection dates in 'YYYY-MM-DD' format.
AnimalRef	A vector of identifier codes of the particular individual that the sample belongs to.
GeneticSex	A vector of genetic sex information ('F' for female, 'M' for male, NA for unknown).
lat	A vector of latitude coordinates in the WGS84 coordinate system (EPSG: 4326).

lng	A vector of longitude coordinates in the WGS84 coordinate system (EPSG: 4326).
SType	A vector of sample types eg.: scat, hair, tissue.
extraCols	A vector of extra column names that the user wants to include in sampledata data frame (see Details).

Details

By specifying the `extraCols` parameter additional information can be included in the `sampledata` dataframe. Such additional information is not required for the functioning of the `wpeR` package functions, but can be useful to the user when interpreting results. When including additional columns the function inputs (`Sample`, `Date`, `AnimalRef`...) have to be defined as a vector extracted from data frame column (eg. `Sample = dataframe$column`) and the `extraCols` parameter is defined, as a vector of column names from the same data frame (eg. `extraCols = c(column1, column2, column3)`).

Value

A data frame with at least 7 columns and a number of rows equal to the length of the input vector. Each column corresponds to one of the input parameters. If the function executes without warnings or errors, the result from `check_sampledata()` can be used as an input parameter for other functions within this package: `get_colony()`, `get_ped()`, `org_fams()` and `plot_table()`.

Examples

```
sampledata <- check_sampledata(
  Sample = wolf_samples$Sample,
  Date = wolf_samples$Date,
  AnimalRef = wolf_samples$AnimalRef,
  GeneticSex = wolf_samples$GeneticSex,
  lat = wolf_samples$lat,
  lng = wolf_samples$lng,
  SType = wolf_samples$SType
)
```

dyn_matrix

Get matrix of apparent survival

Description

Creates a matrix that shows number of captured animals between multiple seasons.

Usage

```
dyn_matrix(animal_id, capture_date, start_dates, end_dates)
```

Arguments

animal_id	A column in the dataframe of all samples that stores individual animal identifier code.
capture_date	A column in the dataframe of all samples that stores the date of sample collection. Must be in Date format.
start_dates	Vector of dates in Date format that define the start of each season.
end_dates	Vector of dates in Date format that define the end of each season.

Value

A matrix with 1 + no. seasons rows and columns.

- diagonal: number of new captures in each session,
- above diagonal: number of recaptures from season x to season y,
- below diagonal: number of animals from season y that skipped season x.

Season x is defined in first row, season y in first column. Column Tot. Capts gives all detected individuals in season y. Row Tot. Skipped gives all individuals skipped in season x but detected later.

Examples

```
# Define start and end dates for sampling seasons.
seasons <- data.frame(
  start = c(
    as.Date("2017-01-01"),
    as.Date("2018-01-01"),
    as.Date("2019-01-01")
  ),
  end = c(
    as.Date("2017-12-31"),
    as.Date("2018-12-31"),
    as.Date("2019-12-31")
  )
)

# Create a dynamics matrix for animal captures.
dyn_matrix(
  animal_id = wolf_samples$AnimalRef,
  capture_date = wolf_samples$Date,
  start_dates = seasons$start,
  end_dates = seasons$end
)
```

get_colony

*Organizes COLONY output***Description**

Extends BestConfig_Ordered output from **COLONY** pedigree reconstruction software with additional data about individuals included in pedigree. The function adds missing parents to OffspringID, assigns sex to each individual included in OffspringID and adds the computed probabilities of paternity and maternity assignments (probability of assignments is visible only if the out parameter is set to "table"). The function also prepares data so that the output of the function can be directly analyzed with **kinship2**, **pedtools** or **FamAgg** packages.

