

Package: barrks (via r-universe)

November 5, 2024

Type Package

Title Calculate Bark Beetle Phenology Using Different Models

Version 1.0.1

Description Calculate the bark beetle phenology based on raster data or point-related data. There are multiple models implemented for two bark beetle species. The models can be customized and their submodels (onset of infestation, beetle development, diapause initiation, mortality) can be combined. The following models are available in the package: PHENIPS-Clim (first-time release in this package), PHENIPS (Baier et al. 2007) <[doi:10.1016/j.foreco.2007.05.020](https://doi.org/10.1016/j.foreco.2007.05.020)>, RITY (Ogris et al. 2019) <[doi:10.1016/j.ecolmodel.2019.108775](https://doi.org/10.1016/j.ecolmodel.2019.108775)>, CHAPY (Ogris et al. 2020) <[doi:10.1016/j.ecolmodel.2020.109137](https://doi.org/10.1016/j.ecolmodel.2020.109137)>, BSO (Jakoby et al. 2019) <[doi:10.1111/gcb.14766](https://doi.org/10.1111/gcb.14766)>, Lange et al. (2008) <[doi:10.1007/978-3-540-85081-6_32](https://doi.org/10.1007/978-3-540-85081-6_32)>, Jönsson et al. (2011) <[doi:10.1007/s10584-011-0038-4](https://doi.org/10.1007/s10584-011-0038-4)>. The package may be expanded by models for other bark beetle species in the future.

URL <https://jjentschke.github.io/barrks/>,
<https://github.com/jjentschke/barrks/>

BugReports <https://github.com/jjentschke/barrks/issues/>

License GPL (>= 3)

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RdMacros Rdpack

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Collate 'appearance.R' 'barrks-package.R' 'bso-phenology-results.R'
'bso-phenology.R' 'utils.R' 'bso-plot.R' 'create-daylengths.R'
'create-events.R' 'create-suntimes.R' 'data.R'
'utils-model-functions.R' 'model.R' 'model-bso.R'
'model-phenips.R' 'model-rity.R' 'model-chapy.R'
'model-combine.R' 'model-joensson.R' 'model-lange.R'

'model-phenips-clim.R' 'phenology-properties.R'
 'phenology-results.R' 'phenology.R'
 'plot-development-diagram.R' 'stations.R' 'utils-doc.R'
 'utils-log.R' 'utils-storage.R'

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analyse.phenology	<i>Analyse a phenology</i>
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Description

Here, all functions are listed that are available to analyse the results of a `phenology()`-call.

Details

Get phenology properties:

- `prop_dates()`
- `prop_filial_generations()`
- `prop_first_date()`
- `prop_hatched_generations()`
- `prop_last_date()`
- `prop_sister_broods()`
- `prop_stations()`
- `prop_year()`

Get phenology results (raster-based):

- `get_development_rst()`
- `get_diapause_rst()`
- `get_generations_rst()`
- `get_hibernating_generations_rst()`
- `get_mortality_rst()`

- `get_onset_rst()`

Get phenology results (station-based):

- `get_development_df()`
- `get_diapause_df()`
- `get_generations_df()`
- `get_hibernating_generations_df()`
- `get_mortality_df()`
- `get_onset_df()`

Plot phenology results (station-based):

- `plot_development_diagram()`

See Also

[analyse.phenology.bso](#)

analyse.phenology.bso *Analyse a BSO generated phenology*

Description

Here, all functions are listed that are available to analyse the results of a `bso_phenology()` call.

Details

Get BSO phenology properties:

- `prop_dates()`
- `prop_filial_generations()`
- `prop_first_date()`
- `prop_hatched_generations()`
- `prop_last_date()`
- `prop_sister_broods()`
- `prop_stations()`
- `prop_year()`

Get BSO phenology results (raster-based):

- `bso_get_flight_rst()`
- `bso_get_individuals_rst()`

Get BSO phenology results (station-based):

- [bso_get_flight_df\(\)](#)
- [bso_get_individuals_df\(\)](#)

Plot BSO phenology results (station-based):

- [bso_plot_flight_diagram\(\)](#)
- [bso_plot_stage_diagram\(\)](#)

See Also

[analyse.phenology](#)

barrks_colors	<i>Get barrks default color palettes</i>
---------------	--

Description

Get barrks default color palettes.

Usage

```
barrks_colors(type = "raster")
```

Arguments

type Select the desired color palette. There are different variants for particular purposes. Allowed values are 'raster', 'diagram_lines', 'diagram_fill', 'bso_flight' and 'bso_stages'.

Value

A character vector of hex colors.

See Also

[barrks_labels\(\)](#)

Examples

```
colors <- barrks_colors()

# use the colors of 'barrks' for your individual plot...
```

barrks_data *Load sample data*

Description

The package comes with sample data that allow the application of all models available. The following data sets are available:

Usage

```
barrks_data(dataset = "raster")
```

Arguments

dataset Choose the data set that should be returned.

Details

- raster Contains a list of raster weather datasets for a sample area. The data was taken from Deutscher Wetterdienst (DWD).
- stations Contains sample station weather data for some cities in Germany. The data was taken from Deutscher Wetterdienst (DWD). Missing global radiation values were replaced by the mean value of the other stations.
- station_coords Contains the coordinates (longitude/latitude) of the stations that are included in the stations data set. The data was taken from Deutscher Wetterdienst (DWD).

Value

The respective data set. Can be a list of SpatRasters (for dataset = 'raster') or a data frame.

Source

- https://opendata.dwd.de/climate_environment/CDC/grids_germany/daily/hyras_de/
- https://opendata.dwd.de/climate_environment/CDC/observations_germany/climate/daily/kl/historical/
- https://opendata.dwd.de/climate_environment/CDC/observations_germany/climate/daily/solar/

Examples

```
# plot first layer of the minimum temperature of the sample raster data
terra::plot(barrks_data()$tmin[[1]])

# print the first lines of the sample station data
head(barrks_data('stations'), 10)

# print the coordinates of the sample stations
barrks_data('station_coords')
```

barrks_labels	<i>Get barrks default legend labels</i>
---------------	---

Description

Get barrks default legend labels.

Usage

```
barrks_labels(type = "raster")
```

Arguments

type Select the desired legend labels. There are different variants for particular purposes. Allowed values are 'raster', 'diagram', 'bso_flight' and 'bso_stages'.

Value

A character vector of labels.

See Also

[barrks_colors\(\)](#)

Examples

```
labels <- barrks_labels()

# use the labels of 'barrks' for your individual plot...
```

bso_get_flight_rst	<i>Get flight of individuals (BSO only)</i>
--------------------	---

Description

Get the number of individuals that are flying.

Usage

```
bso_get_flight_rst(pheno, generation, flight = 1, dates = prop_dates(pheno))

bso_get_flight_df(
  pheno,
  generation,
  stations = prop_stations(pheno),
  flight = 1,
  dates = prop_dates(pheno)
)
```

Arguments

pheno	A BSO phenology (see bso_phenology())
generation	Generation of interest. For sister broods, 0.5 should be added.
flight	Specifies which flight of the respective generation should be returned. Can be 1 (first flight) or 2 (second flight).
dates	Select dates that should be present in the output.
stations	Pass a character vector to choose stations assigned to pheno by their names, or pass different stations. See stations_create() for details.

Value

- `bso_get_flight_rst`: a multi-layer SpatRaster.
- `bso_get_flight_df`: a data frame.

Examples

```
# This may take a few minutes...

# calculate phenology
p <- bso_phenology('bso', barrks_data('stations'), .quiet = TRUE)

# get the number of individuals of the hibernating generation on their first flight
bso_get_flight_df(p, 0, 'Freiburg', flight = 1)
```

```
bso_get_individuals_rst
      Get individuals (BSO only)
```

Description

Get the number of individuals of a generation that are in a specific development stage.

Usage

```
bso_get_individuals_rst(
  pheno,
  generation,
  stage = "all",
  dates = prop_dates(pheno)
)

bso_get_individuals_df(
  pheno,
  generation,
  stations = prop_stations(pheno),
```



```

    stage = "all",
    dates = prop_dates(pheno)
  )

```

Arguments

pheno	A BSO phenology (see bso_phenology())
generation	Generation of interest. For sister broods, 0.5 should be added.
stage	If it is a numeric, the individuals of the slots specified will be retrieved. Otherwise it could be one of the following values: all, egg, larva, pupa, white (egg + larva + pupa), maturation, preflight, reproduction, brown (maturation + preflight + reproduction)
dates	Select dates that should be present in the output.
stations	Pass a character vector to choose stations assigned to pheno by their names, or pass different stations. See stations_create() for details.

Value

- `bso_get_individuals_rst()`: a multi-layer SpatRaster.
- `bso_get_individuals()`: a data frame.

Examples

```

# This may take a few minutes...

# calculate phenology
p <- bso_phenology('bso', barrks_data('stations'), .quiet = TRUE)

# get the number of individuals of the hibernating generation that are waiting to fly
bso_get_individuals_df(p, 0, 'Freiburg', stage = 'preflight')

```

```
bso_plot_flight_diagram
```

Plot a flight diagram (BSO only)

Description

A flight diagram illustrates the daily share of flying individuals over time.

Usage

```

bso_plot_flight_diagram(
  .pheno,
  .station = prop_stations(.pheno)[1],
  .colors = barrks_colors("bso_flight"),
  .labels = barrks_labels("bso_flight"),

```

```

    .xlim = NULL,
    .legend = "topright",
    ...
  )

```

Arguments

<code>.pheno</code>	A BSO phenology (see bso_phenology())
<code>.station</code>	Pass a character vector to choose a station assigned to <code>.pheno</code> by its name, or pass a different station. See stations_create() for details.
<code>.colors, .labels</code>	Vectors of colors/labels starting from the first and the second flight of the hibernating generation followed consecutively by elements for the filial generations (first and second flight).
<code>.xlim</code>	Date vector of length to that limits the dates plotted.
<code>.legend</code>	Pass FALSE if no legend should be plotted. Otherwise the value will be passed to <code>legend()</code> as first argument. Look there for more information.
<code>...</code>	arguments passed to <code>graphics::barplot()</code> .

Value

None

Examples

```

# This may take a few minutes...

# calculate phenology
p <- bso_phenology('bso', barrks_data('stations'), .quiet = TRUE)

bso_plot_flight_diagram(p)

```

`bso_plot_stage_diagram`

Plot a stage diagram (BSO only)

Description

A stage diagram illustrates the share of individuals that are in a specific developmental stage of a particular generation over time.

