

# Package ‘pep725’

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**Title** Pan-European Phenological Data Analysis

**Version** 1.1.0

**Description** Provides a framework for quality-aware analysis of ground-based phenological data from the PEP725 Pan-European Phenology Database (Templ et al. (2018) <[doi:10.1007/s00484-018-1512-8](https://doi.org/10.1007/s00484-018-1512-8)>; Templ et al. (2026) <[doi:10.1111/nph.70869](https://doi.org/10.1111/nph.70869)>) and similar observation networks. Implements station-level data quality grading, outlier detection, phenological normals (climate baselines), anomaly detection, elevation and latitude gradient estimation with robust regression, spatial synchrony quantification, partial least squares (PLS) regression for identifying temperature-sensitive periods, and sequential Mann-Kendall trend analysis. Supports data import from PEP725 files, conversion of user-supplied data, and downloadable synthetic datasets for teaching without barriers of registration. All analysis outputs provide 'print', 'summary', and 'plot' methods. Interactive spatial visualization is available via 'leaflet'.

**License** GPL-3

**Encoding** UTF-8

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---

add_country	<i>Add Country Information to a Dataset Based on Latitude/Longitude</i>
-------------	---

---

## Description

This function adds a new column `country` by performing a fast spatial join between point coordinates (`lat`, `lon`) and world country polygons from **rnaturalearth**. Geometry is handled via **sf**.

## Usage

```
add_country(dt, keep_geometry = FALSE)
```

## Arguments

`dt` A `data.frame` or `data.table` containing columns `lat` and `lon`.

`keep_geometry` Logical. If `TRUE`, the `sf` geometry column is kept. Default is `FALSE`.

## Details

The input data must contain numeric columns named `lat` and `lon` in WGS84 (EPSG:4326). Rows with NA coordinates receive `country = NA`.

After assigning countries, Germany is automatically split into "Germany-North" and "Germany-South" using a latitude threshold:

- `lat >= 50` → "Germany-North"
- `lat < 50` → "Germany-South"

If the input already contains a `country` column it is removed and replaced by the spatially derived one.

## Value

A `data.table` with a new column `country`.

## Author(s)

Matthias Templ

**Examples**

```
small <- data.table(  
  lat = c(48.2, 51.0, 47.5),  
  lon = c(16.3, 10.1, 8.5)  
)  
  
pep_with_country <- add_country(small)  
pep_with_country
```

---

`add_daylength`*Add Daylength to Phenological Data*

---

**Description**

Adds a daylength column to a phenological dataset based on the observation day and station latitude.

**Usage**

```
add_daylength(pep)
```

**Arguments**

`pep` A pep object or data.table with day and lat columns.

**Value**

The input data with an added daylength column (hours).

**Author(s)**

Matthias Templ

**See Also**

[calc\\_daylength](#) for the underlying calculation

**Examples**

```
pep <- pep_download()  
pep <- add_daylength(pep)  
head(pep[, .(day, lat, daylength)])
```

---

as.pep *Coerce to PEP725 Data Object*

---

**Description**

Coerce to PEP725 Data Object

**Usage**

```
as.pep(x, ...)
```

**Arguments**

x                    A data.frame or data.table to coerce.  
 ...                 Additional arguments (currently unused).

**Value**

An object of class pep.

**Author(s)**

Matthias Templ

---

bbch\_description *Get BBCH Phase Description*

---

**Description**

Returns human-readable descriptions for BBCH phenological codes.

**Usage**

```
bbch_description(codes, na.rm = TRUE, sort = TRUE)
```

**Arguments**

codes                Integer vector of BBCH codes.  
 na.rm                Logical. If TRUE (default), removes NA codes and codes without a known description from the output.  
 sort                 Logical. If TRUE (default), sorts the output by phase code.

**Value**

A data.frame with columns phase\_id and description, ordered by phase code (if sort = TRUE).

**Author(s)**

Matthias Templ

**Examples**

```
bbch_description(c(60, 65, 100))
bbch_description(c(100, 60, 65, NA), na.rm = TRUE, sort = TRUE)
```

---

calc\_daylength

*Calculate Daylength (Photoperiod)*


---

**Description**

Computes the astronomical daylength (hours of daylight) for a given day of year and latitude. Photoperiod is an important environmental driver of plant phenology.

**Usage**

```
calc_daylength(doy, lat, method = c("brock", "cbm"))
```

**Arguments**

doy	Integer or numeric vector. Day of year (1-365/366).
lat	Numeric. Latitude in decimal degrees. Positive for Northern Hemisphere, negative for Southern Hemisphere.
method	Character. Daylength model: "brock" (Default) Brock (1981) / Spencer (1971) simple declination model: $\delta = 23.45^\circ \cdot \sin(2\pi(284 + \text{DOY})/365)$ . Fast and adequate at mid-latitudes; differs from CBM by up to a few minutes/day near the poles. "cbm" Forsythe et al. (1995) Climate-Budget-Model formulation. Includes the eccentricity correction of the Earth's orbit and is generally the most accurate of the four models Forsythe et al. compared.

**Details**

Both methods assume a "flat horizon" (sunrise/sunset when the geometric centre of the sun crosses the horizon) and do not account for atmospheric refraction, elevation, or local topography.

At polar latitudes ( $|lat| > 66.5^\circ$ ), continuous daylight or darkness may occur around solstices.

**Value**

A list with components:

**daylength** Daylength in hours

**declination** Solar declination angle in degrees

If doy is a vector, returns a data.frame with these columns.

### Phenological Relevance

Many phenological events are triggered by photoperiod thresholds:

- Spring bud burst often requires both temperature accumulation and minimum daylength
- Autumn leaf senescence may be triggered by shortening days
- Flowering in many species is photoperiod-dependent

### Author(s)

Matthias Templ

### References

Spencer, J. W. (1971). Fourier series representation of the position of the sun. *Search* 2(5):172. (Declination formula used by the "brock" method.)

Brock, T. D. (1981). Calculating solar radiation for ecological studies. *Ecological Modelling* 14(1-2):1-19. doi:10.1016/03043800(81)900119

Forsythe, W. C., Rykiel, E. J., Stahl, R. S., Wu, H., and Schoolfield, R. M. (1995). A model comparison for daylength as a function of latitude and day of year. *Ecological Modelling* 80(1):87-95. doi:10.1016/03043800(94)00034F (CBM daylength model used by the "cbm" method.)

### Examples

```
# Daylength at spring equinox (DOY 80) at 50°N
calc_daylength(80, 50)

# Summer solstice at different latitudes
calc_daylength(172, c(30, 45, 60))

# Daylength through the year at 45°N
yearly <- calc_daylength(1:365, 45)
plot(yearly$doy, yearly$daylength, type = "l",
      xlab = "Day of Year", ylab = "Daylength (hours)")
```

---

calc\_max\_daylength      *Maximum Daylength at a Latitude*

---

### Description

Calculates the maximum possible daylength (at summer solstice) for a given latitude.

### Usage

```
calc_max_daylength(lat)
```

**Arguments**

lat                    Numeric. Latitude in decimal degrees.

**Value**

Numeric. Maximum daylength in hours.

**Author(s)**

Matthias Templ

**Examples**

```
# Maximum daylength at different latitudes
calc_max_daylength(c(0, 30, 45, 60, 66.5))
```

---

calc\_thermal\_sum            *Calculate Thermal Sum at Phenological Events*

---

**Description**

Computes the accumulated thermal units (GDD) from a start date to the observed phenological event date. Useful for determining the thermal requirements of different phenophases.

**Usage**

```
calc_thermal_sum(
  pep,
  temp_data,
  t_base = 5,
  t_start = 1,
  by = NULL,
  method = "average"
)
```

**Arguments**

pep                    A pep object or data.table with phenological observations. Must contain year and day columns.

temp\_data            A data.frame or data.table with daily temperature data. Must contain columns: year, doy (day of year), tmin, tmax (or tmean).

t\_base                Numeric. Base temperature (°C). Default 5.

t\_start               Integer. Start day of year for accumulation. Default 1 (Jan 1). Use 60 for March 1, etc.

by	Character vector. Columns to match temperature data to phenology (e.g., "s_id" for station-specific temperatures). Default NULL uses same temperature for all observations in a year.
method	Character. GDD calculation method (see <a href="#">calc_thermal_units</a> ).

### Details

This function joins phenological observations with temperature data and calculates the thermal sum (accumulated GDD) at each observation date. This is useful for:

- Determining thermal requirements of phenophases
- Comparing thermal sums across years, locations, or species
- Validating phenological models

### Value

The input data with an additional thermal\_sum column containing the accumulated GDD from t\_start to the observed DOY.

### Author(s)

Matthias Templ

### See Also

[calc\\_thermal\\_units](#) for the underlying GDD calculation

### Examples

```
# Load phenology data
data(pep_seed)

# Create example temperature data (normally from weather stations/reanalysis)
temp <- data.frame(
  year = rep(2000:2015, each = 365),
  doy = rep(1:365, 16),
  tmin = rnorm(365 * 16, mean = 5, sd = 8),
  tmax = rnorm(365 * 16, mean = 15, sd = 8)
)

# Add thermal sum to phenology data
pep_thermal <- calc_thermal_sum(pep_seed, temp, t_base = 5)
```

---

calc\_thermal\_units      *Calculate Thermal Units (Growing Degree Days)*

---

### Description

Computes accumulated thermal units (growing degree days, GDD) from daily temperature data. Thermal time is a fundamental concept in phenology, representing the heat accumulation that drives plant development.

### Usage

```
calc_thermal_units(
  tmin,
  tmax = NULL,
  t_base = 5,
  t_cap = NULL,
  method = c("average", "modified", "single_sine"),
  cumulative = TRUE,
  na.rm = TRUE
)
```

### Arguments

tmin	Numeric vector of daily minimum temperatures (°C).
tmax	Numeric vector of daily maximum temperatures (°C). If NULL, tmin is treated as daily mean temperature.
t_base	Numeric. Base temperature (°C) below which no development occurs. Default is 5°C, typical for temperate crops.
t_cap	Numeric. Optional upper temperature cap (°C). Temperatures above this are set to t_cap before GDD calculation. Default NULL (no cap). Use ~30°C for crops where high heat doesn't increase development.
method	Character. Calculation method: <b>"average"</b> (Default) $GDD = \max(0, (t_{max} + t_{min})/2 - t_{base})$ <b>"modified"</b> Sets negative daily GDD to 0 before averaging <b>"single_sine"</b> Sine wave approximation for more accurate heat units
cumulative	Logical. If TRUE (default), returns cumulative sum. If FALSE, returns daily values.
na.rm	Logical. Remove NA values in accumulation? Default TRUE.

### Details

Growing Degree Days (GDD) quantify heat accumulation above a base temperature. Plants require a certain thermal sum to reach phenological stages (e.g., flowering typically requires 500-1500 GDD depending on species).

The base temperature represents the minimum temperature for growth:

- Cool-season crops (wheat, barley): 0-5°C
- Warm-season crops (maize, soybean): 10°C
- Fruit trees (apple, cherry): 4-7°C
- Grapevine: 10°C

### Value

Numeric vector of thermal units (degree-days). Same length as input. If `cumulative = TRUE`, returns running sum.

### Methods

**average** Simple average:  $GDD = \max(0, (T_{max} + T_{min})/2 - T_{base})$ . Most common method, used by many agricultural agencies.

**modified** Like average, but  $T_{min}$  and  $T_{max}$  are first bounded to the range from  $T_{base}$  to  $T_{cap}$  before averaging. Prevents negative contributions.