Usage

```
get_colony(
  colony_project_path,
  sampledata,
  rm_obsolete_parents = TRUE,
  out = "FamAgg"
)
```

Arguments

colony_project_path	Character string. Path to the folder where COLONY output files are saved. Has to include file path and project name (see Details).
sampledata	Data frame. Metadata for all genetic samples that belong to the individuals included in pedigree reconstruction analysis. This data frame should adhere to the formatting and naming conventions outlined in the check_sampledata() documentation.
rm_obsolete_parents	Logical. Should unknown parents be removed from output. Applies just to offspring for which both parents are unknown. Defaults to TRUE.
out	Character string. For use with which package should the output be formatted? kinship2 (out = "kinship2"), pedtools (out = "pedtools"), FamAgg (out = "FamAgg") or the created data.frame can be outputted as is (out = "table"). Defaults to "FamAgg"

Details

COLONY output tables needed for this function (`.BestConfig_Ordered`, `.Maternity` and `.Paternity`) are read directly from the colony output folder and do not need to be imported into R session. The path to the outputs is defined with `colony_project_path` parameter. When defining `colony_project_path` the user needs to define a complete path to the directory where colony outputs are stored and also the file name (file name of COLONY outputs equals the project name eg. `./path/to/the/COLONY/output/folder/COLONY_project_name`).

Value

A data frame describing a common pedigree structure. Each individual included in pedigree represents one row. Columns describe individual identifier code, identifier code for mother and father, sex and family of individual. Column names and arrangement depends on selected output (out parameter).

Examples

```
# Define the path to COLONY output
path <- paste0(system.file("extdata", package = "wpeR"), "/wpeR_samplePed")

# Get pedigree data in FamAgg format
get_colony(
  colony_project_path = path,
  sampledata = wolf_samples
)
```

get_ped

Organizes pedigree data

Description

Offers an alternative to [get_colony\(\)](#) function in cases where the pedigree was not reconstructed with **COLONY** software. It takes a pedigree dataframe and assigns sex to each individual. The function also prepares data so that the output of the function can be directly analyzed with [kinship2](#), [pedtools](#) or [FamAgg](#) packages.

Usage

```
get_ped(ped, sampledata, out = "FamAgg")
```

Arguments

ped	Data frame. Pedigree data frame with the most basic structure. Three columns corresponding to offspring, father and mother (see Details). Unknown parents should be represented by NA values.
sampledata	Data frame. Metadata for all genetic samples that belong to the individuals included in pedigree reconstruction analysis. This data frame should adhere to the formatting and naming conventions outlined in the check_sampledata() documentation.
out	Character string. For use with which package should the output be formatted? kinship2 (out = "kinship2"), pedtools (out = "pedtools") or FamAgg (out = "FamAgg") or the created data.frame can be outputted as is (out = "table"). Defaults to "FamAgg"

Details

The custom pedigree specified through the `ped` parameter should mirror the structure of a COLONY pedigree and share the same column names. It should consist of three columns for each offspring: OffspringID, FatherID, MotherID. When considering unknown parents they should be represented by NA values.

Value

A data frame describing a common pedigree structure. Each individual included in pedigree represents one row. Columns describe individual identifier code, identifier code for mother and father and sex of individual. Column names and arrangement depends on selected output (`out` parameter).

Examples

```
#example pedigree dataframe
ped <- data.frame(
  OffspringID = c(
    "M273P", "M20AM", "M2757", "M2ALK", "M2ETE", "M2EUJ", "MSV00E",
    "MSV018", "MSV05L", "MSV0M6", "MSV0T4", "MSV0T7", "MSV0TJ", "MSV0UL"
  ),
  FatherID = c(
    NA, NA, "M20AM", "M20AM", "M20AM", "M20AM", "M20AM",
    "M20AM", "M20AM", "M20AM", "M20AM", "M20AM", "M20AM"
  ),
  MotherID = c(
    NA, NA, "M273P", "M273P", "M273P", "M273P", "M273P",
    "M273P", "M273P", "M273P", "M273P", "M273P", "M273P"
  )
)
#Get pedigree data in FamAgg format
get_ped(
  ped = ped,
  sampledata = wolf_samples
)
```

nbtw_seasons

Number of detected animals between two sampling seasons

Description

Gives an numeric overview of individuals captured within the second sampling season compared to the first one.

Usage

```
nbtw_seasons(
  animal_id,
  capture_date,
```



```

    season1_start,
    season1_end,
    season2_start,
    season2_end
  )

```

Arguments

animal_id	A column in the dataframe of all samples that stores individual animal identifier code.
capture_date	A column in the dataframe of all samples that stores the date of sample collection. Must be in Date format.
season1_start	String in Date format. Start of first capture season. Start and end date are included in the capture season.
season1_end	String in Date format. End of first capture season. Start and end date are included in the capture season.
season2_start	String in Date format. Start of second capture season. Start and end date are included in the capture season.
season2_end	String in Date format. End of second capture season. Start and end date are included in the capture season.