Usage

```
bso_plot_stage_diagram(
  .pheno,
  .station = prop_stations(.pheno)[1],
  .stages = list("white", "brown"),
  .lty = c("dashed", "solid"),
  .lwd = 2,
  .colors = barrks_colors("bso_stages"),
  .labels = barrks_labels("bso_stages"),
  .legend_col = TRUE,
  .legend_lty = TRUE,
  ...
)
```

Arguments

<code>.pheno</code>	A BSO phenology (see bso_phenology())
<code>.station</code>	Pass a character vector to choose a station assigned to <code>.pheno</code> by its name, or pass a different station. See stations_create() for details.
<code>.stages</code>	List of stages to plot. Elements will be passed to bso_get_individuals_df() . Look there for more information.
<code>.lty, .lwd</code>	Vectors of line types or line widths that are used to plot the different stages. Should have the same length as <code>.stages</code> or 1.
<code>.colors, .labels</code>	Vectors of colors/labels starting from the hibernating generation followed consecutively by elements for the filial generations (not including sisterbroods).
<code>.legend_col, .legend_lty</code>	Manipulate the appearance of the legends for colors and line types. Pass TRUE/FALSE to enable/disable the respective legend. For the customization of the respective legend, a list of parameters for graphics::legend can be passed.
<code>...</code>	arguments passed to base::plot() .

Value

None

Examples

```
# This may take a few minutes...

# calculate phenology
p <- bso_phenology('bso', barrks_data('stations'), .quiet = TRUE)

bso_plot_stage_diagram(p)
```

`bso_translate_phenology`*Translate BSO generated phenology*

Description

A BSO generated phenology cannot be analysed in the same way as other phenology objects. To be able to use the functions that are available for phenology objects returned by `phenology()`, the BSO generated phenology should be translated.

Usage

```
bso_translate_phenology(pheno, threshold = 0.1, .quiet = FALSE)
```

Arguments

<code>pheno</code>	A BSO phenology (see <code>bso_phenology()</code>)
<code>threshold</code>	Share of individuals that must have reached a specific development in the BSO phenology to account for them in the corresponding standard phenology.
<code>.quiet</code>	If TRUE, messages are suppressed.

Value

Returns a standard phenology as a list. Look [here](#) to find out how a phenology can be analysed. It is not recommended to access the list elements directly.

Examples

```
# This may take a few minutes...

# calculate and translate BSO phenology
p <- bso_phenology('bso', barrks_data('stations'), .quiet = TRUE)
pt <- bso_translate_phenology(p, .quiet = TRUE)

# print the generations data frame of station 'Freiburg'
df <- get_generations_df(pt, 'Freiburg')
df
```

`categorize_generations_rst`*Make a numeric generations raster categorical*

Description

Make a numeric generations raster categorical. Useful when mathematical operations were performed with generations rasters (use `get_generations_rst(..., categorical = FALSE)` to get numeric generations rasters).

Usage

```
categorize_generations_rst(  
  rst,  
  colors = barrks_colors(),  
  labels = barrks_labels()  
)
```

Arguments

<code>rst</code>	A numeric <code>SpatRaster</code> that represents bark beetle generations. Sister broods are defined by adding 0.5 to the respective generation.
<code>colors, labels</code>	Vectors of colors/labels starting from zero generations followed consecutively by elements for the respective generations (including sister broods).

Value

A categorical `SpatRaster`.

Examples

```
# calculate phenology with different models  
p1 <- phenology('phenips-clim', barrks_data(), .quiet = TRUE)  
p2 <- phenology('phenips', barrks_data(), .quiet = TRUE)  
  
# get the generation as numerical rasters to allow mathematical operations  
gens1 <- get_generations_rst(p1, categorical = FALSE)  
gens2 <- get_generations_rst(p2, categorical = FALSE)  
  
# calculate the maximum generations from the 2 models  
gens_max <- max(gens1, gens2)  
# categorize the results  
gens_max_cat <- categorize_generations_rst(gens_max)  
  
# plot the categorized raster  
terra::plot(gens_max_cat)  
# plot the uncategorized raster  
terra::plot(gens_max)
```

create_daylength_df *Create a data frame of day lengths*

Description

Generate a data frame of day lengths for given latitudes. The package `geosphere` is required to use this function.

Usage

```
create_daylength_df(lat, dates, .quiet = FALSE)
```

Arguments

lat	Data frame with the fields <code>station</code> and <code>lat</code> . Defines the latitude for the respective stations.
dates	Dates that should be processed.
.quiet	If TRUE, messages are suppressed.

Value

A data frame with the columns `date`, `station` and `daylength`.

See Also

[create_daylength_rst\(\)](#)

Examples

```
# dates of interest
date_start <- as.Date('2020-01-01')
date_end <- as.Date('2020-12-31')

# calculate day length
dl <- create_daylength_df(barrks_data('station_coords'),
  seq(date_start, date_end, by = 'day'),
  .quiet = TRUE)

# print day lengths of station 'Freiburg'
dl[dl$station == 'Freiburg',]
```

create_daylength_rst *Create day length rasters*

Description

Generate a multi-layer SpatRaster of day lengths for a given template. The package `geosphere` is required to use this function.

Usage

```
create_daylength_rst(  
  template,  
  dates = terra::time(template),  
  crs = "EPSG:4258",  
  .quiet = FALSE  
)
```

Arguments

<code>template</code>	(Multi-layer) SpatRaster that determines the spatial extent of the result.
<code>dates</code>	Dates that should be processed. If not specified, the dates of <code>template</code> are used through <code>terra::time()</code> .
<code>crs</code>	Coordinate reference system with longitude/latitude metrics. It is used to project the raster coordinates to be able to retrieve the latitude.
<code>.quiet</code>	If TRUE, messages are suppressed.

Value

A multi-layer SpatRaster. Each layer represents one date.

See Also

[create_daylength_df\(\)](#)

Examples

```
# calculate day length, use barrks_data()$tmin as template  
dl <- create_daylength_rst(barrks_data()$tmin, .quiet = TRUE)  
  
# plot day length on May 1st, 2015  
terra::plot(dl[[terra::time(dl) == '2015-05-01']])
```

create_onset	<i>Create phenological events (onset/diapause/mortality)</i>
--------------	--

Description

Generate onset, diapause or mortality manually to be able to run `phenology()` with observed or arbitrary inputs.

Usage

```
create_onset(
  template,
  doys = NULL,
  stations = NULL,
  dates = NULL,
  .quiet = FALSE
)
```

```
create_diapause(
  template,
  doys = NULL,
  stations = NULL,
  dates = NULL,
  .quiet = FALSE
)
```

```
create_mortality(
  template,
  doys = NULL,
  stations = NULL,
  dates = NULL,
  .quiet = FALSE
)
```

Arguments

template	SpatRaster or data frame that determines the spatial and temporal extent of the result. If a single-layer SpatRaster was passed, the temporal extent should be defined by using dates.
doys	Numeric vector, (multi-layer) SpatRaster or data frame that specifies the days of year when the event is triggered. Vectors will define the events globally whereas SpatRasters allow spatially explicit definitions. For the creation of events based on stations, data frames are used. In that case, the field <code>station</code> specifies the station name and <code>doy</code> indicates the respective day of year.
stations	If <code>template</code> is a SpatRaster and <code>doys</code> is a data frame, <code>stations</code> should be passed to define which cells are affected. See <code>stations_create()</code> for details.

dates	Dates to define the temporal extent of the output if template is a single-layer SpatRaster.
.quiet	If TRUE, messages are suppressed.

Value

A logical multi-layer SpatRaster. Each layer represents one date.

Functions

- create_onset(): Create a onset.
- create_diapause(): Create a diapause.
- create_mortality(): Create mortality events.

Examples

```
# load sample data
d <- barrks_data('stations')

# create onset, diapause, mortality
on <- create_onset(d, lubridate::yday('2015-04-15'))
dia <- create_diapause(d, lubridate::yday('2015-08-15'))
mort <- create_mortality(d, lubridate::yday('2015-11-15'))

# calculate phenology
p <- phenology('phenips-clim', d, .quiet = TRUE,
               .onset = on, .diapause = dia, .mortality = mort)

# plot development
plot_development_diagram(p, .lwd = 4)
```

create_suntimes_df *Create a data frame of sunrises and sunsets*

Description

Generate a data frame that specifies sunrises and sunsets for different coordinates and dates. The package `suncalc` is required to use this function.

Usage

```
create_suntimes_df(coords, dates, tz = Sys.timezone(), .quiet = FALSE)
```

Arguments

coords	Data frame with the fields station, lat and lon. Defines the latitude and longitude for the respective stations.
dates	Dates that should be processed.
tz	Timezone of the results.
.quiet	If TRUE, messages are suppressed.

Value

A data frame with the columns date, station and sunrise and sunset. The values of sunrise and sunset indicate the respective time in minutes.

See Also

[create_suntimes_rsts\(\)](#)

Examples

```
date_start <- as.Date('2020-01-01')
date_end <- as.Date('2020-12-31')

st <- create_suntimes_df(barrks_data('station_coords'),
                        seq(date_start, date_end, by = 'day'),
                        .quiet = TRUE)

# print results of station 'Freiburg'
st[st$station == 'Freiburg',]
```

create_suntimes_rsts *Create rasters that indicate sunrise and sunset*

Description

Generate a list of two multi-layer SpatRasters for a given template that indicate sunrise and sunset for the respective cells. The package `sunCalc` is required to use this function.

Usage

```
create_suntimes_rsts(
  template,
  dates = terra::time(template),
  crs = "EPSG:4258",
  tz = Sys.timezone(),
  .quiet = FALSE
)
```

Arguments

template	(Multi-layer) SpatRaster that determines the spatial extent of the result.
dates	Dates that should be processed. If not specified, the dates of template are used through <code>terra::time()</code> .
crs	Coordinate reference system with longitude/latitude metrics. It is used to project the raster coordinates to be able to retrieve longitude and latitude.
tz	Timezone of the results.
.quiet	If TRUE, messages are suppressed.

Value

A list with the elements sunrise and sunset which are both multi-layer SpatRasters. The values indicate the respective time in minutes. Each layer represents one date.

See Also

`create_suntimes_df()`

Examples

```
# calculate suntimes, use barrks_data()$tmin as template
st <- create_suntimes_rsts(barrks_data()[[1]], .quiet = TRUE)

# plot results on May 1st, 2015
terra::plot(st$sunrise[[terra::time(st$sunrise) == '2015-05-01']])
terra::plot(st$sunset[[terra::time(st$sunset) == '2015-05-01']])
```

get_development_rst *Get the beetles development*

Description

Get the beetles development of specific generations. A value of -1 implies that the generation is not present yet.

Usage

```
get_development_rst(pheno, generation, dates = prop_dates(pheno))

get_development_df(
  pheno,
  stations = prop_stations(pheno),
  generation = prop_hatched_generations(pheno),
  dates = prop_dates(pheno)
)
```

Arguments

pheno	A phenology (see phenology())
generation	Generation of interest. For sister broods, 0.5 should be added. get_development_df() allows multiple generations here.
dates	Select dates that should be present in the output.
stations	Pass a character vector to choose stations assigned to pheno by their names, or pass different stations. See stations_create() for details.

Value

- [get_development_rst\(\)](#): A multi-layer SpatRaster.
- [get_development_df\(\)](#): A data frame which contains a field for each generation (gen_1, gen_1.5, gen_2, gen_2.5, ...) requested.