**single\_sine** Approximates the daily temperature curve as a sine wave, computing the mean excess over  $T_{base}$ . More accurate when temperatures cross the base threshold during the day. Implements the Baskerville-Emin (1969) / Snyder (1985) single-sine formula: with  $T_{avg} = (T_{min} + T_{max})/2$ ,  $T_{amp} = (T_{max} - T_{min})/2$ , and  $\theta = \arccos((T_{base} - T_{avg})/T_{amp})$ , the daily GDD in the mixed case ( $T_{min} < T_{base} < T_{max}$ ) is  $GDD = \frac{1}{\pi} [(T_{avg} - T_{base})\theta + T_{amp} \sin(\theta)]$ .

### Author(s)

Matthias Templ

### References

McMaster, G.S., Wilhelm, W.W. (1997). Growing degree-days: one equation, two interpretations. *Agricultural and Forest Meteorology* 87:291-300.

Baskerville, G.L., Emin, P. (1969). Rapid estimation of heat accumulation from maximum and minimum temperatures. *Ecology* 50:514-517.

Snyder, R.L. (1985). Hand calculating degree days. *Agricultural and Forest Meteorology* 35:353-358.

### See Also

[calc\\_thermal\\_sum](#) for computing thermal sum at phenological observations

### Examples

```
# Daily temperatures for a week
tmin <- c(5, 7, 8, 10, 12, 11, 9)
tmax <- c(15, 18, 20, 22, 25, 23, 19)

# Cumulative GDD with base 10°C
calc_thermal_units(tmin, tmax, t_base = 10)
```

```

# Daily GDD values
calc_thermal_units(tmin, tmax, t_base = 10, cumulative = FALSE)

# With upper cap at 30°C
calc_thermal_units(tmin, tmax, t_base = 10, t_cap = 30)

# Using mean temperature only
tmean <- (tmin + tmax) / 2
calc_thermal_units(tmean, t_base = 10)

```

---

get\_phenology\_doys      *Compute CTRL and SCEN Phenology Day-of-Year Values*

---

## Description

This function computes the mean phenological day-of-year (DOY) for two phenophases (e.g. BBCH 65 and 87) for each country, using either a simple warming-shift model or a GDD-based model.

## Usage

```

get_phenology_doys(
  pep,
  phase1,
  phase2,
  genus_name,
  phenology_method = c("robust_shift", "simple_shift", "gdd_cmp6"),
  ctrl_years = 2011:2021,
  scen_years = 2085:2095,
  calib_years = 1991:2020,
  scen_warming = 3,
  shift_per_deg = -4,
  cmp6_tas = NULL,
  gdd_base = 5,
  gdd_threshold1 = 150,
  gdd_threshold2 = 350,
  regions = NULL,
  delta_Tgl = 2.09,
  giss_data = NULL
)

```

## Arguments

pep	A data frame containing PEP725 phenology observations, including columns country, day, year, phase_id, and genus.
phase1	Integer. First phenophase (e.g. 65).
phase2	Integer. Second phenophase (e.g. 87).

genus_name	Character. Genus to filter (e.g. "Malus").
phenology_method	Either "simple_shift" or "gdd_cmip6".
ctrl_years	Integer vector. Years for CTRL period (default 2011:2021).
scen_years	Integer vector. Years for scenario period (default 2085:2095).
calib_years	Integer vector. Years for calibration period (default 1991:2020).
scen_warming	Numeric. Warming in degC (simple shift model).
shift_per_deg	Numeric. DOY shift per degC (simple shift).
cmip6_tas	Optional list of daily temperature vectors—only required if using "gdd_cmip6".
gdd_base	Numeric. Base temperature for GDD model.
gdd_threshold1	Numeric. GDD threshold for phase1.
gdd_threshold2	Numeric. GDD threshold for phase2.
regions	Character vector. Countries/regions to include. If NULL, all are used.
delta_Tg1	Numeric. Global temperature change (degC) for scenario.
giss_data	Data frame with GISS temperature data (columns: year, dT or Tg1). Required for robust_shift method.

**Value**

A data frame containing:

Country	Country name
doy_start_ctrl	Start DOY for CTRL period
doy_end_ctrl	End DOY for CTRL period
doy_start_scen	Start DOY for SCEN period
doy_end_scen	End DOY for SCEN period

**Author(s)**

Matthias Templ

**Examples**

```

pep <- pep_download()

# Use Alpine subset for faster computation
regions <- c("Austria", "Switzerland")
pep_alpine <- pep[country %in% regions]

# Simple shift method
d1 <- get_phenology_doys(
  pep = pep_alpine,
  phase1 = 65,
  phase2 = 87,
  genus_name = "Malus",

```

```
regions = regions,  
phenology_method = 'simple_shift'  
)  
d1
```

---

is.pep

*Test if Object is a PEP725 Data Object*

---

### **Description**

Test if Object is a PEP725 Data Object

### **Usage**

```
is.pep(x)
```

### **Arguments**

x                    An R object.

### **Value**

Logical; TRUE if x is of class pep.

### **Author(s)**

Matthias Templ

### **Examples**

```
pep <- pep_download()  
is.pep(pep)  
is.pep(data.frame(x = 1))
```

---

`kendall_tau`*Mann-Kendall Z-Statistic (deprecated alias for mann\_kendall\_z)*

---

### Description

This function was misnamed in `pep725 <= 1.0.2`: it returned the Mann-Kendall Z-statistic, not Kendall's  $\tau = S/(n(n-1)/2)$ . Please call `mann_kendall_z` instead for identical (and better: tie-corrected, continuity-corrected) behaviour.

### Usage

```
kendall_tau(x)
```

### Arguments

`x` Numeric vector (time series assumed to be equidistant). NA values are silently removed.

### Details

Note that this function is now a thin wrapper and will continue to return the Z-statistic until a future major release.

### Value

Numeric. Identical output shape to `mann_kendall_z()`.

### Author(s)

Matthias Templ

### See Also

[mann\\_kendall\\_z](#)

### Examples

```
suppressWarnings(kendall_tau(c(120, 118, 115, 112, 110, 108, 105)))
```

---

mann_kendall_z	<i>Mann-Kendall Z-Statistic</i>
----------------	---------------------------------

---

### Description

Computes the Mann-Kendall Z-statistic for a time series, a non-parametric test of monotonic trend. The Z-statistic is approximately standard normal under the no-trend null hypothesis.

### Usage

```
mann_kendall_z(x)
```

### Arguments

x                    Numeric vector (time series assumed to be equidistant). NA values are silently removed.

### Details

The implementation uses the standard tie correction for  $\text{Var}(S)$  (so that tied observations do not inflate the test) and the continuity correction  $Z = (S - \text{sign}(S))/\sqrt{\text{Var}(S)}$ , matching `Kendall::MannKendall()`.

### Value

Numeric. The Mann-Kendall Z-statistic. Positive values indicate an increasing trend, negative values decreasing. Compare against standard normal quantiles (e.g.,  $|Z| > 1.96$  for  $p < 0.05$ ). Returns NA if fewer than three non-missing values.

### Note on `kendall_tau()`

Earlier versions of `pep725` exported a function called `kendall_tau()` that actually returned this Mann-Kendall Z-statistic (not the true Kendall's  $\tau = S/(n(n-1)/2)$ ). `kendall_tau()` is now a deprecated alias of `mann_kendall_z()`; please update callers.

### Author(s)

Matthias Templ

### References

Mann, H.B. (1945). Nonparametric tests against trend. *Econometrica* 13, 245-259.  
Kendall, M.G. (1975). *Rank Correlation Methods*. 4th ed. London: Charles Griffin.

### See Also

[kendall\\_tau](#) (deprecated alias); [pheno\\_trend\\_turning](#) for a full sequential Mann-Kendall analysis.

## Examples

```
# Decreasing trend (earlier phenology)
mann_kendall_z(c(120, 118, 115, 112, 110, 108, 105))

# No clear trend
mann_kendall_z(c(120, 115, 122, 118, 121, 116, 119))
```

---

new\_pep

*Create a PEP725 Phenological Data Object*

---

## Description

Constructor function that creates a validated pep object from a `data.frame` or `data.table`. The object inherits from `data.table` and retains all its functionality.

## Usage

```
new_pep(x, validate = TRUE)
```

## Arguments

<code>x</code>	A <code>data.frame</code> or <code>data.table</code> containing PEP725 phenological data.
<code>validate</code>	Logical. If <code>TRUE</code> (default), validates that required columns are present and have correct types.

## Details

Required columns:

- `s_id` - Station ID
- `lon, lat` - Coordinates
- `genus, species` - Plant taxonomy
- `phase_id` - BBCH phenological phase code
- `year, day` - Observation timing

## Value

An object of class `c("pep", "data.table", "data.frame")`.

## Author(s)

Matthias Templ

**Examples**

```
# From imported data
dt <- pep_download()
pep_data <- new_pep(dt)
```

---

pep-class	<i>PEP725 Phenological Data Class</i>
-----------	---------------------------------------

---

**Description**

S3 class for PEP725 phenological data that inherits from `data.table`. Provides validation, informative print/summary methods, and convenient defaults.

---

pep725_demo	<i>Demonstrate the pep725 Package Functions</i>
-------------	---

---

**Description**

This function provides an interactive demonstration of the main features of the `pep725` package for PEP725 phenological data analysis. It showcases data exploration, filtering, summarization, and visualization capabilities.

**Usage**

```
pep725_demo(which = "all", pause = interactive())
```

**Arguments**

which	Character vector specifying which demos to run. Options are: <ul style="list-style-type: none"> <li>• "class" - Demonstrate the pep class (print, summary)</li> <li>• "filter" - Show data filtering and selection</li> <li>• "plot" - Display visualization capabilities</li> <li>• "analysis" - Show analysis functions (normals, anomalies)</li> <li>• "all" - Run all demonstrations (default)</li> </ul>
pause	Logical. If TRUE (default), pauses between demos waiting for user input. Set to FALSE for non-interactive use.

## Details

The demonstration downloads synthetic PEP725 data using `pep_download()` and walks through typical workflows for phenological data analysis:

1. **Class demo:** Shows the enhanced print and summary methods for the `pep` class, demonstrating how to quickly understand the data.
2. **Filter demo:** Demonstrates selecting specific species, phases, and time periods using `data.table` syntax.
3. **Plot demo:** Creates visualizations including station maps, time series, and DOY distributions.
4. **Analysis demo:** Shows phenological normals calculation, anomaly detection, and data quality assessment.

## Value

Invisibly returns a list containing example outputs from each demo.

## Author(s)

Matthias Templ

## See Also

[pep\\_download](#) for downloading synthetic data, [pheno\\_normals](#) for calculating normals, [pheno\\_anomaly](#) for anomaly detection, [pep\\_quality](#) for quality assessment

## Examples

```
if (interactive()) {  
  # Run all demos interactively  
  pep725_demo()  
  
  # Run specific demos  
  pep725_demo(which = "class")  
  pep725_demo(which = c("filter", "plot"))  
  
  # Run without pausing (for scripts)  
  pep725_demo(pause = FALSE)  
}
```

---

pep\_cache\_clear

*Clear Cached PEP Data*

---

## Description

Removes cached synthetic PEP data from the local cache directory.

**Usage**

```
pep_cache_clear(quiet = FALSE)
```

**Arguments**

quiet                    Logical. If TRUE, suppress messages. Default is FALSE.

**Value**

Invisible TRUE if cache was cleared, FALSE otherwise.

**Author(s)**

Matthias Templ

**See Also**

[pep\\_download](#)

**Examples**

```
# Clear the cache
pep_cache_clear()

# Next download will fetch fresh data
pep <- pep_download()
```

---

pep\_cache\_info                    *Get Cache Information*

---

**Description**

Returns information about the cached PEP data.

**Usage**

```
pep_cache_info()
```

**Value**

A list with cache information, or NULL if no cache exists.

**Author(s)**

Matthias Templ

## Examples

```
# Check cache status
pep_cache_info()
```

---

```
pep_check_connectivity
```

*Check Station-Year Connectivity*

---

## Description

Checks whether station-year combinations in phenological data form a connected set, which is required for valid combined time series estimation.