Value

A data frame with one row and six columns corresponding to season 1 and 2 start and end dates, number of detected animals in season 2 (`total_cap`), number of new detentions in season 2 (`new_captures`), number of animals from season 1 detected within season 2 (recaptured) and number of individuals skipped in season 2 but detected after the end of that season (`skipped`).

Examples

```

# Calculate the number of animals detected between two sampling seasons.
nbtw_seasons(
  animal_id = wolf_samples$AnimalRef,
  capture_date = wolf_samples$Date,
  season1_start = as.Date("2017-01-01"),
  season1_end = as.Date("2017-12-31"),
  season2_start = as.Date("2018-01-01"),
  season2_end = as.Date("2018-12-31")
)

```

 org_fams

Organize animals into families and expand pedigree data

Description

Takes pedigree data from `get_colony()` or `get_ped()` function and groups animals into families. It also expands the pedigree data by adding information about the family that each individual was born in and the family in which the individual is the reproductive animal.

Usage

```
org_fams(ped, sampledata, output = "both")
```

Arguments

ped	Data frame. FamAgg output of <code>get_colony()</code> or <code>get_ped()</code> function. With <code>rm_obsolete_parents</code> parameter set to TRUE.
sampledata	Data frame. Metadata for all genetic samples that belong to the individuals included in pedigree reconstruction analysis. This data frame should adhere to the formatting and naming conventions outlined in the <code>check_sampledata()</code> documentation.
output	Character string. Determines the format of the output. Options are: "ped": returns an extended pedigree data frame. "fams": returns a table of all families present in the pedigree. "both": returns a list with two data frames: "ped" and "fams". (Default)

Details

The result of `org_fams()` function introduces us to two important concepts within the context of this package: family and half-sib group. A family in the output of this function is defined as a group of animals where at least one parent and at least one offspring is known. A half-sib group refers to a group of half-siblings, either maternally or paternally related. In the function output the `DadHSgroup` groups paternal half-siblings and `MomHSgroup` maternal half-siblings.

The `fams` output dataframe contains `famStart` and `famEnd` columns, which estimate a time window for the family based solely on sample collection dates provided in `sampledata`. `famStart` marks the date of the earliest sample collected from any offspring belonging to that family. `famEnd` indicates the date of the latest sample collected from either the mother or the father of that family. It is important to recognize that this method relies on observation (sampling) times. Consequently, `famEnd` (last parental sample date) can precede `famStart` (first offspring sample date), creating a biologically impossible sequence and a negative calculated family timespan. Users should interpret the interval between `famStart` and `famEnd` with this understanding.

Value

Depending on the output parameter, the function returns either a data frame (ped or fams) or a list containing both data frames (ped and fams).

- ped data frame. An extended version of the pedigree data from `get_colony()/get_ped()`. In addition to common pedigree information (individual, mother, father, sex, family), ped includes columns for:
 - parents: Identifier codes of both parents separated with `_`.
 - FamID: Numeric identifier for the family to which the individual belongs (see fams below).
 - FirstSeen: Date of first sample of individual.
 - LastSeen: Date of last sample of individual.
 - IsDead: Logical value (TRUE/FALSE) that identifies if the individual is dead.
 - DadHSgroup: Identifier of paternal half-sib group (see Details).
 - MomHSgroup: Identifier of maternal half-sib group (see Details).
 - hsGroup: Numeric value indicating if the individual is part of a half-sib group (see Details).
- fams data frame includes information on families that individuals in the pedigree belong to. The families are described by:
 - parents: Identifier codes of both parents separated with `_`.
 - father: Identifier code of the father.
 - mother: Identifier code of the mother.
 - FamID: Numeric identifier for the family.
 - famStart: Date when the first sample of one of the offspring from this family was collected (see Details).
 - famEnd: Date when the last sample of mother or father of this family was collected (see Details).
 - FamDead: Logical value (TRUE/FALSE) indicating if the family no longer exists.
 - DadHSgroup: Identifier connecting families that share the same father.
 - MomHSgroup: Identifier connecting families that share the same mother.
 - hsGroup: Numeric value connecting families that share one of the parents.