Examples

```
# calculate station-based phenology
p <- phenology('phenips-clim', barrks_data('stations'), .quiet = TRUE)

# print the development data frame of station 'Freiburg'
df <- get_development_df(p, 'Freiburg')
df[,4:ncol(df)] <- round(df[,4:ncol(df)], 3) # round results
df
```

get_generations_rst *Get generations*

Description

Find out how many generations are present (or have reached a development threshold).

Usage

```
get_generations_rst(
  pheno,
  dates = prop_last_date(pheno),
  threshold = 0,
  generations = prop_hatched_generations(pheno),
  categorical = TRUE,
  colors = barrks_colors("raster"),
  labels = barrks_labels("raster")
)

get_generations_df(
  pheno,
```

```

    stations = prop_stations(pheno),
    dates = prop_dates(pheno),
    threshold = 0,
    generations = prop_hatched_generations(pheno)
  )

get_hibernating_generations_rst(
  pheno,
  categorical = TRUE,
  colors = barrks_colors("raster"),
  labels = barrks_labels("raster")
)

get_hibernating_generations_df(pheno, stations = prop_stations(pheno))

```

Arguments

pheno	A phenology (see phenology())
dates	Select dates that should be present in the output.
threshold	Threshold of the beetle development to account for a generation.
generations	Numeric vector that determines which generations should be included in the result.
categorical	Set FALSE if the <code>SpatRaster</code> that is returned should be numeric. Otherwise, it will be categorical.
colors, labels	Vectors of colors/labels starting from zero generations followed consecutively by elements for the respective generations (including sister broods).
stations	Pass a character vector to choose stations assigned to pheno by their names, or pass different stations. See stations_create() for details.

Value

- `get_generations_rst()`: A multi-layer `SpatRaster`.
- `get_hibernating_generations_rst()`: A `SpatRaster`. Only available if the model's end date has been reached. Otherwise all values will be NA.
- `get_generations_df()`: A data frame.
- `get_hibernating_generations_df()`: A data frame. Only available if the model's end date has been reached. Otherwise all values will be NA.

Examples

```

# calculate phenology
p <- phenology('phenips-clim', barrks_data(), .quiet = TRUE)

# get the generations raster
gens <- get_generations_rst(p)

# plot the generations raster

```

```
terra::plot(gens)
```

get_input_data	<i>Get (preprocessed) input data</i>
----------------	--------------------------------------

Description

The function returns a list that contains the input data of the phenology as well as some intermediate results that are needed as preprocessed inputs for the model. The result can be used as input for `phenology()` to avoid redundant calculations.

Usage

```
get_input_data(pheno)
```

Arguments

pheno A phenology (see [phenology\(\)](#))

Value

A list of `SpatRasters`.

Examples

```
# setup phenology
p <- phenology('phenips-clim', barrks_data(), .setup_only = TRUE, .quiet = TRUE)

# get the (preprocessed) input data
inputs <- get_input_data(p)

# print the names to show which input data is available
names(inputs)
```

get_onset_rst	<i>Get onset, diapause or mortality</i>
---------------	---

Description

Get onset, diapause or mortality as day of year or raw output. Note that multiple mortality events are possible over the season.

Usage

```

get_onset_rst(pheno, as_doy = TRUE, dates = prop_dates(pheno))

get_onset_df(
  pheno,
  stations = prop_stations(pheno),
  as_doy = TRUE,
  dates = prop_dates(pheno)
)

get_diapause_rst(pheno, as_doy = TRUE, dates = prop_dates(pheno))

get_diapause_df(
  pheno,
  stations = prop_stations(pheno),
  as_doy = TRUE,
  dates = prop_dates(pheno)
)

get_mortality_rst(pheno, as_doy = TRUE, dates = prop_dates(pheno))

get_mortality_df(
  pheno,
  stations = prop_stations(pheno),
  as_doy = TRUE,
  dates = prop_dates(pheno)
)

```

Arguments

pheno	A phenology (see phenology())
as_doy	If TRUE, the day(s) of year will be returned. If FALSE the phenological events will be returned in a raw format. Then, the return value could be used as input for phenology()/bso_phenology() (parameters <code>.onset</code> , <code>.diapause</code> and <code>.mortality</code>).
dates	Select dates that should be present in the output.
stations	Pass a character vector to choose stations assigned to pheno by their names, or pass different stations. See stations_create() for details.

Value

- `get_onset_rst()`, `get_diapause_rst()`, `get_mortality_rst()`: A (multi-layer) SpatRaster.
- `get_onset_df()`, `get_diapause_df()`, `get_mortality_df()`: A data frame.

Functions

- `get_onset_rst()`: Returns a (multi-layer) SpatRaster of the onset.

- `get_onset_df()`: Returns a data frame of the onset.
- `get_diapause_rst()`: Returns a (multi-layer) `SpatRaster` of the diapause.
- `get_diapause_df()`: Returns a data frame of the diapause.
- `get_mortality_rst()`: Returns a (multi-layer) `SpatRaster` of the mortality.
- `get_mortality_df()`: Returns a data frame of the mortality.

Examples

```
# calculate phenology
p <- phenology('phenips-clim', barrks_data(), .quiet = TRUE)

# plot onset, diapause, mortality
get_onset_rst(p) |> terra::plot()
get_diapause_rst(p) |> terra::plot()
get_mortality_rst(p)[[1]] |> terra::plot()
```

list_models

List all models

Description

Get the names of all available models.

Usage

```
list_models()
```

Value

A character vector.

Examples

```
# print all available models
list_models()
```

model	<i>Get a phenology model</i>
-------	------------------------------

Description

Returns a (customized) phenology model.

Usage

```
model(m, ...)
```

Arguments

`m` Name of the model or the return value of another `model()`-call.
`...` List of parameters to customize the model.

Value

A phenology model. Can be passed to [phenology\(\)](#).

See Also

Look at the customization manuals, to find out which parameters can be customized for a specific model: [model.bso.customize](#), [model.phenips.customize](#), [model.rity.customize](#), [model.chapy.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips_clim.customize](#).

Examples

```
# customize the temperature beetles need to fly for PHENIPS-Clim
m <- model('phenips-clim', tfly = 16)

# calculate phenology
p <- phenology(m, barrks_data(), .quiet = TRUE)

# plot generations
gens <- get_generations_rst(p)
terra::plot(gens)
```

model.bso.apply	Use BSO
-----------------	---------

Description

This page describes the usage of BSO with `phenology()`. The model-specific inputs are listed and its basic functionality is explained. BSO was published by Jakoby et al. (2019) and parametrized for *Ips typographus* in Switzerland. Note that the onset and the development submodel do not support the usage of a storage (except for some precalculations).

Arguments

...	See <code>phenology()</code> for a detailed description of the function.
tmin, tmax	Daily minimum/maximum air temperatures in °C.
sunrise, sunset	Time of sunrise/sunset in minutes from midnight. Can be created with <code>create_suntimes_rsts()</code> or <code>create_suntimes_df()</code> .
n	number of individuals to simulate.
max_generations	maximum number of generations to calculate.

Details

In barrks, `phenology()` is used to apply a model. The following code illustrates which inputs are required to apply BSO and which additional parameters are available.

```
bso_phenology("bso", ..., tmin, tmax, sunrise, sunset,
              n = 1e+09, max_generations = 4)

# calculate submodels separately
bso_phenology("bso", ..., .submodels = 'onset',
              tmin, tmax, sunrise, sunset, n = 1e+09)
bso_phenology("bso", ..., .submodels = 'diapause', tmin, tmax)
bso_phenology("bso", ..., .submodels = 'development',
              .onset, .diapause = NULL, .mortality = NULL,
              tmin, tmax, sunrise, sunset,
              max_generations = 4)
```

Value

The function returns a BSO phenology. Look [here](#) to find out how it can be analysed.

Functioning of the BSO

In the following, the basic functioning of BSO is explained.

- **Onset:** The onset of swarming will start when the degree days of the mean temperature reach a specific threshold and regeneration feeding of the individuals has finished (Look at development for details).
- **Development:** The development of single individuals is simulated. The simulation of each individual is realized by passing a multitude of slots that are grouped in stages. The hourly probability for an individual to enter the next slot depends on the current stage and the phloem temperature. The hourly temperature is derived from the minimum and maximum temperatures using a sine interpolation. The hourly phloem temperature is calculated using Newton's Law of Cooling (see Trân et al. 2007).
- **Diapause:** Specific photoperiod-related dates define when the diapause is initiated at the earliest and at the latest. In between these dates, the diapause is initiated when the mean temperature falls below a specific threshold.
- **Mortality:** BSO does not have a mortality submodel implemented.

Look [here](#) to find out how the model parameters affect the actual calculations and which values are used by default.

References

Jakoby O, Lischke H, Wermelinger B (2019). "Climate change alters elevational phenology patterns of the European spruce bark beetle (*Ips typographus*).” *Global Change Biology*, **25**(12), 4048-4063. doi:10.1111/gcb.14766.

Trân JK, Ylioja T, Billings RF, Régnière J, Ayres MP (2007). "Impact of minimum winter temperatures on the population dynamics of *Dendroctonus frontalis*.” *Ecological Applications*, **17**(3), 882-899. doi:10.1890/060512.

See Also

[model\(\)](#), [bso_phenology\(\)](#), [model.bso.customize](#)

Other phenology applications: [model.chapy.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips.apply](#), [model.phenips_clim.apply](#), [model.rity.apply](#)

model.bso.customize *Customize BSO*

Description

This page describes the parameters that can be used to customize BSO. The model was developed by Jakoby et al. (2019). Look [here](#) to find out how to apply the model.

Arguments

dd_onset_start_date

The date, when the degree days start to sum up ('MM-DD').

dd_onset_base Base temperature to calculate degree days to trigger the onset.

dd_onset_threshold	Degree days that are required before the individuals start regeneration feeding in slot_dia of the maturation stage. When the regeneration feeding has finished, the onset is triggered.
slot_dia	Maturation feeding slot where the individuals start regeneration feeding after diapause.
k	Factor for the calculation of the phloem temperature.
alpha, tlo, tup	Parameters used to calculate the transition probabilities for each stage (except preflight) in the following order: development, maturation feeding, reproduction.
tfly_min, tfly_max, pfly_max, beta	Parameters used to calculate the transition probabilities for the preflight stage.
num_slots	Named vector that defines the number of slots for each stage. The development stage is subdivided into the stages egg, larva and pupa.
psis	Probability that a sister brood will be established.
slot_sis	Maturation feeding slot where the individuals start regeneration feeding before they establish a sister brood.
model_end_date	Date when the model ends (no further development will be modeled).
diapause_first	The day of year when the diapause could start at the earliest.
diapause_last	The day of year when the diapause could start at the latest.
tdia_min	The diapause will be initiated when the average daily temperature falls below that value.