## Usage

```
pep_check_connectivity(pep, by = NULL)
```

## Arguments

<code>pep</code>	A pep object or data.table with year, day, and s_id columns.
<code>by</code>	Optional grouping variables to check connectivity within groups.

## Details

Connectivity means that there exist overlapping years between stations, allowing year effects to be separated from station effects. If data is disconnected, the combined series cannot be uniquely estimated across the disconnected parts.

## Value

A list with connectivity information:

**is\_connected** Logical. TRUE if all data forms one connected set.

**n\_sets** Number of disconnected sets found.

**set\_sizes** Number of observations in each set.

**largest\_set** Stations and years in the largest connected set.

## Author(s)

Matthias Templ

## See Also

[pheno\\_combine](#) which uses this internally

**Examples**

```

pep <- pep_download()
pep_alpine <- pep[country %in% c("Switzerland", "Austria")]
conn <- pep_check_connectivity(pep_alpine)
if (!conn$is_connected) {
  warning("Data has ", conn$n_sets, " disconnected sets")
}

```

---

pep\_check\_phases      *Check for Expected Phenological Phases*

---

**Description**

Validates that expected phenological phases (BBCH codes) are present in the data. Issues informative warnings for missing phases without stopping execution, making it useful for pipeline validation.

**Usage**

```

pep_check_phases(
  pep,
  expected = NULL,
  species = NULL,
  year_min = NULL,
  year_max = NULL,
  warn = TRUE,
  label = "Data"
)

```

**Arguments**

pep	A pep object or data.table containing phenological observations with a phase_id column.
expected	Integer vector of expected BBCH phase codes. If NULL (default), checks for common phases: 10 (leaf), 60 (heading/flowering), 65 (full flowering), and 100 (harvest).
species	Optional character string to filter by species/genus before checking. If specified, only checks phases for this species.
year_min	Optional integer. Only check phases for years $\geq$ this value. Useful for checking data availability within an analysis window.
year_max	Optional integer. Only check phases for years $\leq$ this value.
warn	Logical. If TRUE (default), issues warnings for missing phases. If FALSE, only returns the result silently.
label	Character. Optional label to include in warning messages, useful when checking multiple datasets. Default is "Data".

**Details**

This function is designed to be used at the start of analysis pipelines to validate that required phenological phases are available. It issues warnings but does not stop execution, allowing users to be informed of potential issues while still proceeding with available data.

**Value**

A list with class `phase_check` containing:

**expected** The expected phase codes

**present** Phase codes found in the data

**missing** Phase codes not found in the data

**extra** Phase codes in data but not in expected list

**complete** Logical. TRUE if all expected phases are present

**n\_obs** Named integer vector of observation counts per present phase

**coverage** Data.table with phase-level statistics

**Common BBCH Phases**

**10** Leaf development

**60** Beginning of flowering / Heading

**65** Full flowering / Anthesis

**69** End of flowering

**87** Hard dough (cereals)

**100** Harvest

**Author(s)**

Matthias Templ

**See Also**

[bbch\\_description](#) for phase code descriptions, [pep\\_completeness](#) for detailed completeness assessment, [pep\\_quality](#) for quality grading

**Examples**

```
pep <- pep_download()

# Use Swiss subset for faster checking
pep_ch <- pep[country == "Switzerland"]

# Basic check for common phases
check <- pep_check_phases(pep_ch)

# Check specific phases for grapevine (longest historical records)
check <- pep_check_phases(pep_ch,
```

```

        expected = c(60, 65, 81),
        species = "Vitis vinifera")

# Check within analysis window
check <- pep_check_phases(pep_ch,
                          expected = c(60, 65),
                          year_min = 1991,
                          year_max = 2020)

# Use in pipeline with custom label
vine_data <- pep_ch[species == "Vitis vinifera"]
if (nrow(vine_data) > 0) {
  pep_check_phases(vine_data, expected = c(60, 65),
                  label = "Grapevine analysis")
}

# Silent check (no warnings)
result <- pep_check_phases(pep_ch, warn = FALSE)
if (!result$complete) {
  # Handle missing phases programmatically
}

```

---

```
pep_check_phases_multi
```

*Check Multiple Phases Across Species*

---

## Description

A convenience function to check phase availability across multiple species at once.

## Usage

```

pep_check_phases_multi(
  pep,
  species_list,
  expected = c(60, 65, 100),
  year_min = NULL,
  year_max = NULL,
  warn = TRUE
)

```

## Arguments

pep	A pep object or data.table
species_list	Character vector of species/genus names to check
expected	Integer vector of expected BBCH phase codes

year_min	Optional minimum year filter
year_max	Optional maximum year filter
warn	Logical. Issue warnings for missing phases? Default TRUE.

**Value**

A data.table summarizing phase availability per species

**Author(s)**

Matthias Templ

**See Also**

[pep\\_check\\_phases](#) for single-species checking

**Examples**

```
# Check multiple species (use subset for speed)
pep <- pep_download()
pep_subset <- pep[country %in% c("Switzerland", "Austria")]
result <- pep_check_phases_multi(pep_subset,
                                species_list = c("Malus domestica",
                                                  "Vitis vinifera"),
                                expected = c(60, 65))

print(result)
```

---

pep\_completeness

*Assess Species and Phase Completeness of Phenological Data*

---

**Description**

Provides comprehensive assessment of data completeness across species, phases, years, and optionally countries. Useful for identifying data gaps and planning analyses.

**Usage**

```
pep_completeness(
  pep,
  by = c("genus", "species", "phase_id"),
  year_range = NULL,
  min_obs = 1,
  include_years = FALSE
)
```

**Arguments**

pep	A pep object or data.table containing phenological observations with columns year, day, genus, species, and phase_id.
by	Character vector specifying grouping dimensions. Options include "species", "genus", "phase_id", "year", "country", and "s_id". Default is c("genus", "species", "phase_id").
year_range	Optional integer vector of length 2 specifying the start and end years for assessment. If NULL (default), uses the full range of years in the data.
min_obs	Integer. Minimum number of observations required to include a group in the output. Default is 1.
include_years	Logical. If TRUE, includes year-level detail in the output (creates a row per group × year). Default is FALSE for summary mode.

**Details**

This function assesses data coverage across multiple dimensions, helping users understand where data is available and where gaps exist. It is particularly useful for:

- Identifying which species × phase combinations have sufficient data
- Finding temporal gaps in time series
- Comparing coverage across countries or regions
- Planning analyses that require minimum observation counts

**Value**

A data.table of class pep\_completeness with columns:

**by variables** Grouping variables as specified in by  
**n\_obs** Total number of observations  
**n\_stations** Number of unique stations  
**n\_years** Number of years with data  
**year\_min** First year of observations  
**year\_max** Last year of observations  
**year\_span** Total span in years  
**completeness\_pct** Percentage of years with data within span  
**median\_doy** Median day of year  
**iqr\_doy** Interquartile range of day of year

**Completeness Calculation**

Completeness is calculated as the percentage of years with at least one observation within the year span ( $\text{year\_max} - \text{year\_min} + 1$ ). A completeness of 100% means every year in the span has data.

**Author(s)**

Matthias Templ

**See Also**

[pep\\_check\\_phases](#) for validating expected phases exist, [pep\\_quality](#) for quality grading, [pep\\_coverage](#) for overall data coverage

**Examples**

```
pep <- pep_download()

# Use Swiss subset for faster computation
pep_ch <- pep[country == "Switzerland"]

# Basic completeness by species and phase
comp <- pep_completeness(pep_ch)
print(comp)

# Filter to well-observed combinations
comp_good <- comp[n_obs >= 100 & completeness_pct >= 80]

# Completeness by country (use Alpine countries)
pep_alpine <- pep[country %in% c("Switzerland", "Austria")]
comp_country <- pep_completeness(pep_alpine,
                                 by = c("country", "genus", "phase_id"))

# Year-level detail for a specific period
comp_yearly <- pep_completeness(pep_ch,
                                year_range = c(1991, 2020),
                                include_years = TRUE)

# Visualize completeness
plot(comp)
```

---

pep\_coverage

*Assess Data Coverage of PEP725 Phenological Data*

---

**Description**

Provides comprehensive coverage statistics for phenological datasets, including temporal, geographical, and species dimensions.

**Usage**

```
pep_coverage(
  x,
  kind = c("all", "temporal", "geographical", "species"),
  top = 10,
  plot = FALSE,
  by = NULL
)
```

**Arguments**

x	A pep object or data.frame with phenological data.
kind	Character. Type of coverage to assess: <ul style="list-style-type: none"> <li>• "all" (default): All coverage types combined</li> <li>• "temporal": Year range, gaps, observations per year</li> <li>• "geographical": Countries, stations, coordinate ranges</li> <li>• "species": Species and genus diversity</li> </ul>
top	Integer. Number of top entries to show in tables (default: 10).
plot	Logical. If TRUE, produces a visualization of the coverage. Default is FALSE.
by	Character. Optional grouping variable for more detailed analysis (e.g., "country", "species"). Only used for temporal coverage.

**Value**

A list of class pep\_coverage containing coverage statistics. The structure depends on kind:

**temporal** Year range, number of years, gaps, observations per year

**geographical** Countries, stations, coordinate bounds, altitude range

**species** Genera, species, subspecies counts and details

**all** All of the above combined

**Author(s)**

Matthias Templ

**Examples**

```
pep <- pep_download()
pep_ch <- pep[country == "Switzerland"]

# Full coverage report
pep_coverage(pep_ch)

# Temporal coverage only
pep_coverage(pep_ch, kind = "temporal")

# Species coverage, top 5
```

```
pep_coverage(pep_ch, kind = "species", top = 5)
```

---

 pep\_download

*Download Synthetic PEP725 Data*


---

## Description

Downloads a pre-generated synthetic version of PEP725 phenological data from an external repository. The data is cached locally after the first download to avoid repeated downloads.

## Usage

```
pep_download(url = NULL, cache = TRUE, force = FALSE, quiet = FALSE)
```

## Arguments

url	Character string specifying the URL to download from. Default uses the official pep725 package data repository.
cache	Logical. If TRUE (default), cache the downloaded data locally for future use.
force	Logical. If TRUE, force re-download even if cached data exists. Default is FALSE.
quiet	Logical. If TRUE, suppress download progress messages. Default is FALSE.

## Details

The synthetic data preserves the statistical properties and structure of real PEP725 data (stations, species, phases, temporal patterns) while ensuring data privacy. It is suitable for:

- Learning and testing package functions
- Reproducing examples from vignettes
- Developing analysis workflows before accessing real data

The data is cached in the platform-appropriate user cache directory determined by `R_user_dir("pep725", which = "cache")`. Typical locations:

- Windows: %LOCALAPPDATA%/R/cache/R/pep725/
- macOS: ~/Library/Caches/org.R-project.R/R/pep725/
- Linux: ~/.cache/R/pep725/

## Value

A pep object containing synthetic phenological data with the same structure as original PEP725 data.

## Data Access

If you have access to real PEP725 data, you can import it using [pep\\_import](#) instead.

**Author(s)**

Matthias Templ

**See Also**[pep\\_simulate](#) for creating your own synthetic data, [pep\\_import](#) for importing real PEP725 data**Examples**

```
# Download synthetic data
pep <- pep_download()

# Explore the data
print(pep)
summary(pep)

# Force re-download
pep <- pep_download(force = TRUE)
```

---

pep\_flag\_outliers      *Flag Phenological Outliers*

---

**Description**

Identifies outlier observations in phenological data using the 30-day rule or statistical methods. Based on the approach described by Schaber & Badeck (2002) for quality control of phenological observations.