Examples

```
# Prepare the data for usage with org_fams() function.
# Get animal timespan data using the anim_timespan() function.
animal_ts <- anim_timespan(
  wolf_samples$AnimalRef,
  wolf_samples$Date,
  wolf_samples$SType,
  dead = c("Tissue")
)
# Add animal timespan to the sampledata
sampledata <- merge(wolf_samples, animal_ts, by.x = "AnimalRef", by.y = "ID", all.x = TRUE)
# Define the path to the pedigree data file.
path <- paste0(system.file("extdata", package = "wpeR"), "/wpeR_samplePed")
# Retrieve the pedigree data from the get_colony function.
ped_colony <- get_colony(path, sampledata, rm_obsolete_parents = TRUE, out = "FamAgg")

# Run the function
# Organize families and expand pedigree data using the org_fams function.
```

```
org_fams(
  ped = ped_colony,
  sampledata = sampledata
)
```

ped_satplot

Temporal plot of pedigree

Description

Creates "capture" history plot of individuals arranged by families included in data frame created by [plot_table\(\)](#) function.

Usage

```
ped_satplot(
  plottable,
  famSpacing = 2,
  hsGroupSpacing = 2,
  xWhiteSpace = 100,
  xlabel = "Date",
  ylabel = "Animal",
  title = "",
  subtitle = "",
  LegendLabel = "Sex",
  xlegend = 0.2,
  ylegend = 0.94,
  text_size = 2.5,
  fam_label_size = 2
)
```

Arguments

plottable	Data frame. Output of plot_table() function.
famSpacing	Y-axis spacing between families. Should be even number!
hsGroupSpacing	Y-axis spacing between half-sib groups. Should be even number!
xWhiteSpace	Spacing on the X-axis at the beginning and end of the plot.
xlabel	X-axis label.
ylabel	Y-axis label.
title	Plot title.
subtitle	Plot subtitle.
LegendLabel	Title of the legend.
xlegend	Horizontal position of the legend.

ylegend Vertical position of the legend.
 text_size Plot text size.
 fam_label_size Family label text size.

Value

A graphical representation of detected family members through time.

Examples

```

# Prepare the data for usage with plot_table() function.
# Get animal timespan data using the anim_timespan() function.
animal_ts <- anim_timespan(wolf_samples$AnimalRef,
  wolf_samples$Date,
  wolf_samples$SType,
  dead = c("Tissue")
)
# Add animal timespan to the sampledata
sampledata <- merge(wolf_samples, animal_ts, by.x = "AnimalRef", by.y = "ID", all.x = TRUE)
# Define the path to the pedigree data file.
path <- paste0(system.file("extdata", package = "wpeR"), "/wpeR_samplePed")
# Retrieve the pedigree data from the get_colony function.
ped_colony <- get_colony(path, sampledata, rm_obsolete_parents = TRUE, out = "FamAgg")
# Organize families and expand pedigree data using the org_fams function.
org_tables <- org_fams(ped_colony, sampledata, output = "both")
# Prepare data for plotting.
pt <- plot_table(plot_fams = 1,
  org_tables$fams,
  org_tables$ped,
  sampledata,
  deadSample = c("Tissue")
)

# Run the function.
# Get a temporal pedigree plot.
ped_satplot(plottable = pt)

```

ped_spatial

Get files for spatial representation of pedigree

Description

Creates georeferenced data for spatial pedigree representation from the output of `plot_table()` function.

Usage

```
ped_spatial(
  plottable,
  na.rm = TRUE,
  output = "list",
  fullsibdata = NULL,
  sibthreshold = 0,
  path = "",
  filename = "",
  out.format = "geopackage",
  time.limits = c(as.Date("1900-01-01"), as.Date("2100-01-01")),
  time.limit.rep = FALSE,
  time.limit.offspring = FALSE,
  time.limit.moves = FALSE
)
```