Details

In `barrks`, `model()` is used to customize a model. The following code illustrates which parameters are available for BSO and specifies their default values.

```
model("bso",

      # ==== onset ====

      dd_onset_start_date = '01-01',
      dd_onset_base = 5.124198,
      dd_onset_threshold = 100,
      slot_dia = 6,

      # ==== onset + development ====

      k = 2.853738e-02,
      alpha = c(2.549060e-05, 0.0000789, 1.009450e-05),
      tlo = c(-1.297644e+01, 4.760089e+00, -4.424628e+00),
      tup = c(3.600070e+01, 4.002483e+01, 3.999390e+01),
      tfly_min = 16.1064,
      tfly_max = 31.2901,
      pfly_max = 9.863263e-03,
```

```
beta = 1.363763,  
  
num_slots = c(  
  'reproduction' = 11,  
  'egg' = 18,  
  'larva' = 45,  
  'pupa' = 8,  
  'maturation' = 8,  
  'preflight' = 1  
)  
  
# ==== development ====  
  
psis = 2.994450e-01,  
slot_sis = 4,  
  
model_end_date = '12-30',  
  
# ==== diapause ====  
  
diapause_first = 210,  
diapause_last = 232,  
tdia_min = 1.645209e+01  
)
```

References

Jakoby O, Lischke H, Wermelinger B (2019). “Climate change alters elevational phenology patterns of the European spruce bark beetle (*Ips typographus*).” *Global Change Biology*, **25**(12), 4048-4063. [doi:10.1111/gcb.14766](https://doi.org/10.1111/gcb.14766).

See Also

[model\(\)](#), [phenology\(\)](#), [model.bso.apply](#)

Other model customizations: [model.chapy.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips.customize](#), [model.phenips_clim.customize](#), [model.rity.customize](#)

model.chapy.apply *Use CHAPY*

Description

This page describes the usage of CHAPY with [phenology\(\)](#). The model specific inputs are listed and its basic functionality is explained. CHAPY was published by Ogris et al. (2020) and parametrized for *Pityogenes chalcographus* in Slovenia.

Arguments

tmin, tmean, tmax	Daily minimum/mean/maximum air temperatures in °C. For the development submodel, the parameter that is obligatory depends on mode.
daylength	Length of the day in hours. Can be created with create_daylength_rst() or create_daylength_rst() .
mode	Specifies which temperature should be used to calculate the development. Can be min, mean or max.
.submodels, .onset, .diapause, .mortality, ...	See phenology() for a detailed description of the function.

Details

In barrks, [phenology\(\)](#) is used to apply a model. The following code illustrates which inputs are required to apply CHAPY and which additional parameters are available.

```
phenology("chapy", ..., tmin = NULL, tmean = NULL, tmax, daylength, mode = 'max')

# calculate submodels separately
phenology("chapy", ..., .submodels = 'onset', tmax)
phenology("chapy", ..., .submodels = 'diapause', daylength)
phenology("chapy", ..., .submodels = 'mortality', tmax)
phenology("chapy", ..., .submodels = 'development',
          .onset, .diapause = NULL, .mortality = NULL,
          tmin = NULL, tmean = NULL, tmax = NULL, mode = 'max')
```

Functioning

The functioning of CHAPY is identical to [RITY](#) but it is has a different [parametrization](#).

References

Ogris N, Ferlan M, Hauptman T, Pavlin R, Kavčič A, Jurc M, de Groot M (2020). "Sensitivity analysis, calibration and validation of a phenology model for *Pityogenes chalcographus* (CHAPY)." *Ecological Modelling*, **430**, 109137. ISSN 0304-3800, [doi:10.1016/j.ecolmodel.2020.109137](https://doi.org/10.1016/j.ecolmodel.2020.109137).

See Also

[model\(\)](#), [phenology\(\)](#), [model.chapy.customize](#)

Other phenology applications: [model.bso.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips.apply](#), [model.phenips_clim.apply](#), [model.rity.apply](#)

 model.chapy.customize *Customize CHAPY*

Description

This page describes the parameters that can be used to customize CHAPY. The model was developed by Ogris et al. (2020). Look [here](#) to find out how to apply the model.

Arguments

dd_onset_start_date	The date, when the degree days start to sum up ('MM-DD').
dd_onset_base	Base temperature to calculate degree days to trigger the onset.
dd_onset_threshold	Degree days that are required to trigger the onset of infestation. Additionally, the maximum temperature must exceed tfly.
tfly	Minimum temperature that beetles need to fly.
dd_development_base	Base temperature to calculate degree days for development.
dd_total_dev	Degree days that are required for a generation to fully develop
dev_start, dev_end	Share in total development when the egg development starts and the juvenile beetle's development ends respectively. Usable if the development below/above these thresholds should account for mating, oviposition etc.
dev_sister_brood	Share in the total development, when a sister brood will be established.
dev_mortal_min, dev_mortal_max	The beetles are considered to be in white stages (egg, larva, pupa) if their development exceeds dev_mortal_min and subceeds dev_mortal_max. During these stages, the beetles could die due to a mortality event. NULL means that no lower/upper threshold is defined.
func_ftmin, func_ftmean, func_ftmax	Functions to caclulate the air temperature in forest stands (see Ogris et al. 2019, equations 1 - 3). Each parameter will be passed as SpatRaster: <ul style="list-style-type: none"> • tmin: min air temperature • tmean: mean air temperature • tmax: maximum air temperature
func_btmin, func_btmean, func_btmax	Functions to caclulate the bark temperature (see Ogris et al. 2019, equations 4 - 6). Each parameter will be passed as SpatRaster: <ul style="list-style-type: none"> • ftmin: min air temperature in forest stands • ftmean: mean air temperature in forest stands • ftmax: maximum air temperature in forest stands

dt_low, dt_up, topt, tmax, alpha, beta, gamma
 Parameters to calculate the effective bark temperature (see Ogris et al. 2020, equations A.7 - A.9).

model_end_date Date when the model ends (no further development will be modeled).

daylength_dia When the daylength falls below this threshold, diapause will be initiated.

mortality_date Date when all white stages (egg, larva, pupa) die.

Details

In barrks, `model()` is used to customize a model. The following code illustrates which parameters are available for CHAPY and specifies their default values.

```
model("chapy",

      # ==== onset ====

      dd_onset_start_date = '03-09',
      dd_onset_base = 7.4,
      dd_onset_threshold = 216.5,

      # ==== onset + development ====

      tfly = 15.6,

      # ==== development ====

      dd_development_base = 7.4,
      dd_total_dev = 635.4,
      dev_start = 0,
      dev_end = 1,
      dev_sister_brood = 0.5,
      dev_mortal_min = NULL,
      dev_mortal_max = 0.8,

      func_ftmin = function(tmin) { 1.44 + 0.82 * tmin },
      func_ftmean = function(tmean) { 0.50 + 0.81 * tmean },
      func_ftmax = function(tmax) { 1.03 + 0.86 * tmax },

      func_btmin = function(atmin) { 0.56 + 0.99 * atmin },
      func_btmean = function(atmean) { -0.48 + 1.03 * atmean },
      func_btmax = function(atmax) { 0.03 + 0.99 * atmax },

      dt_low = 7.4,
      dt_up = 39.4,
      topt = 30,
      tmax = 41.97,
      alpha = 0.031,
      beta = 5.3,
```



```

    gamma = 1.25,
    model_end_date = '12-31',
    # ==== diapause ====
    daylength_dia = 13.6,
    # ==== mortality ====
    mortality_date = '12-31'
  )

```

References

Ogris N, Ferlan M, Hauptman T, Pavlin R, Kavčič A, Jurc M, De Groot M (2019). “RITY–A phenology model of *Ips typographus* as a tool for optimization of its monitoring.” *Ecological Modelling*, **410**, 108775. doi:10.1016/j.ecolmodel.2019.108775.

Ogris N, Ferlan M, Hauptman T, Pavlin R, Kavčič A, Jurc M, de Groot M (2020). “Sensitivity analysis, calibration and validation of a phenology model for *Pityogenes chalcographus* (CHAPY).” *Ecological Modelling*, **430**, 109137. ISSN 0304-3800, doi:10.1016/j.ecolmodel.2020.109137.

See Also

[model\(\)](#), [phenology\(\)](#), [model.chapy.apply](#)

Other model customizations: [model.bso.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips.customize](#), [model.phenips_clim.customize](#), [model.rity.customize](#)

model.joensson.apply *Use the Jönsson model*

Description

This page describes the usage of the the Jönsson model with [phenology\(\)](#). The model specific inputs are listed and its basic functionality is explained. The Jönsson model was published by Jönsson et al. (2011) and parametrized for *Ips typographus* in southern Sweden.

Arguments

...	See phenology() for a detailed description of the function.
tmean, tmax	Daily mean/maximum temperatures in °C.
daylength	Length of the day in hours. Can be created with create_daylength_rst() or create_daylength_rst() .
mode	Can be 'fast' (default) or 'slow'. Determines if the lower ('fast') or upper ('slow') limit for the development of generation should be used.

Details

In `barrks`, `phenology()` is used to apply a model. The following code illustrates which inputs are required to apply the Jönsson model and which additional parameters are available.

```
phenology("joensson", ..., tmean, tmax, daylength, mode = 'fast')

# calculate submodels separately
phenology("joensson", ..., .submodels = 'onset', tmax)
phenology("joensson", ..., .submodels = 'diapause', tmax, daylength)
phenology("joensson", ..., .submodels = 'development',
          .onset, .diapause = NULL, .mortality = NULL,
          tmean, mode = 'fast')
```

Value

The function returns a phenology. Look [here](#) to find out how it can be analysed.

Functioning

In the following, the basic functioning of the Jönsson model is explained.

- **Onset:** The onset of swarming is triggered when the degree days of the maximum temperature reach a specific threshold and the maximum temperature exceeds the minimum flight temperature. The onset of infestation is triggered seven days later to account for a pre-oviposition period.
- **Development:** The development progresses proportional to the degree days of the mean temperature. To account for varying sun exposures, two different thermal thresholds are defined that reflect the lower and the upper limit of development. A generation starts swarming when it has finished its development and the maximum temperature exceeds the minimum flight temperature. Seven days later, the development of a new generation starts.
- **Diapause:** The diapause is initiated when the daylength falls below a threshold. It is recommended to adjust the daylength threshold when applying elsewhere (e.g. values from literature). Jönsson et al. (2011) proposes a model to calculate the daylength threshold based on long-term climate data.
- **Mortality:** The Jönsson model does not have a mortality submodel implemented.

Look [here](#) to find out how the model parameters affect the actual calculations and which values are used by default.

References

Jönsson AM, Harding S, Krokene P, Lange H, Åke Lindelöw A, Økland B, Ravn HP, Schroeder LM (2011). "Modelling the potential impact of global warming on *Ips typographus* voltinism and reproductive diapause." *Climatic Change*, **109**, 695-718. doi:10.1007/s1058401100384.