**Usage**

```
pep_flag_outliers(
  pep,
  by = c("s_id", "genus", "species", "phase_id"),
  method = c("30day", "mad", "iqr", "zscore", "gam_residual", "mahalanobis"),
  threshold = NULL,
  center = c("median", "mean"),
  min_obs = 5,
  formula = day ~ s(year) + s(alt) + s(lat) + s(s_id, bs = "re"),
  min_n_per_group = 50,
  flag_only = TRUE
)
```

**Arguments**

pep	A pep object or data.table with phenological observations. Must contain year, day, and grouping columns.
by	Character vector of columns defining groups for outlier detection. Default c("s_id", "genus", "species", "phase_id") detects outliers within each station-species-phase combination.
method	<p>Character. Outlier detection method:</p> <p><b>"30day"</b> (Default) Flag observations &gt;30 days from group mean/median. The classic Schaber-Badeck approach for phenological data.</p> <p><b>"mad"</b> Use Median Absolute Deviation. Flag if <math> deviation  &gt; k * MAD</math>, where <math>k</math> is set by threshold.</p> <p><b>"iqr"</b> Use Interquartile Range. Flag if outside <math>(Q1 - kIQR, Q3 + kIQR)</math>.</p> <p><b>"zscore"</b> Flag if <math> z\text{-score}  &gt; \text{threshold}</math> (default 3).</p> <p><b>"gam_residual"</b> Fit a GAM (via <code>mgcv::gam()</code>) per group using the supplied formula — by default accounting for year trend, elevation, latitude, and a station random intercept — and flag observations whose robust-z-scored residual exceeds threshold. Detects covariate-inconsistent anomalies that the within-group "30day" rule misses. Falls back to "30day" on groups with fewer than <code>min_n_per_group</code> observations or when the model fails to converge.</p> <p><b>"mahalanobis"</b> Treat each station-year as a vector of DOYs across phases and flag station-years whose <b>robust</b> Mahalanobis distance exceeds the threshold. The centre and covariance are estimated by the Minimum Covariance Determinant (MCD) estimator of Rousseeuw (1984), via <code>robustbase::covMcd()</code> — using the classical, non-robust covariance here would let the outliers contaminate the estimate and mask themselves. Under MCD the squared distance is still approximately <math>\chi_p^2</math>, so the default threshold is <math>\sqrt{\chi_{0.975,p}^2}</math>, where <math>p</math> is the number of phases present in the group. Detects station-years whose pattern across phases is jointly inconsistent (e.g. BBCH 60 and 65 impossibly close) even when each marginal DOY looks fine. All rows within a flagged station-year are marked; the <code>deviation</code> column stores the robust Mahalanobis distance. Groups with too few complete station-years (fewer than <math>\max(p + 5, 15)</math>) or a singular MCD fit fall back to the 30-day rule.</p>
threshold	<p>Numeric. Threshold for outlier detection.</p> <p><b>For "30day":</b> Days deviation to flag as outlier. Default 30.</p> <p><b>For "mad":</b> Number of MADs. Default 3 (~2.5 SD equivalent).</p> <p><b>For "iqr":</b> IQR multiplier. Default 1.5 (standard Tukey rule).</p> <p><b>For "zscore":</b> Z-score threshold. Default 3.</p> <p><b>For "gam_residual":</b> Robust-z threshold on residuals. Default 3.5.</p>
center	Character. Central tendency measure: "median" (default, more robust) or "mean".
min_obs	Integer. Minimum observations per group required for outlier detection. Groups with fewer observations are skipped. Default 5.

formula	Optional formula for method = "gam_residual". Default is $\text{day} \sim \text{s}(\text{year}) + \text{s}(\text{alt}) + \text{s}(\text{lat}) + \text{s}(\text{s\_id}, \text{bs} = \text{"re"})$ ; any smooth term referencing a column missing from pep is silently dropped.
min_n_per_group	Integer. Minimum group size required before fitting the GAM in method = "gam_residual". Groups below this threshold fall back to the "30day" rule. Default 50.
flag_only	Logical. If TRUE (default), adds outlier flag columns to data. If FALSE, removes flagged outliers from data.

### Details

The 30-day rule (Schaber & Badeck 2002) is widely used for phenological data quality control. It flags observations that deviate more than 30 days from the expected value for that station-species-phase combination. This threshold is based on the observation that legitimate phenological variation rarely exceeds one month.

### Value

If flag\_only = TRUE: The input data with additional columns:

**is\_outlier** Logical. TRUE for flagged outliers.

**deviation** Numeric. Days deviation from expected value.

**expected\_doy** Numeric. Expected DOY for the group.

If flag\_only = FALSE: Data with outliers removed.

### When to Use Each Method

**30day** Standard for phenological QC. Use when the 30-day biological threshold is meaningful.

**mad** Robust to existing outliers. Good for initial screening of potentially contaminated data.

**iqr** Standard statistical approach. Useful when comparing with other quality metrics.

**zscore** Parametric approach. Use when data is approximately normal.

### Author(s)

Matthias Templ

### References

Schaber J, Badeck F-W (2002). Evaluation of methods for the combination of phenological time series and outlier detection. *Tree Physiology* 22:973-982.

Rousseeuw P J (1984). Least median of squares regression. *Journal of the American Statistical Association* 79(388):871-880. (Minimum Covariance Determinant estimator used by method = "mahalanobis".)

**See Also**

[pep\\_quality](#) for comprehensive quality assessment, [pheno\\_combine](#) which uses residuals for outlier detection

**Examples**

```
data(pep_seed)

# Flag outliers using 30-day rule
pep_flagged <- pep_flag_outliers(pep_seed)
table(pep_flagged$is_outlier)

# View flagged observations
pep_flagged[is_outlier == TRUE]

# Remove outliers instead of flagging
pep_clean <- pep_flag_outliers(pep_seed, flag_only = FALSE)

# Use MAD method with stricter threshold
pep_flagged <- pep_flag_outliers(pep_seed, method = "mad", threshold = 2.5)

# Group by country instead of station
pep_flagged <- pep_flag_outliers(pep_seed,
                                by = c("country", "genus", "phase_id"))
```

---

 pep\_import

---

*Import and preprocess PEP725 phenological data*


---

**Description**

This function imports all CSV files from the specified folder, reads them efficiently using **data.table**, and combines them into a single data table. It performs several preprocessing steps, including:

- Converting selected columns to factors (provider\_id, s\_id, gss\_id, genus, species, subspecies).
- Replacing missing altitude values coded as -9999 with NA.
- Creating a combined altitude variable alt2 using alt and alt\_dem.
- Converting the date column to Date class.
- Recoding cultivation season (cult\_season) and quality control flags (qc\_ori\_flag) into labeled factors.
- Removing unused or problematic columns (affected\_flag, qc\_flag).

**Usage**

```
pep_import(path = "data/Data_PEP725_all", flags = FALSE, add_country = TRUE)
```

### Arguments

path	path to the folder containing PEP725 CSV files (default is "data/Data_PEP725_all/").
flags	Logical indicating whether the pep data contains quality control flags (default is FALSE).
add_country	Logical indicating whether to add country information based on station coordinates (default is TRUE).

### Details

This function is intended for importing raw PEP725 station data into a standardized format suitable for further phenological analysis, such as sensitivity and trend estimation.

### Value

A `pep` object (extends `data.table`) containing the combined and preprocessed PEP725 data.

### Author(s)

Matthias Templ (FHNW)

### See Also

[fread](#), [rbindlist](#)

### Examples

```
# Small example files shipped with the package
path <- system.file("extdata", package = "pep725")
pep_data <- pep_import(path = path, add_country = FALSE)
print(pep_data)
```

---

pep\_outliers\_leaflet *Interactive Leaflet Map of Outlier-Flagged Stations*

---

### Description

Renders a Leaflet map of the stations in a `pep_outliers` object, with each station plotted as a circle marker whose size encodes the number of flagged observations and whose colour encodes the maximum absolute residual (or Mahalanobis distance) for the station. Popups list the worst-offending station-years and their residual values — useful for interactive inspection during QC.

**Usage**

```
pep_outliers_leaflet(
  x,
  outlier_only = TRUE,
  top_n = 10,
  radius_range = c(4, 14),
  palette = "plasma",
  ...
)
```

**Arguments**

<code>x</code>	A <code>pep_outliers</code> object from <code>pep_flag_outliers</code> . Must contain <code>lon</code> , <code>lat</code> , <code>s_id</code> , <code>year</code> , <code>phase_id</code> , <code>day</code> , <code>is_outlier</code> , and <code>deviation</code> columns.
<code>outlier_only</code>	Logical. If <code>TRUE</code> (default), only stations with at least one flagged observation are shown. If <code>FALSE</code> , every station appears (non-flagged stations are drawn small and grey for context).
<code>top_n</code>	Integer. Maximum number of station-year records included in a station's popup. Default 10.
<code>radius_range</code>	Numeric length-2. Minimum and maximum circle marker radius (in pixels) mapped to $\log_{1p}(n\_outliers)$ . Default <code>c(4, 14)</code> .
<code>palette</code>	Character. Name of a <code>RColorBrewer</code> or <code>viridis</code> palette for the colour scale. Default <code>"plasma"</code> .
<code>...</code>	Additional arguments reserved for future expansion.

**Details**

Unlike `pheno_leaflet` (which is a selection gadget built on Shiny + miniUI), this is a pure visualisation: it returns a `leaflet` `htmlwidget` so you can embed it in an R Markdown report, include it in a vignette, or call `print()` on it for interactive exploration.

**Colour** encodes the station's maximum absolute residual (for univariate methods:  $\max(|deviation|)$  in days; for `method = "mahalanobis"`: the maximum robust Mahalanobis distance). **Size** encodes  $\log_{1p}(n\_outliers)$  — a station flagged once has a small marker, a station flagged 30 times has a much larger one.

For `method = "gam_residual"` and `"mahalanobis"` this is the natural companion to the Schaber-Badeck-style text tables and the static diagnostic figure, because spatial structure in outliers (a local observer network, a regional climate anomaly) is often only visible on a map.

**Value**

A `leaflet` `htmlwidget`. Call `print()` to display in the RStudio Viewer or embed it in an Rmd file.

**Author(s)**

Matthias Templ

**See Also**

[pep\\_flag\\_outliers](#) — detection, [pep\\_plot\\_outliers](#) — static diagnostic plots, [pheno\\_leaflet](#) — station-selection Shiny gadget.

**Examples**

```
## Not run:
pep <- pep_download()
pep_ch <- pep[country == "Switzerland" & species == "Malus domestica"]
flagged <- pep_flag_outliers(
  pep_ch,
  by = c("genus", "species", "phase_id"),
  method = "gam_residual"
)
pep_outliers_leaflet(flagged)

## End(Not run)
```

---

 pep\_plot\_outliers

*Visualize Phenological Outliers for Inspection*


---

**Description**

Creates diagnostic plots to help distinguish between data errors and biologically meaningful extreme events (e.g., second flowering). Multiple plot types reveal different aspects of outlier patterns.

**Usage**

```
pep_plot_outliers(
  x,
  type = c("overview", "seasonal", "map", "detail", "station", "doy_context",
    "diagnostic", "profile"),
  phase_id = NULL,
  outlier_only = NULL,
  late_threshold = 250,
  n_top = 20,
  ...
)
```

**Arguments**

x	A pep_outliers object from <a href="#">pep_flag_outliers</a> .
type	Character. Type of plot to produce: <b>"overview"</b> (Default) Multi-panel overview with distribution, seasonal pattern, and summary statistics.