Arguments

plottable	Data frame. Output of <code>plot_table()</code> function.
na.rm	Logical (TRUE/FALSE). Remove samples with missing coordinates and/or dates.
output	Character vector specifying the desired output type ('list' - default or 'gis'). Available outputs: list: all spatial data returned as list, gis: all spatial data returned as georeferenced files.
fullsibdata	Data frame with COLONY full-sibling data.
sibthreshold	Numeric. P-value threshold for sibship assignment.
path	System path for storing georeferenced files.
filename	Common name for all georeferenced files.
out.format	Character string. Type of georeferenced files to be generated. Can be either "geopackage" or "shapefile". Default is "geopackage"
time.limits	Vector of two Date values as the time window.
time.limit.rep	Logical (TRUE/FALSE). Apply time limits to reference samples of reproductive animals.
time.limit.offspring	Logical (TRUE/FALSE). Apply time limits to reference samples of offspring.
time.limit.moves	Logical (TRUE/FALSE). Apply time limits to movement data.

Details

The parameters `path`, `filename` and `out.format`, are used only when `output` parameter is set to "gis", since they control which georeferenced files should be created, where they will be saved and which common file name will they have.

Value

Depending on the output parameter the function can return a list of *sf* objects, a georeferenced vector data files or both.

Most of the objects are created separately for mothers, fathers and offspring, this include:

- Reference Points (*motherRpoints*, *fatherRpoints*, and *offspringRpoints*).
 - Each point corresponds to an animal included in the `'plot_table()'` function output.
 - For reproductive animals (mothers and fathers), a reference point is the location of their last sample within the specified time window.
 - For offspring, the reference point is the location of their first sample within the time window.
- Movement Points (*motherMovePoints*, *fatherMovePoints*, and *offspringMovePoints*).
 - These points represent all the samples of the respective animals.
- Movement Lines (*motherMoveLines*, *fatherMoveLines* and *offspringMoveLines*).
 - Movement lines connect all `'...MovePoints'` of a specific animal in chronological order.
- Movement Polygons (*motherMovePolygons*, *fatherMovePolygons* and *offspringMovePolygons*):
 - Movement polygons represent a convex hull that encloses all the samples of an individual.
 - An individual must have more than two samples for this representation.

Besides that the function also produces lines that connect mothers and their offspring (*maternityLines*), fathers and their offspring (*paternityLines*), and if *fullsibdata* parameter is specified, full siblings (*FullsibLines*).

Examples

```
# Prepare the data for usage with ped_spatial() function.
# Get animal timespan data using the anim_timespan() function.
animal_ts <- anim_timespan(wolf_samples$AnimalRef,
  wolf_samples$Date,
  wolf_samples$SType,
  dead = c("Tissue")
)
# Add animal timespan to the sampledata
sampledata <- merge(wolf_samples, animal_ts, by.x = "AnimalRef", by.y = "ID", all.x = TRUE)
# Define the path to the pedigree data file.
path <- paste0(system.file("extdata", package = "wpeR"), "/wpeR_samplePed")
# Retrieve the pedigree data from the get_colony function.
ped_colony <- get_colony(path, sampledata, rm_obsolete_parents = TRUE, out = "FamAgg")
# Organize families and expand pedigree data using the org_fams function.
org_tables <- org_fams(ped_colony, sampledata, output = "both")
# Prepare data for plotting.
pt <- plot_table(plot_fams = 1,
  org_tables$fams,
  org_tables$ped,
  sampledata,
  deadSample = c("Tissue")
)
```

```
# Run the function
# Get files for spatial pedigree representation in list format.
ped_spatial(plottable = pt)
```

plot_table	<i>Prepares pedigree data for plotting and spatial representation</i>
------------	-----------------------------------------------------------------------

Description

Combines extended pedigree (obtained by [org_fams\(\)](#) function) and sample metadata data for visual ([ped_satplot\(\)](#)) and spatial ([ped_spatial\(\)](#)) representation of the pedigree.

Usage

```
plot_table(
  plot_fams = "all",
  all_fams,
  ped,
  sampledata,
  datacolumns = c("Sample", "AnimalRef", "GeneticSex", "Date", "SType", "lat", "lng",
    "FirstSeen", "LastSeen", "IsDead"),
  deadSample = c("Tissue")
)
```

Arguments

plot_fams	Character string or numeric vector. FamID numbers from fams data generated by org_fams() function. If all families want to be plotted it is defined as character string "all". For a subset of families a numeric vector of FamIDs has to be specified. Defaults to "all".
all_fams	Data frame. Family (fams) data generated by org_fams() function.
ped	Data frame. Organized pedigree (ped) generated by org_fams() function.
sampledata	Data frame. Metadata for all genetic samples that belong to the individuals included in pedigree reconstruction analysis. For description of sampledata structure and sample information needed for plot_table() see Details.
datacolumns	Vector of column names included sampledata that are needed to produce this functions output (see Details).
deadSample	Single value or vector of different lethal sample types. Defaults to <code>c("Tissue")</code> .