See Also

[model\(\)](#), [phenology\(\)](#), [model.joensson.customize](#)

Other phenology applications: [model.bso.apply](#), [model.chapy.apply](#), [model.lange.apply](#), [model.phenips.apply](#), [model.phenips_clim.apply](#), [model.rity.apply](#)

model.joensson.customize

Customize the Jönsson model

Description

This page describes the parameters that can be used to customize the Jönsson model. The model was developed by Jönsson et al. (2011). Look [here](#) to find out how to apply the model.

Arguments

`dd_onset_start_date` The date, when the degree days start to sum up ('MM-DD').

`dd_onset_base` Base temperature to calculate degree days to trigger the onset.

`dd_onset_threshold` Degree days that are required to trigger the onset of infestation. Additionally, the maximum temperature must exceed `tfly`.

`tfly` Minimum temperature that beetles need to fly.

`dd_development_base` Base temperature to calculate degree days for development.

`dd_total_dev_lower`, `dd_total_dev_upper` Lower/upper limit of degree days that are required for a generation to fully develop

`dev_start`, `dev_end` Share in total development when the egg development starts and the juvenile beetle's development ends respectively. Usable if the development below/above these thresholds should account for mating, oviposition etc.

`dev_mortal_min`, `dev_mortal_max` The beetles are considered to be in white stages (egg, larva, pupa) if their development exceeds `dev_mortal_min` and subceeds `dev_mortal_max`. During these stages, the beetles could die due to a mortality event. NULL means that no lower/upper threshold is defined.

`model_end_date` Date when the model ends (no further development will be modeled).

`daylength_dia`, `tdia_min` When the daylength falls below `daylength_dia` and the average daily temperature falls below `tdia_min`, diapause will be initiated. The default value for the critical daylength was set to 19.3 hours according to Schroeder and Dalin (2017) who examined the photoperiodic diapause induction in Sweden. If the model is used for other regions, this value should be adjusted.

Details

In `barrks`, `model()` is used to customize a model. The following code illustrates which parameters are available for the Jönsson model and specifies their default values.

```
model("joensson",
      # ==== onset ====
      dd_onset_start_date = '01-01',
      dd_onset_base = 5,
      dd_onset_threshold = 120,
      # ==== onset + development ====
      tfly = 20,
      # ==== development ====
      dd_development_base = 5,
      dd_total_dev_lower = 625,
      dd_total_dev_upper = 750,
      dev_start = 0,
      dev_end = 1,
      dev_mortal_min = NULL,
      dev_mortal_max = NULL,
      model_end_date = '12-31',
      # ==== diapause ====
      daylength_dia = 19.3,
      tdia_min = 15
    )
```

References

Jönsson AM, Harding S, Krokene P, Lange H, Åke Lindelöw A, Økland B, Ravn HP, Schroeder LM (2011). “Modelling the potential impact of global warming on *Ips typographus* voltinism and reproductive diapause.” *Climatic Change*, **109**, 695-718. doi:10.1007/s1058401100384.

Schroeder M, Dalin P (2017). “Differences in photoperiod-induced diapause plasticity among different populations of the bark beetle *Ips typographus* and its predator *Thanasimus formicarius*.” *Agricultural and Forest Entomology*, **19**(2), 146-153. doi:10.1111/afe.12189.

See Also

[model\(\)](#), [phenology\(\)](#), [model.joensson.apply](#)

Other model customizations: [model.bso.customize](#), [model.chapy.customize](#), [model.lange.customize](#), [model.phenips.customize](#), [model.phenips_clim.customize](#), [model.rity.customize](#)

model.lange.apply *Use the Lange model*

Description

This page describes the usage of the the Lange model with [phenology\(\)](#). The model specific inputs are listed and its basic functionality is explained. The model was published by Lange et al. (2008) for *Ips typographus*.

Arguments

tmin, tmean, tmax

Daily minimum/mean/maximum temperatures in °C.

.submodels, .onset, .diapause, .mortality, ...

See [phenology\(\)](#) for a detailed description of the function.

Details

In barrks, [phenology\(\)](#) is used to apply a model. The following code illustrates which inputs are required to apply the Lange model and which additional parameters are available.

```
phenology("lange", ..., tmin, tmean, tmax)

# calculate submodels separately
phenology("lange", ..., .submodels = 'onset', tmean, tmax)
phenology("lange", ..., .submodels = 'mortality', tmin)
phenology("lange", ..., .submodels = 'development',
          .onset, .diapause = NULL, .mortality = NULL,
          tmean, tmax)
```

Value

The function returns a phenology. Look [here](#) to find out how it can be analysed.

Functioning

In the following, the basic functioning of the Lange model is explained.

- **Onset:** The onset of swarming is triggered when the degree days of the maximum temperature reach a specific threshold and the maximum temperature exceeds the minimum flight temperature according to Annala (1969).
- **Development:** The development is calculated using stage-specific temperature sums and thresholds (Wermelinger and Seifert 1998). A new generation starts its development when the last generation finished its development and the maximum temperature exceeds the minimum flight temperature.

- **Diapause:** The Lange model does not have a diapause submodel implemented.
- **Mortality:** White stages (egg to pupa) die when the minimum temperature falls below a specific threshold.

Look [here](#) to find out how the model parameters affect the actual calculations and which values are used by default.

References

Annala E (1969). “Influence of temperature upon the development and voltinism of *Ips typographus* L. (Coleoptera, Scolytidae).” *Annales Zoologici Fennici*, **6**(2), 161–208. <http://www.jstor.org/stable/23731366>.

Lange H, Økland B, Krokene P (2008). “To be or twice to be? The life cycle development of the spruce bark beetle under climate change.” In *Unifying Themes in Complex Systems: Proceedings of the Sixth International Conference on Complex Systems*, 251–258. Springer. doi:10.1007/9783540850816_32.

Wermelinger B, Seifert M (1998). “Analysis of the temperature dependent development of the spruce bark beetle *Ips typographus* (L) (Col., Scolytidae).” *Journal of Applied Entomology*, **122**(1-5), 185-191. doi:10.1111/j.14390418.1998.tb01482.x.

See Also

[model\(\)](#), [phenology\(\)](#), [model.lange.customize](#)

Other phenology applications: [model.bso.apply](#), [model.chapy.apply](#), [model.joensson.apply](#), [model.phenips.apply](#), [model.phenips_clim.apply](#), [model.rity.apply](#)

model.lange.customize *Customize the Lange model*

Description

This page describes the parameters that can be used to customize Lange. The model was developed by Lange et al. (2008). Look [here](#) to find out how to apply the model.

Arguments

dd_onset_start_date	The date, when the degree days start to sum up ('MM-DD').
dd_onset_base	Base temperature to calculate degree days to trigger the onset.
dd_onset_threshold	Degree days that are required to trigger the onset of infestation. Additionally, the maximum temperature must exceed tfly.
tfly	Minimum temperature that beetles need to fly.

dd_base_stages Base temperatures to calculate degree days for the different stages in the following order: egg, larva, pupa, juvenile adult.

dd_threshold_stages Thermal thresholds for the different stages in the following order: egg, larva, pupa, juvenile adult.

model_end_date Date when the model ends (no further development will be modeled).

first_lethal_date Date before which no mortality will be modeled.

tlethal Temperature threshold below which white stages (egg, larva, pupa) will die.

Details

In barrks, `model()` is used to customize a model. The following code illustrates which parameters are available for the Lange model and specifies their default values.

```
model("lange",
      # ==== onset ==== #
      dd_onset_start_date = '01-01',
      dd_onset_base = 5,
      dd_onset_threshold = 110,
      # ==== onset + development ====
      tfly = 19.5,
      # ==== development ==== #
      dd_base_stages = c(10.6, 8.2, 9.9, 3.2),
      dd_threshold_stages = c(51.8, 204.4, 57.7, 238.5),
      model_end_date = '12-31',
      # ==== mortality ==== #
      first_lethal_date = '09-01',
      tlethal = 0
    )
```

References

Lange H, Økland B, Krokene P (2008). "To be or twice to be? The life cycle development of the spruce bark beetle under climate change." In *Unifying Themes in Complex Systems: Proceedings of the Sixth International Conference on Complex Systems*, 251–258. Springer. doi:10.1007/9783-540850816_32.

See Also

[model\(\)](#), [phenology\(\)](#), [model.lange.apply](#)

Other model customizations: [model.bso.customize](#), [model.chapy.customize](#), [model.joensson.customize](#), [model.phenips.customize](#), [model.phenips_clim.customize](#), [model.rity.customize](#)

model.phenips.apply *Use PHENIPS*

Description

This page describes the usage of PHENIPS with [phenology\(\)](#). The model specific inputs are listed and its basic functionality is explained. PHENIPS was published by Baier et al. (2007) and parametrized at the Kalkalpen National Park in Austria for *Ips typographus*.

Arguments

...	See phenology() for a detailed description of the function.
tmean, tmax	Daily mean/maximum temperatures in °C.
rad	Daily radiation in W * h / m ² .
daylength	Length of the day in hours. Can be created with create_daylength_rst() or create_daylength_df() .
exposure	Specifies the sun exposure. Can be 'sunny' (default) or 'shaded'.
sister_broods	Set FALSE if sister broods should not be calculated.

Details

In barrks, [phenology\(\)](#) is used to apply a model. The following code illustrates which inputs are required to apply PHENIPS and which additional parameters are available.

```
phenology("phenips", ..., tmean, tmax, rad, daylength,
          exposure = 'sunny', sister_broods = TRUE)

# calculate submodels separately
phenology("phenips", ..., .submodels = 'onset', tmax)
phenology("phenips", ..., .submodels = 'diapause', daylength)
phenology("phenips", ..., .submodels = 'mortality', tmax)
phenology("phenips", ..., .submodels = 'development',
          .onset, .diapause = NULL, .mortality = NULL,
          tmean, tmax, rad,
          exposure = 'sunny', sister_broods = TRUE)
```

Value

The function returns a phenology. Look [here](#) to find out how it can be analysed.

Functioning

In the following, the basic functioning of PHENIPS is explained.

- **Onset:** The onset is triggered when the degree days of the maximum temperature reach a specific threshold and the maximum temperature exceeds the minimum flight temperature.
- **Development:** The beetles develop according to a slightly modified version of the optimum curve described by Wermelinger and Seifert (1998) depending on the bark temperature. The bark temperature is modeled based on mean and maximum temperature, global radiation and sun exposure. A new generation will emerge when the last generation is fully developed and the maximum temperature exceeds the minimum flight temperature.
- **Diapause:** The diapause is initiated when the daylength falls below a threshold.
- **Mortality:** White stages (egg to pupa) die on a fixed date.

Look [here](#) to find out how the model parameters affect the actual calculations and which values are used by default.

References

Baier P, Pennerstorfer J, Schopf A (2007). “PHENIPS—A comprehensive phenology model of *Ips typographus* (L.)(Col., Scolytinae) as a tool for hazard rating of bark beetle infestation.” *Forest Ecology and Management*, **249**(3), 171–186. doi:10.1016/j.foreco.2007.05.020.