	<b>"seasonal"</b> Distribution of outliers by month/season, useful for detecting second flowering (late-season outliers for spring phases).
	<b>"map"</b> Geographic distribution of outliers (requires lon/lat).
	<b>"detail"</b> Detailed view of individual outlier events with context.
	<b>"station"</b> Station-level outlier patterns over time.
	<b>"doy_context"</b> Shows outliers in context of full DOY distribution per phase, highlighting potential second events.
	<b>"diagnostic"</b> Paper- and vignette-ready 4-panel figure of model fit quality: (A) residuals vs fitted; (B) Q-Q plot of residuals; (C) residual vs altitude (if available) or year; (D) spatial map of maximum residual per station. Designed to accompany <code>method = "gam_residual"</code> but usable for every method (uses the deviation column as the residual). For <code>method = "mahalanobis"</code> the panels automatically switch to MD-specific diagnostics (sorted MD with chi-square threshold; Q-Q against $\chi_p^2$ ; mean / max MD over time; per-station worst-case MD map).
	<b>"profile"</b> Parallel-coordinates plot of the phase profile (one line per station-year across phases), with flagged station-years highlighted and the robust central profile overlaid. Designed for <code>method = "mahalanobis"</code> where the outlieriness lives in the <i>shape</i> of a station-year across phases, not in any single DOY. The primary paper figure for the multivariate method.
<code>phase_id</code>	Optional integer vector to filter specific phases for plotting.
<code>outlier_only</code>	Logical. If TRUE (default for some types), show only outlier observations. If FALSE, show all data with outliers highlighted.
<code>late_threshold</code>	Integer. DOY threshold for classifying "late" outliers as potential second events (default 250, ~September 7).
<code>n_top</code>	Integer. For detail view, number of most extreme outliers to show. Default 20.
<code>...</code>	Additional arguments passed to plotting functions.

## Details

This function is designed to support quality control workflows where the goal is to distinguish:

1. **Data errors:** Typos, wrong dates, mis-coded phases
2. **Observer errors:** Misidentified species or phases
3. **Biologically meaningful extremes:** Second flowering, delayed development, climate-driven anomalies

For detecting potential **second flowering events**:

- Use `type = "seasonal"` to see if late-season outliers cluster in autumn months
- Use `type = "doy_context"` to visualize outliers relative to the main flowering distribution
- Focus on flowering phases (60, 65) with late DOY values (>250)

## Value

A ggplot object (or list of ggplots for "overview").

**Author(s)**

Matthias Templ

**See Also**[pep\\_flag\\_outliers](#) for detecting outliers, [pep\\_quality](#) for comprehensive quality assessment**Examples**

```
pep <- pep_download()

# Use Swiss subset for faster outlier detection
pep_ch <- pep[country == "Switzerland"]
pep_flagged <- pep_flag_outliers(pep_ch)

# Overview of all outliers
pep_plot_outliers(pep_flagged, type = "overview")

# Seasonal distribution - look for autumn outliers in flowering phases
pep_plot_outliers(pep_flagged, type = "seasonal", phase_id = c(60, 65))

# Geographic pattern
pep_plot_outliers(pep_flagged, type = "map")

# Detailed view of most extreme outliers
pep_plot_outliers(pep_flagged, type = "detail", n_top = 30)

# DOY context for flowering phase - spot potential second flowering
pep_plot_outliers(pep_flagged, type = "doy_context", phase_id = 65)
```

---

pep\_quality

*Assess Data Quality of Phenological Observations*

---

**Description**

Provides a comprehensive assessment of data completeness, temporal coverage, outlier prevalence, and assigns quality grades to phenological time series.

**Usage**

```
pep_quality(
  pep,
  by = c("s_id", "genus", "phase_id"),
  year_range = NULL,
  species = NULL,
  phase_id = NULL,
  outlier_method = c("tukey", "zscore"),
```

```

    na.rm = TRUE
  )

```

### Arguments

<code>pep</code>	A <code>pep</code> object or <code>data.table</code> containing phenological observations with columns <code>year</code> , <code>day</code> , and grouping variables.
<code>by</code>	Character vector of column names to group by for quality assessment. Default is <code>c("s_id", "genus", "phase_id")</code> for station-level assessment.
<code>year_range</code>	Optional integer vector of length 2 specifying the start and end years for assessment. If <code>NULL</code> (default), uses the full range of years in the data.
<code>species</code>	Optional character string to filter by species/genus.
<code>phase_id</code>	Optional integer vector to filter by BBCH phase codes.
<code>outlier_method</code>	Character string specifying outlier detection method. Either <code>"tukey"</code> (default, uses $1.5 \times \text{IQR}$ rule) or <code>"zscore"</code> (flags values with $ z  > 3$ ).
<code>na.rm</code>	Logical. Should missing values be removed? Default <code>TRUE</code> .

### Details

This function evaluates phenological time series quality based on multiple criteria important for trend analysis and climatological calculations.

### Value

A `data.table` with the following columns:

<b>by variables</b>	Grouping variables as specified in <code>by</code>
<b>n_obs</b>	Total number of observations
<b>year_min</b>	First year of observations
<b>year_max</b>	Last year of observations
<b>year_span</b>	Total span in years ( <code>year_max - year_min + 1</code> )
<b>n_years</b>	Number of distinct years with data
<b>completeness_pct</b>	Percentage of years with data within the span
<b>max_gap_years</b>	Length of the longest gap (consecutive missing years)
<b>n_gaps</b>	Number of gaps in the time series
<b>n_outliers</b>	Number of statistical outliers detected
<b>outlier_pct</b>	Percentage of observations that are outliers
<b>doy_mean</b>	Mean day-of-year
<b>doy_sd</b>	Standard deviation of day-of-year
<b>doy_range</b>	Range of day-of-year values
<b>quality_grade</b>	Overall quality grade (A, B, C, or D)

### Quality Grading Criteria

**Grade A** Completeness  $\geq 80\%$ , no gaps  $> 2$  years, outliers  $< 2\%$

**Grade B** Completeness  $\geq 60\%$ , no gaps  $> 5$  years, outliers  $< 5\%$

**Grade C** Completeness  $\geq 40\%$ , gaps  $\leq 10$  years, outliers  $< 10\%$

**Grade D** Below grade C thresholds

### Outlier Detection

The Tukey method (default) flags observations outside the interquartile fences:  $[Q1 - 1.5 \cdot IQR, Q3 + 1.5 \cdot IQR]$ . This is robust to non-normal distributions common in phenological data.

### Author(s)

Matthias Templ

### References

Schaber, J. and Badeck, F.-W. (2002). Evaluation of methods for the combination of phenological time series and outlier detection. *Tree Physiology*, 22, 973–982. doi:10.1093/treephys/22.14.973

### See Also

[pheno\\_normals](#) for calculating climatological normals, [pheno\\_anomaly](#) for anomaly detection

### Examples

```
pep <- pep_download()

# Subset to one country for speed
pep_ch <- pep[country == "Switzerland"]

# Assess quality for stations in Switzerland
quality <- pep_quality(pep_ch)
print(quality)

# Filter to high-quality stations only
high_quality <- quality[quality_grade %in% c("A", "B")]
high_quality

# Assess specific species and phase
apple_quality <- pep_quality(pep_ch,
                             species = "Malus",
                             phase_id = 60)
apple_quality

# Country-level assessment (use subset of countries)
pep_subset <- pep[country %in% c("Switzerland", "Austria")]
country_quality <- pep_quality(pep_subset,
                               by = c("country", "genus", "phase_id"))
country_quality
```

```
# Assess quality within a specific time window
modern_quality <- pep_quality(pep_ch,
                             year_range = c(1991, 2020))
modern_quality
```

---

pep\_second\_events      *Detect Second Flowering and Other Repeated Phenological Events*

---

### Description

Identifies potential second flowering events and other irregular repeated phenological observations. These may represent biologically meaningful responses to climate disruption rather than data errors.

### Usage

```
pep_second_events(
  pep,
  phases = c(60, 65),
  method = c("late_season", "multiple_per_year", "both"),
  late_threshold = 250,
  gap_threshold = 60,
  reference_period = 1991:2020,
  z_threshold = 3,
  min_obs = 10
)
```

### Arguments

pep	A pep object or data.table with phenological observations.
phases	Integer vector of phase IDs to analyze. Default is c(60, 65) for flowering phases. Use NULL to analyze all phases.
method	Character. Detection method: <b>"late_season"</b> (Default) Flag observations occurring after the expected phenological window (DOY > late_threshold). <b>"multiple_per_year"</b> Detect multiple observations of the same phase at the same station within a single year. <b>"both"</b> Apply both methods.
late_threshold	Integer. DOY threshold for "late season" events. Default is 250 (~September 7). Events after this date for spring phases are flagged as potential second events.
gap_threshold	Integer. For "multiple_per_year" method, minimum days between observations to consider them separate events. Default is 60.

reference_period	Integer vector of years to calculate expected phenological windows. Default is 1991:2020.
z_threshold	Numeric. Z-score threshold for identifying late events relative to the reference period. Default is 3.
min_obs	Integer. Minimum observations in reference period to calculate expected windows. Default is 10.

### Details

Second flowering and other repeated phenological events are increasingly reported as climate patterns become more irregular. These events may indicate:

- Disrupted coordination between environmental cues and development
- Response to unusual autumn warmth after summer dormancy
- Failure of normal seasonal inhibition mechanisms

The function uses two complementary approaches:

1. **Late season detection:** Identifies observations occurring well after the normal phenological window (e.g., flowering in autumn)
2. **Multiple events:** Finds stations reporting the same phase multiple times in one year with sufficient gap between observations

### Value

A list of class `second_events` containing:

**events** Data.table of detected second/repeated events with columns: `s_id`, `species`, `phase_id`, `year`, `day`, `event_type`, expected timing info

**summary** Summary statistics by species, phase, and event type

**by\_year** Annual counts of detected events

**by\_month** Monthly distribution of events

**total\_by\_year** Total observations per year (for relative scaling in plots)

**method** Detection method used

**params** Parameters used for detection

### Interpreting Results

Not all detected events are necessarily "true" second flowering. Consider:

- **High confidence:** Same station reports phase twice with 60+ day gap; late observation is clearly outside normal window
- **Medium confidence:** Single late observation at a station; could be second event or data entry error
- **Requires verification:** Check original data sources, geographic context, species biology

**Author(s)**

Matthias Templ

**References**

Related to emerging research on phenological irregularity and climate disruption of seasonal timing.

**See Also**

[pep\\_flag\\_outliers](#) for general outlier detection, [pep\\_plot\\_outliers](#) for outlier visualization

**Examples**

```
pep <- pep_download()

# Use grapevine flowering in Alpine region (long records, second events possible)
pep_subset <- pep[species == "Vitis vinifera" & phase_id %in% c(60, 65) &
  country %in% c("Germany-South", "Switzerland", "Austria")]

if (nrow(pep_subset) > 0) {
  # Method 1: Detect late-season flowering events (default)
  second <- pep_second_events(pep_subset, phases = c(60, 65))
  print(second)
  summary(second)

  # Plot from 1980 onwards for clearer trends
  plot(second, from_year = 1980)
  plot(second, type = "overview", scale = "relative", from_year = 1980)

  # Method 2: Detect repeated observations at same station
  repeated <- pep_second_events(pep_subset, method = "multiple_per_year")
  print(repeated)
  plot(repeated, type = "timeline", from_year = 1980)

  # Method 3: Combine both detection methods
  all_events <- pep_second_events(pep_subset, method = "both")
  summary(all_events)
  plot(all_events)
  plot(all_events, scale = "relative")
}
```

---

 pep\_seed

---

*Minimal Seed Dataset for Synthetic Data Generation*


---

**Description**

A small subset of phenological data suitable for use with [pep\\_simulate](#) to generate synthetic phenological datasets. This dataset contains real structure but limited scope, making it suitable for package examples and testing.