Details

- sampledata has to include columns that contain information on:
 - unique identifier of each sample; character or numeric (default column name = Sample, see [check_sampledata\(\)](#) function),

- date of sample collection in Date format (default = Date),
- assignment of sample to particular individual; character or numeric (default = AnimalRef, see [check_sampledata\(\)](#) function),
- sex of the animal coded as F, M or NA; character (default = GeneticSex, see [check_sampledata\(\)](#) function),
- longitude and latitude coordinates of sample collection location; numeric (default = lng and lat, see [check_sampledata\(\)](#) function),
- type of particular sample eg. scat, tissue, saliva; character (default = SType, see [check_sampledata\(\)](#) function),
- date of first and last sample of individual in Date format (default = FirstSeen and LastSeen, see [anim_timespan\(\)](#) function),
- value identifying if if the individual is dead; logical (default = IsDead, see [anim_timespan\(\)](#) function).

Value

Extended sampledata data frame that includes all columns defined in datacolumns parameter and adds information needed for visual and spatial representation of pedigree:

- plottingID: Numeric. Identifier number for temporal pedigree plot [ped_satplot\(\)](#). In case of polygamous animals same individual can be included in more than one family.
- FamID: Numeric. Identifier number of family that individual belongs to.
- hsGroup: Numeric. Identifier number for the half-sib group of individual.
- rep: Logical. Is individual reproductive in current family, (current family defined with FamID for a particular entry).
- later_rep: Logical. Is individual reproductive in any other (later) families.
- isPolygamous: Logical. Does the individual have more than one mate.
- dead: Logical. Is individual dead.
- first_sample: Logical. Is this particular sample the first sample of the individual.
- last_sample: Logical. Is this particular sample the last sample of the individual.
- isReference: Logical. Is this particular sample reference sample of individual.

Examples

```
# Prepare the data for usage with plot_table() function.
# Get animal timespan data using the anim_timespan() function.
animal_ts <- anim_timespan(wolf_samples$AnimalRef,
  wolf_samples$Date,
  wolf_samples$SType,
  dead = c("Tissue")
)
# Add animal timespan to the sampledata
sampledata <- merge(wolf_samples, animal_ts, by.x = "AnimalRef", by.y = "ID", all.x = TRUE)
# Define the path to the pedigree data file.
path <- paste0(system.file("extdata", package = "wpeR"), "/wpeR_samplePed")
# Retrieve the pedigree data from the get_colony function.
ped_colony <- get_colony(path, sampledata, rm_obsolete_parents = TRUE, out = "FamAgg")
```

```

# Organize families and expand pedigree data using the org_fams function.
org_tables <- org_fams(ped_colony, sampledata, output = "both")

# Run the function
# Prepare data for plotting.
plot_table(plot_fams = "all",
  org_tables$fams,
  org_tables$ped,
  sampledata,
  deadSample = c("Tissue")
)

```

wolf_samples

Wolf monitoring genetic samples metadata

Description

Metadata of selected genetic samples of wolves collected between 2015 and 2021, in the scope of Slovenian National Wolf Monitoring

Usage

wolf_samples

Format

A data frame with 407 rows and 7 columns:

Sample Sample unique identifier code

Date Date of sample collection (format: YYYY-MM-DD)

AnimalRef Identification string for particular animal

GeneticSex Sex of animal to which the sample belong (format: M = male, F = female)

lat latitude (N-S) of the sample (CRS: WGS84; EPSG: 4326)

lng longitude (W-E) of the sample (CRS: WGS84; EPSG: 4326)

SType Type of the sample. (Direct Saliva, Scat, Urine, Saliva, Tissue, Decomposing Tissue, Blood)

Source

Slovenian National Wolf Monitoring

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