Wermelinger B, Seifert M (1998). “Analysis of the temperature dependent development of the spruce bark beetle *Ips typographus* (L) (Col., Scolytidae).” *Journal of Applied Entomology*, **122**(1-5), 185-191. doi:10.1111/j.14390418.1998.tb01482.x.

See Also

[model\(\)](#), [phenology\(\)](#), [model.phenips.customize](#)

Other phenology applications: [model.bso.apply](#), [model.chapy.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips_clim.apply](#), [model.rity.apply](#)

model.phenips.customize

Customize PHENIPS

Description

This page describes the parameters that can be used to customize PHENIPS. The model was developed by Baier et al. (2007). Look [here](#) to find out how to apply the model.

Arguments

dd_onset_start_date	The date, when the degree days start to sum up ('MM-DD').
dd_onset_base	Base temperature to calculate degree days to trigger the onset.
dd_onset_threshold	Degree days that are required to trigger the onset of infestation. Additionally, the maximum temperature must exceed tfly.
tfly	Minimum temperature that beetles need to fly.
dd_development_base	Base temperature to calculate degree days for calculating the beetles development.
dd_total_dev	Degree days that are required for a generation to fully develop
dev_start, dev_end	Share in total development when the egg development starts and the juvenile beetle's development ends respectively. Usable if the development below/above these thresholds should account for mating, oviposition etc.
dev_sister_brood	Share in the total development when a sister brood will be established.
dev_mortal_min, dev_mortal_max	The beetles are considered to be in white stages (egg, larva, pupa) if their development exceeds dev_mortal_min and subceeds dev_mortal_max. During these stages, the beetles could die due to a mortality event. NULL means that no lower/upper threshold is defined.
topt	Temperature for optimal development.
tlow, tup	Temperature below/above which no development happens.
func_btmean, func_btmax, func_btdiff	Functions to calculate the effective bark temperature (see Baier et al. 2007, equations A.3 to A5). Each parameter will be passed as SpatRaster: <ul style="list-style-type: none"> • tmean: mean air temperature • tmax: maximum air temperature • rad: radiation • btmax: maximum bark temperature
model_end_date	Date when the model ends (no further development will be modeled).
daylength_dia	When the daylength falls below this threshold, diapause will be initiated.
mortality_date	Date when all white stages (egg, larva, pupa) die.

Details

In barrks, `model()` is used to customize a model. The following code illustrates which parameters are available for PHENIPS and specifies their default values.

```
model("phenips",
      # ==== onset ====
```

```

dd_onset_start_date = '04-01',
dd_onset_base = 8.3,
dd_onset_threshold = 140,

# ==== onset + development ====

tfly = 16.5,

# ==== development ====

dd_development_base = 8.3,
dd_total_dev = 557,
dev_start = 0,
dev_end = 1,
dev_sister_brood = 0.5,
dev_mortal_min = NULL,
dev_mortal_max = 0.6,

topt = 30.4,
tlow = 8.3,
tup = 38.9,

func_btmean = function(tmean, rad) {
  -0.173 + 0.0008518 * rad + 1.054 * tmean
},
func_btmax = function(tmax, rad) {
  1.656 + 0.002955 * rad + 0.534 * tmax + 0.01884 * tmax ^ 2
},
func_btdiff = function(btmax) {
  (-310.667 + 9.603 * btmax) / 24
},

model_end_date = '10-31',

# ==== diapause ====

daylength_dia = 14.5,

# ==== mortality ====

mortality_date = '10-31'
)

```

References

Baier P, Pennerstorfer J, Schopf A (2007). "PHENIPS—A comprehensive phenology model of *Ips typographus* (L.)(Col., Scolytinae) as a tool for hazard rating of bark beetle infestation." *Forest Ecology and Management*, **249**(3), 171–186. doi:10.1016/j.foreco.2007.05.020.

See Also

[model\(\)](#), [phenology\(\)](#), [model.phenips.apply](#)

Other model customizations: [model.bso.customize](#), [model.chapy.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips_clim.customize](#), [model.rity.customize](#)

model.phenips_clim.apply

Use PHENIPS-Clim

Description

This page describes the usage of PHENIPS-Clim with [phenology\(\)](#). The model specific inputs are listed and its basic functionality is explained. PHENIPS-Clim is not published yet. This manual will be updated when a publication is available. It was parametrized for *Ips typographus* in southern Germany.

Arguments

...	See phenology() for a detailed description of the function. See phenology() for details.
tmin, tmean, tmax	Daily minimum/mean/maximum temperatures in °C. tmin is optional. If available it will be used to calculate the temperature amplitude. If not, (tmax - tmean) * 2 will be used as amplitude.
rad	Daily radiation in W * h / m ² .
daylength	Length of the day in hours. Can be created with create_daylength_rst() or create_daylength_rst() .
sister_broods	Set FALSE to disable the calculation of sister broods.
scenario	Choose a scenario to use a suitable combination of parameters for specific situations. The scenario defines a default value for each value that can be overwritten by specifying a value for the respective parameter. The following scenarios are available: <ul style="list-style-type: none"> • mean: <code>list(exposure = 'sunny', onset_mode = 0.5, diapause_mode = 'photoperiodic', oviposition_mode = 0.5)</code> • max: <code>list(exposure = 'sunny', onset_mode = 0.1, diapause_mode = 'thermal', oviposition_mode = 0.1)</code>
exposure	Specifies the sun exposure. Can be 'sunny' (default) or 'shaded'.
onset_mode	Share of beetles that are already infesting trees necessary to trigger the onset. Must be 0.1, 0.5 or 0.9 if not customized.
oviposition_mode	Share of beetles that should have finished oviposition to trigger the beginning of the development. Must be 0.1, 0.5 or 0.9 if not customized.
diapause_mode	Determines how the diapause is initiated. Can be one of the following options:

- 'photoperiodic': The diapause is initiated when the daylength falls below a specific threshold.
- 'thermal': The diapause is initiated by a logistic model that depends on the daylength and the maximum temperature.

Share of beetles that already stopped reproducing necessary to trigger the diapause. Must be thermal or 'photoperiodic' if not customized. If 'photoperiodic' is chosen, the diapause is controlled by a daylength threshold (see parameter `daylength_dia` [here](#)).

Details

In `barrks`, `phenology()` is used to apply a model. The following code illustrates which inputs are required to apply PHENIPS-Clim and which additional parameters are available.

```
phenology("phenips-clim", ..., tmin, tmean, tmax, rad, daylength,
         sister_broods = TRUE, scenario = 'max', exposure = NULL,
         onset_mode = NULL, oviposition_mode = NULL, diapause_mode = NULL)

# calculate submodels separately
phenology("phenips-clim", ..., .submodels = 'onset', tmax, scenario = 'max', onset_mode = NULL)
phenology("phenips-clim", ..., .submodels = 'diapause', tmax, daylength, scenario = 'max', diapause_mode = NULL)
phenology("phenips-clim", ..., .submodels = 'mortality', tmin)
phenology("phenips-clim", ..., .submodels = 'development',
         .onset, .diapause = NULL, .mortality = NULL,
         tmin, tmean, tmax, rad, sister_broods = TRUE,
         scenario = 'max', exposure = NULL, oviposition_mode = NULL)
```

Functioning

In the following, the basic functioning of PHENIPS-Clim is explained.

- **Onset:** A base onset is triggered by a logistic model that relates to the maximum temperature and the respective degree days. Beginning from the base onset, a specific level of degree days (depending on the share of individuals that should be accounted for) and maximum air temperature must be reached to trigger the actual onset.
- **Development:** While the bark temperature and the emergence of new generations are determined according to [PHENIPS](#), the calculation of the beetles' development rates is refined. Rather than implying a constant development within a day, temperature fluctuations are incorporated by taking the daily temperature amplitude into account. Additionally, the first part of development represents the pre-oviposition period and will not appear in the resulting output.
- **Diapause:** The diapause can be initiated due to the photoperiod according to [PHENIPS](#) or by a logistic model that depends on the daylength and the maximum temperature and accounts for beetles that reproduce even on shorter days if the temperatures are favorable. In the second case, PHENIPS-Clim detects a reproductive arrest, due to adverse abiotic parameters, and not an actual diapause as the output can be adjusted, if conditions improve and allow for further reproduction later in the season.
- **Mortality:** White stages (egg to pupa) die when the minimum temperature falls below a specific threshold.

Look [here](#) to find out how the model parameters affect the actual calculations and which values are used by default.

See Also

[model\(\)](#), [phenology\(\)](#), [model.phenips_clim.customize](#)

Other phenology applications: [model.bso.apply](#), [model.chapy.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips.apply](#), [model.rity.apply](#)

model.phenips_clim.customize

Customize PHENIPS-Clim

Description

In barrks, [model\(\)](#) is used to customize a model. Here, the parameters are described that can be used to customize PHENIPS-Clim. The model is currently unpublished. This manual will be updated as soon as a publication is available. Look [here](#) to find out how to apply the model.

Arguments

dd_onset_start_date	The date, when the degree days start to sum up ('MM-DD').
dd_onset_base	Base temperature to calculate degree days to trigger the onset.
onset_func	Function with the SpatRasters tmax (maximum temperature) and dd_tmax (degree days of maximum temperature) as parameters. The function should return TRUE when the base onset is triggered. See onset_add_dd for the actual onset of infestation.
onset_add_dd	Vector of options to calculate the actual onset of infestation. The vector should be named after the share of beetles that already started breeding when the onset is triggered (choose an option via phenology(..., onset_mode = [option]) when applying the model). The values specify the degree days that are required starting at the first positive return value of onset_func.
tfly	Minimum temperature that beetles need to fly.
dd_total_dev	Degree days that are required for a generation to fully develop
dev_oviposition	Named numeric vector of shares in the total development when the oviposition is finished. The vector should be named after the share of beetles that should be taken into account (choose an option via phenology(..., oviposition_mode = [option]) when applying the model).
dev_end	Share in total development when the juvenile beetle's development ends. Usable if the development above this threshold should account for mating, oviposition etc.
dev_sister_brood	Share in the total development, when a sister brood will be established.

dev_mortal_min, dev_mortal_max	The beetles are considered to be in white stages (egg, larva, pupa) if their development exceeds dev_mortal_min and subceeds dev_mortal_max. During these stages, the beetles could die due to a mortality event. NULL means that no lower/upper threshold is defined.
topt	Temperature for optimal development.
func_btmean, func_btmax, func_btdiff	Functions to calculate the bark temperatures (see Baier et al. 2007, equations A.3 to A.5). Each parameter will be passed as SpatRaster: <ul style="list-style-type: none"> • tmean: mean air temperature • tmax: maximum air temperature • rad: radiation • btmax: maximum bark temperature
dev_rates	Data frame that specifies the development rates per day depending on the mean temperature and the temperature amplitude. Column names are the mean temperatures and row names the temperature amplitudes both with one decimal place. base onset (see onset_func) to trigger the actual onset.
model_end_date	Date when the model ends (no further development will be modeled).
first_diapause_date	Date before which an initiation of the diapause is impossible ('MM-DD').
diapause_thermal_func	Function to calculate the initiation of the diapause if the model was applied using phenology(..., diapause_mode = 'thermal'). The diapause will be initiated the last time when the function returns TRUE.
daylength_dia	When the daylength falls below this threshold, diapause will be initiated if the model was applied using phenology(..., diapause_mode = 'photoperiodic').
tlethal	Temperature threshold below which white stages will die.