## Format

A data.table with 1,319 rows and 10 variables:

**s\_id** Station identifier (factor)  
**lon** Longitude in decimal degrees  
**lat** Latitude in decimal degrees  
**alt** Altitude in meters  
**genus** Plant genus (factor)  
**species** Full species name (factor)  
**phase\_id** BBCH phenological phase code (integer)  
**year** Observation year  
**day** Day of year (DOY) of the phenological event  
**country** Country name

## Details

This dataset includes observations for:

- **Species:** Triticum aestivum (wheat), Vitis vinifera (grapevine), Malus domestica (apple)
- **Phases:** BBCH 10 (emergence), 60 (flowering), 65 (full flowering)
- **Countries:** Austria, Germany-South
- **Years:** 1990-2015
- **Stations:** 30 stations

For a full synthetic dataset, use [pep\\_download](#) to download pre-generated synthetic data from the package repository.

## Source

Derived from PEP725 Pan European Phenology Database <https://pep725.eu/>

## See Also

[pep\\_simulate](#) for generating synthetic data, [pep\\_download](#) for downloading full synthetic dataset

## Examples

```
# Load the seed dataset
data(pep_seed)

# Examine structure
str(pep_seed)

# Use with pep_simulate to create synthetic data
pep_synthetic <- pep_simulate(pep_seed)
```

pep\_simulate

*Simulate Synthetic PEP Data***Description**

Generates synthetic phenological observations by fitting a GAM smooth ( $\text{day} \sim \text{s}(\text{year})$ ) per station-species-phase group and replacing the observed day values with predictions plus Gaussian noise. The station-year-species-phase structure of the input data is kept intact.

**Usage**

```
pep_simulate(pep, min_obs = 20, seed = 42, progress = TRUE)
```

**Arguments**

pep	A pep object or data.table with at least the columns species, phase_id, s_id, year, and day.
min_obs	Integer. Minimum number of observations required per station-species-phase group to attempt GAM fitting. Groups with fewer observations keep their original day values unchanged. Default is 20.
seed	Integer. Random seed for reproducibility. Default is 42.
progress	Logical. Show a progress bar? Default TRUE.

**Details**

For each qualifying group (station x species x phase with at least min\_obs observations), a GAM is fitted and synthetic day-of-year values are generated as  $\text{round}(\text{predicted} + \text{rnorm}(n, \text{sd} = \text{residual\_sd}))$ .

Groups where the GAM fit fails receive a fallback: the species-phase median day plus Gaussian jitter ( $\text{sd} = 5$  days).

Groups below the min\_obs threshold are left unchanged. A message reports how many rows were not simulated.

**Value**

A pep object with the same rows and columns as the input. The day column is overwritten with synthetic values for groups that meet the min\_obs threshold; other rows retain their original values.

**Author(s)**

Matthias Templ

**Examples**

```
pep <- pep_download()
# Single species + country for fast example
vine_ch <- pep[species == "Vitis vinifera" & country == "Switzerland"]
pep_synth <- pep_simulate(vine_ch)
```

pheno\_anomaly

*Calculate Phenological Anomalies***Description**

Computes deviations from long-term phenological baselines to identify extreme years and detect climate signals. Returns both raw anomalies (in days) and standardized anomalies for cross-species comparison.

**Usage**

```
pheno_anomaly(
  pep,
  baseline_period = 1961:1990,
  target_years = NULL,
  by = c("country", "genus", "phase_id"),
  species = NULL,
  phase_id = NULL,
  robust = TRUE,
  min_years = 15,
  extreme_threshold = 2,
  normals = NULL,
  na.rm = TRUE
)
```

**Arguments**

pep	A pep object or data.table containing phenological observations with columns year, day, and grouping variables.
baseline_period	Integer vector specifying the years to use for baseline calculation. Default is 1961:1990 (pre-acceleration warming reference period).
target_years	Integer vector specifying years to calculate anomalies for. If NULL (default), calculates for all years in the data.
by	Character vector of column names to group by. Must match columns in the data. Default is c("country", "genus", "phase_id").
species	Optional character string to filter by species/genus. If NULL (default), all species are included.
phase_id	Optional integer vector to filter by BBCH phase codes. If NULL (default), all phases are included.
robust	Logical. If TRUE (default), uses median and MAD for baseline and standardization. If FALSE, uses mean and SD.
min_years	Minimum number of years required in the baseline period to calculate valid anomalies. Default is 15.

<code>extreme_threshold</code>	Numeric. Z-score threshold for flagging extreme events. Default is 2 (approximately 95th percentile for normal distribution).
<code>normals</code>	Optional pre-computed <code>pheno_normals</code> object. If provided, baseline statistics are taken from this object instead of being computed.
<code>na.rm</code>	Logical. Should missing values be removed? Default TRUE.

## Details

Phenological anomalies quantify how much a given year deviates from long-term expectations. This is crucial for:

- Detecting climate change signals in phenology
- Identifying extreme phenological years
- Comparing anomalies across different species/regions

The function supports two approaches:

- **Robust** (default): Uses median for central tendency and MAD (Median Absolute Deviation) for spread. More resistant to outliers.
- **Classical**: Uses mean and standard deviation. More sensitive to extreme values but provides interpretable z-scores under normality.

## Value

A `data.table` with the following columns:

**by variables** Grouping variables as specified in `by`  
**year** Year of observation  
**observed\_doy** Observed mean day-of-year for the group/year  
**baseline\_doy** Long-term baseline (median if robust, mean otherwise)  
**baseline\_spread** Baseline spread (MAD if robust, SD otherwise)  
**anomaly\_days** Deviation in days (negative = early, positive = late)  
**z\_score** Standardized anomaly (anomaly / spread)  
**percentile** Percentile rank relative to baseline distribution  
**is\_extreme** Logical flag for extreme events ( $|z\_score| > \text{threshold}$ )  
**direction** Character: "early", "late", or "normal"

## Interpretation

- **anomaly\_days**: Direct interpretation - "X days earlier/later than normal"
- **z\_score**: Standardized - values  $> 2$  or  $< -2$  are typically considered extreme
- **percentile**: Normal approximation from z-score - e.g., 5th percentile means "earlier than 95\

## Author(s)

Matthias Templ

## References

- Menzel, A. et al. (2006). European phenological response to climate change matches the warming pattern. *Global Change Biology*, 12(10), 1969–1976. doi:10.1111/j.13652486.2006.01193.x
- Templ, M., Templ, B., and Filzmoser, P. (2018). Modelling and prediction of phenological data of the beginning of flowering: Austrian indicators of climate change. *International Journal of Biometeorology*, 62, 1319–1334. doi:10.1007/s0048401815342

## See Also

[pheno\\_normals](#) for calculating baseline statistics, [pep\\_download](#) for obtaining the main dataset

## Examples

```
pep <- pep_download()

# Grapevine in Switzerland (longest historical records back to 1775)
vine_ch <- pep[country == "Switzerland" & species == "Vitis vinifera"]

if (nrow(vine_ch) > 0) {
  # Calculate anomalies relative to 1961-1990 baseline
  anomalies <- pheno_anomaly(vine_ch,
                             baseline_period = 1961:1990,
                             phase_id = 65)

  print(anomalies)

  # Find extreme early years
  extreme_early <- anomalies[is_extreme == TRUE & direction == "early"]
  extreme_early

  # Anomalies for recent years only
  recent <- pheno_anomaly(vine_ch,
                          baseline_period = 1961:1990,
                          target_years = 2000:2020,
                          phase_id = 65)

  recent
}
```

## Description

Estimates a combined (regional/national) phenological time series from multi-station observations by separating year effects from station effects. This is essential for creating representative time series from networks with varying station coverage over time.

**Usage**

```
pheno_combine(
  pep,
  by = NULL,
  method = c("robust", "mixed", "ols"),
  min_years = 5,
  min_stations = 3,
  check_connectivity = TRUE
)
```

**Arguments**

pep	A pep object or data.table containing phenological observations with columns year, day, and s_id.
by	Character vector of grouping variables (e.g., c("genus", "species", "phase_id")). A separate combined series is created for each group. Default is NULL (treat all data as one group).
method	Character. Estimation method: <b>"robust"</b> (Default) Least Absolute Deviations (LAD/L1) regression. Minimizes sum of absolute residuals, robust to outliers. <b>"mixed"</b> Mixed-effects model with year as fixed effect and station as random effect. Uses REML estimation. <b>"ols"</b> Ordinary Least Squares. Sensitive to outliers but provides standard errors and confidence intervals.
min_years	Integer. Minimum years of overlap required. Default 5.
min_stations	Integer. Minimum stations required. Default 3.
check_connectivity	Logical. If TRUE (default), checks whether station-year combinations form a connected set. Warns if data is disconnected (which prevents valid combined series estimation).

**Details**

The model fitted is:

$$DOY_{ij} = \mu_i + \sigma_j + \epsilon_{ij}$$

where  $\mu_i$  is the year effect (combined series),  $\sigma_j$  is the station effect, and  $\epsilon_{ij}$  is the residual.

Station effects represent systematic differences between stations (e.g., due to altitude, microclimate, or observer differences). The combined series removes these effects to reveal the underlying temporal trend.

**Value**

An object of class pheno\_combined containing:

**combined** Data.table with the combined time series: year, year\_effect (the combined DOY), se (standard error, if available)

**station\_effects** Data.table of station effects: s\_id, effect, se  
**residuals** Data.table with original data plus fitted values and residuals for outlier detection  
**method** Estimation method used  
**connectivity** Connectivity check results (if performed)  
**by** Grouping variables used  
**group\_results** For grouped analysis, results per group

### Connectivity

A prerequisite for valid estimation is that station-year combinations form a "connected" set - meaning there must be overlapping years between stations to separate year and station effects. The function checks this automatically and warns if data is disconnected.

### Outlier Detection

With method = "robust", large residuals (e.g., > 30 days) indicate potential outliers or data errors. The residuals table can be used to identify and investigate these.

### Author(s)

Matthias Templ

### References

Schaber J, Badeck F-W (2002). Evaluation of methods for the combination of phenological time series and outlier detection. *Tree Physiology* 22:973-982.

### See Also

[pheno\\_normals](#) for climatological baselines, [pheno\\_trend\\_turning](#) for trend turning point detection, [pep\\_check\\_connectivity](#) for connectivity assessment

### Examples

```
pep <- pep_download()

# Grapevine flowering in Switzerland (longest historical records)
vine_ch <- pep[species == "Vitis vinifera" &
              phase_id == 65 & country == "Switzerland"]

if (nrow(vine_ch) > 0) {
  # Create combined series using OLS
  combined <- pheno_combine(vine_ch, method = "ols")
  print(combined)

  # Identify potential outliers (residuals > 30 days)
  outliers <- combined$residuals[abs(residual) > 30]
  outliers
}
```

---

pheno\_gradient      *Analyze Phenological Gradients with Elevation or Latitude*

---

### Description

Quantifies how phenological timing shifts with altitude or latitude, calculating the phenological lapse rate using robust regression methods.

### Usage

```
pheno_gradient(
  pep,
  variable = c("alt", "lat"),
  species = NULL,
  phase_id = NULL,
  year_range = NULL,
  by = NULL,
  method = c("robust", "ols", "quantile"),
  aggregate = TRUE,
  na.rm = TRUE
)
```

### Arguments

pep	A pep object or data.table containing phenological observations with columns year, day, and either alt (altitude) or lat (latitude).
variable	Character string specifying the gradient variable. Either "alt" (altitude/elevation) or "lat" (latitude). Default is "alt".
species	Optional character string to filter by species/genus.
phase_id	Optional integer to filter by a single BBCH phase code.
year_range	Optional integer vector of length 2 specifying years to include.
by	Optional character vector of grouping variables (e.g., "country"). If provided, separate gradients are calculated for each group.
method	Character string specifying regression method: <ul style="list-style-type: none"> <li>"robust" (default): Robust regression using robustbase::lmrob</li> <li>"ols": Ordinary least squares regression</li> <li>"quantile": Quantile regression for median using quantreg::rq</li> </ul>
aggregate	Logical. If TRUE (default), first aggregate to station means before fitting. If FALSE, use all individual observations.
na.rm	Logical. Should missing values be removed? Default TRUE.