Details

In `barrks`, `model()` is used to customize a model. The following code illustrates which parameters are available for PHENIPS-Clim and specifies their default values.

```
model("phenips-clim",

      # ==== onset ====

      dd_onset_start_date = '03-01',
      dd_onset_base = 12,
      onset_func = function(tmax, dd_tmax) {
        0.564071 * tmax + 0.006434 * dd_tmax - 12.37046 > 0
      },
      onset_add_dd = c('0.1' = 0, '0.5' = 90, '0.9' = 190),

      # ==== development ====
```

```

tfly = 16.5,

dd_total_dev = 557,

dev_oviposition = c('0.1' = 0.1,
                    '0.5' = 0.15,
                    '0.9' = 0.26),

dev_end = 1,
dev_sister_brood = 0.3,

dev_mortal_min = NULL,
dev_mortal_max = 0.6,

topt = 30.4,

func_btmean = function(tmean, rad) {
  -0.173 + 0.0008518 * rad + 1.054 * tmean
},
func_btmax = function(tmax, rad) {
  1.656 + 0.002955 * rad + 0.534 * tmax + 0.01884 * tmax ^ 2
},
func_btdiff = function(tmax) {
  (-310.667 + 9.603 * tmax) / 24
},

# dev_rates too large to show here, type `params('phenips-clim')$dev_rates`
# to get the dev_rates that are used by default
# dev_rates = matrix(...),

model_end_date = '12-31',

# ==== diapause ====

first_diapause_date = '08-12',
diapause_thermal_func = function(daylength, tmax) {
  0.8619156 * daylength + 0.5081128 * tmax - 23.63691 > 0
},
daylength_dia = 14.5,

# ==== mortality ====

tlethal = -5
)

```

References

Baier P, Pennerstorfer J, Schopf A (2007). “PHENIPS—A comprehensive phenology model of *Ips typographus* (L.) (Col., Scolytinae) as a tool for hazard rating of bark beetle infestation.” *Forest Ecology and Management*, **249**(3), 171–186. doi:10.1016/j.foreco.2007.05.020.

See Also

[model\(\)](#), [phenology\(\)](#), [model.phenips_clim.apply](#)

Other model customizations: [model.bso.customize](#), [model.chapy.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips.customize](#), [model.rity.customize](#)

model.rity.apply	<i>Use RITY</i>
------------------	-----------------

Description

This page describes the usage of RITY with [phenology\(\)](#). The model specific inputs are listed and its basic functionality is explained. RITY (also called RITY-2) was published by Ogris et al. (2019) and parametrized for *Ips typographus* in Slovenia.

Arguments

tmin, tmean, tmax	Daily minimum/mean/maximum temperatures in °C. For the development sub-model, the parameter that is obligatory depends on mode.
daylength	Length of the day in hours. Can be created with create_daylength_rst() or create_daylength_rst() .
mode	Specifies which temperature should be used to calculate the development. Can be min, mean or max.
.submodels, .onset, .diapause, .mortality, ...	See phenology() for a detailed description of the function.

Details

In `barrks`, [phenology\(\)](#) is used to apply a model. The following code illustrates which inputs are required to apply RITY and which additional parameters are available.

```
phenology("rity", ..., tmin = NULL, tmean = NULL, tmax, daylength, mode = 'max')

# calculate submodels separately
phenology("rity", ..., .submodels = 'onset', tmax)
phenology("rity", ..., .submodels = 'diapause', daylength)
phenology("rity", ..., .submodels = 'mortality', tmax)
phenology("rity", ..., .submodels = 'development',
          .onset, .diapause = NULL, .mortality = NULL,
          tmin = NULL, tmean = NULL, tmax = NULL, mode = 'max')
```

Value

The function returns a phenology. Look [here](#) to find out how it can be analysed.

Functioning

In the following, the basic functioning of RITY is explained.

- **Onset:** See [PHENIPS](#).
- **Development:** Based on [PHENIPS](#) with a few modifications:
 - The optimum curve is calculated according to Wermelinger and Seifert (1998) without simplification.
 - The minimum, mean or maximum bark temperature can be used to calculate the development. These temperatures depend only on the respective air temperatures.
- **Diapause:** See [PHENIPS](#).
- **Mortality:** See [PHENIPS](#).

Look [here](#) to find out how the model parameters affect the actual calculations and which values are used by default.

References

Ogris N, Ferlan M, Hauptman T, Pavlin R, Kavčič A, Jurc M, De Groot M (2019). “RITY–A phenology model of *Ips typographus* as a tool for optimization of its monitoring.” *Ecological Modelling*, **410**, 108775. doi:10.1016/j.ecolmodel.2019.108775.

Wermelinger B, Seifert M (1998). “Analysis of the temperature dependent development of the spruce bark beetle *Ips typographus* (L) (Col., Scolytidae).” *Journal of Applied Entomology*, **122**(1-5), 185-191. doi:10.1111/j.14390418.1998.tb01482.x.

See Also

[model\(\)](#), [phenology\(\)](#), [model.rity.customize](#)

Other phenology applications: [model.bso.apply](#), [model.chapy.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips.apply](#), [model.phenips_clim.apply](#)

model.rity.customize *Customize RITY*

Description

This page describes the parameters that can be used to customize RITY (also called RITY-2). The model was developed by Ogris et al. (2019). Look [here](#) to find out how to apply the model.

Arguments

dd_onset_start_date

The date, when the degree days start to sum up ('MM-DD').

dd_onset_threshold

Degree days that are required to trigger the onset of infestation. Additionally, the maximum temperature must exceed tfly.

tfly	Minimum temperature that beetles need to fly.
dd_onset_base	Base temperature to calculate degree days for development.
dd_total_dev	Degree days that are required for a generation to fully develop
dev_start, dev_end	Share in total development when the egg development starts and the juvenile beetle's development ends respectively. Usable if the development below/above these thresholds should account for mating, oviposition etc.
dev_sister_brood	Share in the total development, when a sister brood will be established.
dev_mortal_min, dev_mortal_max	The beetles are considered to be in white stages (egg, larva, pupa) if their development exceeds dev_mortal_min and subceeds dev_mortal_max. During these stages, the beetles could die due to a mortality event. NULL means that no lower/upper threshold is defined.
func_ftmean, func_ftmax, func_atdiff	Functions to caclulate the air temperature in forest stands (see Ogris et al. 2019, equations 1 - 3). Each parameter will be passed as SpatRaster: <ul style="list-style-type: none"> • tmin: min air temperature • tmean: mean air temperature • tmax: maximum air temperature
func_btmean, func_btmax, func_btdiff	Functions to caclulate the bark temperature (see Ogris et al. 2019, equations 4 - 6). Each parameter will be passed as SpatRaster: <ul style="list-style-type: none"> • ftmin: min air temperature in forest stands • ftmean: mean air temperature in forest stands • ftmax: maximum air temperature in forest stands
dt_low, dt_up, topt, tmax, alpha, beta, gamma	Parameters to calculate the effective bark temperature (see Ogris et al. 2019, equations 7 - 9).
model_end_date	Date when the model ends (no further development will be modeled).
daylength_dia	When the daylength falls below this threshold, diapause will be initiated.
mortality_date	Date when all white stages (egg, larva, pupa) die.

Details

In barrks, `model()` is used to customize a model. The following code illustrates which parameters are available for RITY and specifies their default values.

```
model("rity",

      # ==== onset ====

      dd_onset_start_date = '03-07',
      dd_onset_base = 8.3,
      dd_onset_threshold = 155.6,
```

```

# ==== onset + development ====

tfly = 14.5,

# ==== development ====

dd_development_base = 8.3,
dd_total_dev = 557,
dev_start = 0,
dev_end = 1,
dev_sister_brood = 0.5,
dev_mortal_min = NULL,
dev_mortal_max = 0.6,

func_ftmin = function(tmin) { 1.44 + 0.82 * tmin },
func_ftmean = function(tmean) { 0.50 + 0.81 * tmean },
func_ftmax = function(tmax) { 1.03 + 0.86 * tmax },

func_btmin = function(ftmin) { 0.56 + 0.99 * ftmin },
func_btmean = function(ftmean) { -0.48 + 1.03 * ftmean },
func_btmax = function(ftmax) { 0.03 + 0.99 * ftmax },

dt_low = 8.3,
dt_up = 38.9,
topt = 30.4,
tmax = 40.9958913,
alpha = 0.02876507,
beta = 3.5922336,
gamma = 1.24657367,

model_end_date = '10-31',

# ==== diapause ====

daylength_dia = 14.5,

# ==== mortality ====

mortality_date = '10-31'
)

```

References

Ogris N, Ferlan M, Hauptman T, Pavlin R, Kavčič A, Jurc M, De Groot M (2019). “RITY–A phenology model of *Ips typographus* as a tool for optimization of its monitoring.” *Ecological Modelling*, **410**, 108775. doi:10.1016/j.ecolmodel.2019.108775.

See Also

[model\(\)](#), [phenology\(\)](#), [model.rity.apply](#)

Other model customizations: [model.bso.customize](#), [model.chapy.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips.customize](#), [model.phenips_clim.customize](#)

model_combine	<i>Combine different (sub-)models</i>
---------------	---------------------------------------

Description

Combine different (sub-)models.

Usage

```
model_combine(...)
```

Arguments

... Phenology models, model names or lists with the keys `model` and `submodels`. In the last case, only the submodels specified are used (one of 'onset', 'diapause', 'mortality' or 'development') of the respective model. If multiple models are supplied for the same submodel, the last one overwrites all others.

Value

A phenology model. Can be passed to [phenology\(\)](#).

See Also

- [model\(\)](#), [phenology\(\)](#)
- Customize (sub-)models: [model.bso.customize](#), [model.phenips.customize](#), [model.rity.customize](#), [model.chapy.customize](#), [model.joensson.customize](#), [model.lange.customize](#), [model.phenips_clim.customize](#)
- Use (sub-)models: [model.bso.apply](#), [model.phenips.apply](#), [model.rity.apply](#), [model.chapy.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips_clim.apply](#)

Examples

```
# combine PHENIPS with the diapause submodel of PHENIPS-Clim
m <- model_combine('phenips',
                  list(model = 'phenips-clim', submodels = 'diapause'))

# calculate phenology
p <- phenology(m, barrks_data(), .quiet = TRUE)

# plot calculated generations
gens <- get_generations_rst(p)
terra::plot(gens)
```

params	<i>Get model parameters</i>
--------	-----------------------------

Description

Get the parameters of a model.

Usage

```
params(m, ...)
```

Arguments

m	Name of the model or the return value of another model()-call.
...	List of parameters to customize the model.

Value

A list.