## Details

Phenological gradients describe how the timing of biological events changes with environmental factors. This function calculates:

- **Altitude gradient:** Typically 2-4 days delay per 100m elevation gain in temperate regions
- **Latitude gradient:** Typically 2-5 days delay per degree north in Europe

The robust regression method (default) is recommended because phenological data often contains outliers from observation errors or microclimate effects.

## Value

A list with class `pheno_gradient` containing:

**summary** A `data.table` with slope, intercept, R-squared, RMSE, `n_stations`, and interpretation

**model** The fitted model object(s)

**data** The data used for fitting

**variable** The gradient variable used

**method** The regression method used

## Interpretation

The slope represents:

- For altitude: days delay per 100 meters elevation increase
- For latitude: days delay per degree latitude increase

Positive slopes indicate later phenology at higher elevations/latitudes.

## Author(s)

Matthias Templ

## References

Defila, C. and Clot, B. (2001). Phytophenological trends in Switzerland. *International Journal of Biometeorology*, 45, 203–207. doi:[10.1007/s004840100101](https://doi.org/10.1007/s004840100101)

Ziello, C. et al. (2009). Influence of altitude on phenology of selected plant species in the Alpine region (1971–2000). *Climate Research*, 39, 227–234. doi:[10.3354/cr00822](https://doi.org/10.3354/cr00822)

## See Also

[pheno\\_normals](#) for calculating climatological normals, [lmrob](#) for robust regression details

**Examples**

```

pep <- pep_download()

# Subset to Alpine countries (good elevation variation)
pep_subset <- pep[country %in% c("Switzerland", "Austria")]

# Altitude gradient for apple flowering
grad_alt <- pheno_gradient(pep_subset,
                           variable = "alt",
                           species = "Malus",
                           phase_id = 60)

print(grad_alt)

# Latitude gradient
grad_lat <- pheno_gradient(pep_subset,
                           variable = "lat",
                           species = "Malus",
                           phase_id = 60)

print(grad_lat)

# Compare regression methods
grad_ols <- pheno_gradient(pep_subset, species = "Malus",
                           phase_id = 60, method = "ols")

print(grad_ols)

grad_robust <- pheno_gradient(pep_subset, species = "Malus",
                              phase_id = 60, method = "robust")

print(grad_robust)

# Gradient by country (separate regression per country)
grad_by_country <- pheno_gradient(pep_subset,
                                  variable = "alt",
                                  species = "Malus",
                                  phase_id = 60,
                                  by = "country")

print(grad_by_country)

```

---

pheno\_leaflet

*Interactive Leaflet Map to Select PEP725 Stations*


---

**Description**

Launches a Shiny gadget that allows selection of PEP phenological stations by drawing a bounding box or clicking individual points on the map.

**Usage**

```
pheno_leaflet(pep, label_col = NULL, quiet = FALSE)
```

**Arguments**

pep	A data frame or data.table with columns lon, lat, and optionally species.
label_col	Optional column name to show as label in popups (e.g. "species" or "s_id").
quiet	Logical. If FALSE (default), displays a message about expected loading time.

**Details**

**Note:** Loading time depends on the size of the dataset. For the full PEP725 dataset with millions of observations, expect 10-20 seconds for the map to fully render. Consider filtering the data before calling this function to speed up loading:

```
# Filter to a specific species first
wheat <- pep[genus == "Triticum"]
selected <- pheno_leaflet(wheat)
```

**Value**

A data.frame with selected stations (columns: lon, lat, and metadata).

**Author(s)**

Matthias Templ

**Examples**

```
if (interactive()) {
  pep <- pep_download()

  # For faster loading, filter the data first (recommended)
  # Grapevine has the longest historical records
  pep_subset <- pep[species == "Vitis vinifera"]

  if (nrow(pep_subset) > 0) {
    # Launch interactive selection gadget
    selected <- pheno_leaflet(pep_subset, label_col = "species")

    # Inspect selected subset
    print(selected)
  }
}
```

pheno\_map

*Plot Phenology Station Maps***Description**

Generates a map of PEP725 phenological station locations with optional coloring by various statistics including mean phenological timing, trends, and species variation.

**Usage**

```
pheno_map(
  pep,
  location = c(lon = 6.233, lat = 46.4),
  zoom = 4,
  background = c("google", "none"),
  color_by = c("none", "n_obs", "n_species", "mean_doy", "trend", "species_cv"),
  phase_id = NULL,
  period = NULL,
  min_years = 10,
  min_species = 3,
  point_size = 0.8,
  output_file = NULL,
  key = NULL
)
```

**Arguments**

pep	A data.table or data.frame with phenological observations. Required columns depend on color_by: <ul style="list-style-type: none"> <li>• Always required: lon, lat</li> <li>• For "n_species", "mean_doy", "species_cv": species</li> <li>• For "mean_doy", "trend": year, day</li> </ul>
location	Named vector with center longitude and latitude for the map. Defaults to near Changins: c(lon = 6.233, lat = 46.400). Used primarily for background = "google".
zoom	Zoom level for the map. For background = "google", this is the Google Maps zoom level (4 = Europe, 7 = regional). For background = "none", this controls the padding around data extent (4 = wide view, 7 = tight view). Default is 4.
background	Character. Map background type: <p><b>"google"</b> Google Maps satellite/terrain background. Requires API key (set via ggmap::register_google() or key parameter).</p> <p><b>"none"</b> Simple map with country borders from Natural Earth data. No API key required. Good for publications and vignettes.</p>
color_by	Character. What to color stations by:

	" <b>none</b> "	Black points showing station locations only
	" <b>n_obs</b> "	Number of observations per station
	" <b>n_species</b> "	Number of species recorded per station
	" <b>mean_doy</b> "	Mean day-of-year (phenological timing) per station
	" <b>trend</b> "	Trend in days/year per station (positive = later, negative = earlier)
	" <b>species_cv</b> "	Coefficient of variation in timing across species at each station (higher = more variation among species)
phase_id	Integer.	B BCH phase code to filter by. Recommended for "mean_doy" and "trend" to ensure meaningful comparisons. If NULL (default), all phases are included.
period	Integer vector	of years to include. Default is all years. For trend calculation, at least 10 years are recommended.
min_years	Integer.	Minimum years required for trend calculation. Stations with fewer years show as NA. Default is 10.
min_species	Integer.	Minimum species required for species_cv. Default is 3.
point_size		Size of station points (default: 0.8).
output_file		Optional file path to export the plot (e.g. "map.pdf").
key		Google Maps API key. Only needed for background = "google". You can set globally via <code>ggmap::register_google()</code> instead.

## Details

The function supports two background types:

**Google Maps background** (`background = "google"`): Provides detailed satellite or terrain imagery but requires a Google Maps API key. Register at <https://console.cloud.google.com/> and enable the Maps Static API.

**No background** (`background = "none"`): Uses country outlines from Natural Earth data. No API key required, making it suitable for:

- CRAN package vignettes (no external API calls)
- Reproducible research (no authentication needed)
- Publication-quality maps with clean appearance

The color options provide insights into:

- **mean\_doy**: Spatial patterns in phenological timing. Earlier timing (lower DOY) typically in southern/lowland areas.
- **trend**: Where phenology is advancing (negative = earlier over time, shown in blue) or delaying (positive, shown in red). Uses Kendall's tau normalized statistic for robustness.
- **species\_cv**: Stations where different species show similar timing (low CV) vs. divergent timing (high CV). High variation may indicate species-specific responses to local conditions.

## Value

A ggplot map object.

### Interpreting Trends

The trend is calculated as Kendall's normalized tau statistic, which ranges roughly from -3 to +3 for typical data:

- Negative values (blue): Phenology is getting earlier over time
- Positive values (red): Phenology is getting later over time
- Values near 0: No clear trend
- $|t_{\text{aul}}| > 1.96$ : Statistically significant at 95% level

### Author(s)

Matthias Templ

### See Also

[pheno\\_normals](#) for detailed normal calculations, [mann\\_kendall\\_z](#) for trend calculation, [pheno\\_synchrony](#) for synchrony analysis

### Examples

```
pep <- pep_download()
pep_alpine <- pep[country %in% c("Switzerland", "Austria")]
pheno_map(pep_alpine, background = "none", color_by = "n_obs")

## Not run:
# With Google Maps background (requires API key)
ggmap::register_google(key = "your_api_key_here")
pheno_map(pep, background = "google", color_by = "n_species", zoom = 5)

## End(Not run)
```

---

pheno\_normals

*Calculate Phenological Normals (Climatology)*

---

### Description

Computes reference "normal" phenology for a specified period, analogous to WMO climate normals. Returns central tendency, spread, and percentile statistics for phenological day-of-year (DOY) values.

### Usage

```
pheno_normals(
  pep,
  period = 1991:2020,
  by = c("country", "genus", "phase_id"),
  species = NULL,
```

```

    phase_id = NULL,
    min_years = 20,
    probs = c(0.05, 0.1, 0.25, 0.75, 0.9, 0.95),
    na.rm = TRUE
  )

```

### Arguments

pep	A pep object or data.table containing phenological observations with columns year, day, and grouping variables.
period	Integer vector specifying the years to include in the normal calculation. Default is 1991:2020 (current WMO standard period).
by	Character vector of column names to group by. Common choices: <ul style="list-style-type: none"> <li>• c("country", "genus", "phase_id") - Regional normals by genus/phase</li> <li>• c("s_id", "species", "phase_id") - Station-level normals</li> <li>• c("genus", "phase_id") - Overall normals ignoring geography</li> </ul> Default is c("country", "genus", "phase_id").
species	Optional character string to filter by species column. Can be genus name (e.g., "Triticum") or full species (e.g., "Triticum aestivum"). If NULL (default), all species in the data are included.
phase_id	Optional integer vector to filter by BBCH phase codes. If NULL (default), all phases in the data are included.
min_years	Minimum number of years required to calculate valid normals. Default is 20 (WMO standard). Groups with fewer years return NA.
probs	Numeric vector of probabilities for percentile calculation. Default is c(0.05, 0.10, 0.25, 0.75, 0.90, 0.95).
na.rm	Logical. Should missing values be removed? Default TRUE.

### Details

Phenological normals provide a baseline for comparing individual years or detecting trends. This function calculates both classical statistics (mean, SD) and robust alternatives (median, MAD, IQR) that are less sensitive to outliers.

The function can be used in two ways:

1. **Full dataset:** Pass the complete pep object and use species and phase\_id parameters to filter internally.
2. **Pre-filtered subset:** Filter the data first using data.table syntax, then pass the subset to the function.

### Value

A data.table with the following columns:

**by variables** Grouping variables as specified in by  
**n\_years** Number of years with data in the period

**n\_obs** Total number of observations  
**mean\_doy** Arithmetic mean DOY  
**median\_doy** Median DOY (more robust to outliers)  
**sd\_doy** Standard deviation  
**iqr\_doy** Interquartile range  
**mad\_doy** Median absolute deviation (robust spread)  
**q05, q10, q25, q75, q90, q95** Percentiles with the default probs. Custom probs produce column names derived from the actual levels — integer percents pad to two digits (e.g., q05, q10), fractional percents use an underscore in place of the decimal point (e.g., q02\_5 for the 2.5\ The mapping is also stored in `attr(result, "q_names")`.  
**period** Character string describing the reference period

### Standard Reference Periods

- **1961-1990**: Historical reference (pre-acceleration of warming)
- **1991-2020**: Current WMO standard normal period

### Author(s)

Matthias Templ

### References

WMO (2017). *WMO Guidelines on the Calculation of Climate Normals*. WMO-No. 1203, World Meteorological Organization, Geneva.