Examples

```
# print the first parameters of `phenips-clim`  
head(params('phenips-clim'))
```

phenology	<i>Central function to calculate a phenology</i>
-----------	--

Description

Calculate a phenology (or its subparts) with a specific model.

Usage

```
phenology(  
  .model,  
  .data = NULL,  
  .dates = NULL,  
  .win = NULL,  
  .ext = "tif",  
  .onset = NULL,  
  .diapause = NULL,
```

```

    .mortality = NULL,
    .submodels = c("onset", "diapause", "mortality", "development"),
    .setup_only = FALSE,
    .stations = NULL,
    .storage = NULL,
    .quiet = FALSE,
    ...
)

bso_phenology(
  .model = "bso",
  .data = NULL,
  .dates = NULL,
  .win = NULL,
  .ext = "tif",
  .onset = NULL,
  .diapause = NULL,
  .mortality = NULL,
  .submodels = c("onset", "diapause", "mortality", "development"),
  .setup_only = FALSE,
  .stations = NULL,
  .storage = NULL,
  .quiet = FALSE,
  ...
)

```

Arguments

<code>.model</code>	A phenology model or a model name (see model() , model_combine()).
<code>.data</code>	Data that will be passed to the model. It can be one of the following: <ul style="list-style-type: none"> • Character string: The raster data will be loaded from the path specified. The files have to be named like the respective model inputs. • Named list: Each element contains the input data according to its name. • Data frame (station data): Should have the columns <code>date</code> and <code>station</code> (name of the station). Additional columns have to be named like the respective model inputs. • Additionally, data can be passed through the <code>...</code> argument. <p>Look at the model application manuals to find out which inputs are required by a specific model: model.bso.apply, model.phenips.apply, model.rity.apply, model.chapy.apply, model.joensson.apply, model.lange.apply, model.phenips_clim.apply.</p>
<code>.dates</code>	Vector of dates that the data should be restricted to.
<code>.win</code>	SpatExtent to set a window (area of interest) if <code>.data</code> is a path to load the raster data from.
<code>.ext</code>	Extension of the files that should be used if <code>.data</code> is a path to load the raster data from.

<code>.onset, .diapause, .mortality</code>	Pass custom or precalculated phenological events to the model. See create_events to find out how to create events manually. Alternatively, the return value of get_onset_rst() , get_diapause_rst() or get_mortality_rst() could be used (with <code>as_doy = FALSE</code>) to extract the respective phenological event from another phenology. In that case, that phenology must match the temporal and spatial extent of the other inputs.
<code>.submodels</code>	Character vector. Specifies which submodels should be calculated. Can be a subset of <code>c('onset', 'diapause', 'mortality', 'development')</code> .
<code>.setup_only</code>	If TRUE only the inputs will be preprocessed without calculating any submodels. The preprocessed data can be used as input for other phenology() calls and can be accessed via get_input_data() .
<code>.stations</code>	Assign stations to the phenology. See stations_create() for details.
<code>.storage</code>	If set, the path specified here will be used to save the (intermediate) results. If phenology() is called successively with a growing amount of data, the calculations will continue where they stopped. This can save calculation time especially for large raster inputs. Note that this will only work of raster inputs and if <code>terra::sources()</code> is not empty. Otherwise the results of the calculations will be saved but successive calculations are not possible. If no input data is passed, the phenology will be loaded from the storage.
<code>.quiet</code>	If TRUE, messages are suppressed.
<code>...</code>	Parameters that will be passed to the model. Must be named according to the model inputs. See <code>.data</code> for alternative ways to pass data to the model.

Value

A phenology as a list. Look [here](#) to find out how a phenology can be analysed. It is not recommended to access the list elements directly.

Functions

- `bso_phenology()`: As BSO works a bit different than the other models, a separate phenology function is implemented for this model. Note that while the onset and the development submodels are needed to be taken from BSO, the diapause and the mortality submodels are compatible with other models.

The function returns a BSO phenology as a list. Look [here](#) to find out how a BSO phenology can be analysed. It is not recommended to access the list elements directly. To be able to use the functions that are available for phenology objects returned by [phenology\(\)](#), call [bso_translate_phenology\(\)](#).

See Also

[model.bso.apply](#), [model.phenips.apply](#), [model.rity.apply](#), [model.chapy.apply](#), [model.joensson.apply](#), [model.lange.apply](#), [model.phenips_clim.apply](#)

Examples

```
# calculate phenology
p <- phenology('phenips-clim', barrks_data())

# plot calculated generations
gens <- get_generations_rst(p)
terra::plot(gens)
```

plot_development_diagram

Plot a development diagram

Description

A development diagram illustrates the beetles' development of all appearing generations within a year.

Usage

```
plot_development_diagram(
  .phenos,
  .station = prop_stations(.phenos[[1]])[1],
  .generations = NULL,
  .colors = barrks_colors("diagram_lines"),
  .fill = barrks_colors("diagram_fill"),
  .labels = barrks_labels("diagram"),
  .legend_col = TRUE,
  .legend_lty = TRUE,
  .group = TRUE,
  .minmax_only = FALSE,
  .fun_bg = NULL,
  .lty = "solid",
  .lwd = 2,
  .date_split = NULL,
  .date_stop = NULL,
  .lty2 = "dotted",
  .lwd2 = 2,
  .fill2 = NA,
  ...
)
```

Arguments

.phenos List of (named) phenology objects or a single phenology that will be plotted (see [phenology\(\)](#)).

<code>.station</code>	Pass a character vector to choose a station assigned to pheno by its name, or pass a different station. See stations_create() for details.
<code>.generations</code>	Generations that will be shown.
<code>.colors, .fill, .labels</code>	Character vectors. Change the line colors, fill or labels of the generations starting from the first generation followed consecutively by elements for the other generations (including sister broods).
<code>.legend_col, .legend_lty</code>	Manipulate the appearance of the legends for colors and line types. Pass TRUE/FALSE to enable/disable the respective legend. For the customization of the respective legend, a list of parameters for graphics::legend can be passed.
<code>.group</code>	Select the phenology objects that will be used to draw the filling. It can be a character vector of the phenology names, an integer vector of the phenology numbers, or TRUE if all phenology objects should be used.
<code>.minmax_only</code>	If TRUE, only the minimum and the maximum development line will be plotted.
<code>.fun_bg</code>	Function to draw a background.
<code>.lty, .lwd</code>	Use specific line types and line widths. Vectors of the same length as <code>.phenos</code> will assign the values to the respective phenology.
<code>.date_split, .lty2, .lwd2, .fill2</code>	When <code>.date_split</code> is reached, the appearance of the plot will change according to the respective values.
<code>.date_stop</code>	If specified, no data will be plotted after the respective date.
<code>...</code>	Parameters passed to base::plot() .

Value

None

See Also[stations](#)**Examples**

```
# calculate phenology
p <- phenology('phenips-clim', barrks_data('stations'), .quiet = TRUE)

# plot development diagram of the station 'Mannheim'
plot_development_diagram(p, 'Mannheim', .lwd = 4, .legend_lty = FALSE)
```

properties	<i>Get phenology properties</i>
------------	---------------------------------

Description

To examine a phenology, there are different functions to query its properties.

Usage

```
## get the year the phenology was calculated for
prop_year(pheno)

## get all dates that are covered by the phenology
prop_dates(pheno)

## get the first date that is covered by the phenology
prop_first_date(pheno)

## get the last date that is covered by the phenology
prop_last_date(pheno)

## get all hatched generations as numeric vector
prop_hatched_generations(pheno)

## get all hatched filial generations as numeric vector
prop_filial_generations(pheno)

## get all hatched sister broods as numeric vector
prop_sister_broods(pheno)

## get the stations assigned to the phenology
prop_stations(pheno)
```

Arguments

pheno A phenology (see [phenology\(\)](#))

Value

The requested property.

Examples

```
# calculate phenology
p <- phenology('phenips-clim', barrks_data(), .quiet = TRUE)

# print all generations that were hatched
prop_hatched_generations(p)
```

save_phenology	<i>Save/load a phenology</i>
----------------	------------------------------

Description

Saves/loads a phenology to/from a path.

Usage

```
save_phenology(  
  pheno,  
  .storage,  
  .submodels = c("onset", "diapause", "mortality", "development"),  
  .overwrite = FALSE,  
  .ext = ".tif",  
  .quiet = FALSE  
)  
  
load_phenology(  
  .storage,  
  .submodels = c("onset", "diapause", "mortality", "development"),  
  .ext = ".tif",  
  .quiet = FALSE  
)
```

Arguments

pheno	A phenology, calculated with phenology() .
.storage	Path to save/load the phenology.
.submodels	Which submodels should be saved/loaded.
.overwrite	Should an existing storage be overwritten?
.ext	Extension for raster files.
.quiet	If TRUE, messages are suppressed.

Value

- `save_phenology()`: None
- `load_phenology()`: A phenology as a list. Look [here](#) to find out how a phenology can be analysed. It is not recommended to access the list elements directly.

Functions

- `save_phenology()`: Saves a phenology to a path.
- `load_phenology()`: Loads a phenology from a path.

Examples

```
# calculate phenology
p <- phenology('phenips-clim', barrks_data(), .quiet = TRUE)

# choose path to save the phenology
path <- file.path(tempdir(), 'pheno')

# save phenology
save_phenology(p, path, .overwrite = TRUE, .quiet = TRUE)

###

# load phenology from path
p2 <- load_phenology(path, .quiet = TRUE)

# plot generations
gens <- get_generations_rst(p2)
terra::plot(gens)
```

stations_create

Work with stations

Description

In `barrks`, stations are references to specific raster cells. Thus, they can be used to extract point-related data from a phenology. Look [here](#) to find out which station-based functions are available to analyse a phenology.

Usage

```
stations_create(station_names, cells)
```

```
stations_assign(pheno, stations)
```

```
stations_names(stations)
```

```
stations_cells(stations)
```

Arguments

<code>station_names</code>	Character vector that specifies the names of the stations.
<code>cells</code>	Numbers of the cells that should be represented by the stations.
<code>pheno</code>	A phenology (see phenology())
<code>stations</code>	Stations created with <code>stations_create()</code> or obtained by prop_stations() .

Value

- `stations_create()`: A named numeric vector.
- `stations_assign()`: A phenology object (see `phenology()`).
- `stations_names()`: A character vector.
- `stations_cells()`: A numeric vector.

Functions

- `stations_create()`: Create stations.
- `stations_assign()`: Assign stations to a phenology. Returns the phenology that was passed with respective stations assigned.
- `stations_names()`: Get the names of stations.
- `stations_cells()`: Get the raster cells of stations.

Examples

```
# calculate phenology
p <- phenology('phenips-clim', barrks_data(), .quiet = TRUE)

# create stations and assign them to the phenology object
stations <- stations_create(c('station a', 'station b'),
                           c(234, 345))
p <- stations_assign(p, stations)

# plot the development of 'station b'
plot_development_diagram(p, 'station b', .lwd = 4, .legend_lty = FALSE)
```

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