Templ, M., Templ, B., and Filzmoser, P. (2018). Modelling and prediction of phenological data of the beginning of flowering: Austrian indicators of climate change. *International Journal of Biometeorology*, 62, 1319–1334. doi:10.1007/s0048401815342

### See Also

[pheno\\_anomaly](#) for calculating anomalies relative to normals, [pep\\_download](#) for obtaining the main dataset

### Examples

```
# Download synthetic data first
pep <- pep_download()

# Use Swiss subset for faster computation
pep_ch <- pep[country == "Switzerland"]

# Calculate normals for all species and phases
normals_all <- pheno_normals(pep_ch)

# Normals for apples, flowering phase (60)
apple_normals <- pheno_normals(pep_ch,
                              species = "Malus",
```

```
                                phase_id = 60)

# Using a pre-filtered subset
apples <- pep_ch[genus == "Malus" & phase_id %in% c(60, 65)]
apple_normals <- pheno_normals(apples,
                              by = c("species", "phase_id"))

# Station-level normals for detailed analysis
station_normals <- pheno_normals(pep_ch,
                                species = "Malus",
                                phase_id = 60,
                                by = c("s_id"))

# Historical reference period
historical <- pheno_normals(pep_ch,
                           period = 1961:1990,
                           species = "Malus")

# Compare two periods
period1 <- pheno_normals(pep_ch, period = 1961:1990, species = "Malus")
period2 <- pheno_normals(pep_ch, period = 1991:2020, species = "Malus")
shift <- period2$mean_doy - period1$mean_doy
```

---

pheno\_plot

*Plot Phenological Time Series*

---

## Description

Generates time series plots from processed phenology data. Shows DOY trends with faceting by phenophase and spatial scope.

## Usage

```
pheno_plot(data_list, alpha_lines = 0.6, linewidth = 0.7)
```

## Arguments

data_list	A named list of prepared data objects, typically output from <a href="#">pheno_regional</a> . Must contain element <code>ts_tidy</code> .
alpha_lines	Alpha transparency for time series lines (default is 0.6).
linewidth	Line width for time series lines (default is 0.7).

## Details

This function visualizes DOY trends with faceting by phenophase and spatial scope.

**Value**

A ggplot object for visual inspection and further customization.

**Author(s)**

Matthias Templ

**See Also**

[pheno\\_regional](#)

**Examples**

```
# Construct a minimal data_list with ts_tidy
ts_tidy <- data.frame(
  year = rep(1990:1995, 2),
  DOY = c(120, 118, 115, 119, 113, 110, 125, 122, 120, 124, 118, 115),
  source = rep(c("PEP725 (aggregated)", "PEP725 (near Changins)"), each = 6),
  phase = "Heading",
  panel = "DOY"
)
pheno_plot(list(ts_tidy = ts_tidy))
```

---

pheno\_plot\_hh

*Plot Phenological Time Series for Heading and Harvest*

---

**Description**

Visualizes phenological time series for heading and harvest phases, comparing aggregated and spatially filtered PEP725 data.

**Usage**

```
pheno_plot_hh(
  data_list,
  phase_select = NULL,
  alpha_lines = 0.6,
  linewidth = 0.7
)
```

**Arguments**

data_list	A named list returned by <a href="#">pheno_regional_hh</a> , containing ts_tidy.
phase_select	Character. Optional filter for a specific phenological phase (e.g., "Heading", "Harvest"). If NULL, all phases are shown.
alpha_lines	Numeric. Transparency level for lines (default is 0.6).
linewidth	Numeric. Line width for time series plots (default is 0.7).

**Details**

This function shows DOY trends by phase and data source across years, with month labels on the y-axis for easier interpretation.

**Value**

A ggplot object.

**Author(s)**

Matthias Templ

**See Also**

[pheno\\_regional\\_hh](#)

**Examples**

```
pep <- pep_download()
out <- pheno_regional_hh(pep,
  species_name = "Triticum aestivum")
pheno_plot_hh(out)
```

---

pheno\_plot\_timeseries *Plot Phenological Time Series*

---

**Description**

Visualizes phenological day-of-year (DOY) trends over time since 1961. Designed for aggregated or site-level phenology data from PEP725.

**Usage**

```
pheno_plot_timeseries(
  data,
  color_by = "species",
  facet_by = NULL,
  alpha_lines = 0.7,
  linewidth = 0.8,
  smooth = TRUE,
  se = FALSE,
  year_min = 1961,
  title = "Phenological time series of flowering (BBCH 65)"
)
```

**Arguments**

<code>data</code>	A <code>data.frame</code> or <code>data.table</code> containing at least <code>year</code> , <code>day</code> (day-of-year), and one grouping variable (e.g. <code>species</code> , <code>phase</code> , or <code>functional_group</code> ).
<code>color_by</code>	Character. Column name used for color grouping (default = "species").
<code>facet_by</code>	Character or <code>NULL</code> . Optional column name for faceting (e.g. "functional_group").
<code>alpha_lines</code>	Numeric. Transparency level for time-series lines (default = 0.7).
<code>linewidth</code>	Numeric. Line width (default = 0.8).
<code>smooth</code>	Logical. If <code>TRUE</code> , adds a linear trend smoother (default = <code>TRUE</code> ).
<code>se</code>	Logical. Whether to display confidence interval for smoother (default = <code>FALSE</code> ).
<code>year_min</code>	Numeric. Minimum year to plot (default = 1961).
<code>title</code>	Character. Optional plot title.

**Value**

A `ggplot` object.

**Author(s)**

Matthias Templ

**Examples**

```

pep <- pep_download()

# Use Alpine subset for faster computation
pep_alpine <- pep[country %in% c("Switzerland", "Austria")]

# Example: flowering DOY by genus
pheno_plot_timeseries(data = pep_alpine[phase_id == 65],
                      color_by = "genus",
                      facet_by = NULL, smooth = TRUE)

# Example: single species, colored by site (grapevine has longest records)
vine_data <- pep_alpine[species == "Vitis vinifera" & phase_id == 65]
if (nrow(vine_data) > 0) {
  pheno_plot_timeseries(
    data = vine_data,
    color_by = "s_id",
    smooth = TRUE,
    title = "Grapevine flowering (BBCH 65) across sites"
  )
}

# Example: facets by genus
pheno_plot_timeseries(
  data = pep_alpine[phase_id == 65],
  color_by = "species",
  facet_by = "genus",

```

```

    smooth = TRUE
  )

```

---

 pheno\_pls

*Partial Least Squares Analysis for Phenology-Temperature Relationships*


---

### Description

Performs PLS regression to identify temperature-sensitive periods affecting phenological timing. This is useful for understanding which parts of the year have the strongest influence on phenological events.

### Usage

```

pheno_pls(
  pep,
  temp_data,
  by = NULL,
  method = c("robust", "standard"),
  split_month = 6,
  runn_mean = 11,
  ncomp = NULL,
  expl_var = 30,
  max_iter = 20,
  tol = 1e-04
)

```

### Arguments

pep	A pep object or data.frame with phenological observations. Must contain year and day columns.
temp_data	A data.frame with daily temperature data. Must contain columns year, doy (day of year), and either Tmean or both Tmin and Tmax.
by	Character vector of grouping columns to match phenology and temperature data (e.g., "s_id" for station-specific analysis). Default NULL uses the same temperature for all observations per year.
method	Character. PLS method to use: <b>"robust"</b> (Default) Iteratively reweighted PLS using bisquare weights. Resistant to outliers in both phenology and temperature data. <b>"standard"</b> Classical PLS regression without robust weighting.
split_month	Integer (1-12). The last month to include in the phenological year. Default 6 (June) means July-June phenological years. This is important for spring phenology where relevant temperatures span the previous autumn/winter.

<code>runn_mean</code>	Integer (odd). Window size for running mean smoothing of temperatures. Default 11 days. Larger values produce smoother results.
<code>ncomp</code>	Integer or NULL. Number of PLS components. If NULL (default), automatically determined to explain at least <code>expl_var</code> percent of variance.
<code>expl_var</code>	Numeric. Minimum percentage of variance to explain when automatically selecting components. Default 30.
<code>max_iter</code>	Integer. Maximum iterations for robust method. Default 20.
<code>tol</code>	Numeric. Convergence tolerance for robust method. Default 1e-4.

### Details

PLS regression is particularly useful for phenology-temperature analysis because it handles the high collinearity between daily temperatures (adjacent days are highly correlated) while identifying which periods have the strongest influence on phenological timing.

The robust method uses iteratively reweighted PLS with bisquare weights, which downweights observations with large residuals. This is important for phenological data which may contain outliers from observation errors or unusual weather events.

### Value

An object of class `pheno_pls` containing:

**vip** Data.frame with Variable Importance in Projection scores for each day of the phenological year.

**coefficients** PLS regression coefficients for each day.

**model** The fitted PLS model object.

**weights** Observation weights (all 1 for standard method).

**r\_squared** Model R-squared.

**ncomp** Number of components used.

**method** Method used ("robust" or "standard").

**n\_obs** Number of observations.

**n\_years** Number of years analyzed.

### Interpreting Results

**VIP scores** Values > 0.8 indicate important variables. Days with high VIP scores have strong influence on phenology timing.

**Coefficients** Negative coefficients indicate that warmer temperatures during that period lead to earlier phenology (lower DOY).

### Phenological Year

Spring phenology (e.g., flowering) is influenced by temperatures from the previous autumn through spring. The `split_month` parameter defines where to split the calendar year. With `split_month = 6`, a flowering event in April 2020 is analyzed against temperatures from July 2019 through June 2020.

**Author(s)**

Matthias Templ

**References**

Luedeling E, Gassner A (2012). Partial Least Squares Regression for analyzing walnut phenology in California. *Agricultural and Forest Meteorology* 158-159:104-113.

Wold S, Sjoström M, Eriksson L (2001). PLS-regression: a basic tool of chemometrics. *Chemometrics and Intelligent Laboratory Systems* 58:109-130.

**See Also**

[pheno\\_trend\\_turning](#) for trend analysis, [pheno\\_anomaly](#) for anomaly detection

**Examples**

```
data(pep_seed)

# Create example temperature data (fewer years for speed)
years <- 2005:2012
temp <- data.frame(
  year = rep(years, each = 365),
  doy = rep(1:365, length(years)),
  Tmin = rnorm(365 * length(years), mean = 5, sd = 8),
  Tmax = rnorm(365 * length(years), mean = 15, sd = 8)
)

# Run standard PLS (fast: no CV, fixed components)
result <- pheno_pls(pep_seed, temp, method = "standard", ncomp = 2)
print(result)
```

---

pheno\_regional

*Compile Regional Phenology Data and Climate Sensitivity Inputs*

---

**Description**

Extracts and aggregates phenological observations for a selected species and phenological phase from the PEP725 dataset (globally and spatially filtered). Optionally links them with global temperature anomalies (GISS data) for climate impact studies.

**Usage**

```
pheno_regional(
  pep,
  giss = NULL,
  lon_min = 4.2,
```

```

lon_max = 8.1,
lat_min = 44.7,
lat_max = 48.1,
species_name = "Triticum aestivum",
functional_group = NULL,
year_min = 1961,
pep_for_giss = c("near", "aggregated"),
phase = 60
)

```

### Arguments

pep	A data.table containing the full PEP725 dataset. Must include at least columns: species, year, phase_id, day, lat, lon.
giss	Optional. A data.table with GISS global temperature anomalies, containing columns year, dT, and dT_sm. If NULL, GISS-related outputs are omitted.
lon_min	Minimum longitude for spatial filtering of PEP data (default is 4.2)
lon_max	Maximum longitude for spatial filtering of PEP data (default is 8.1)
lat_min	Minimum latitude for spatial filtering of PEP data (default is 44.7)
lat_max	Maximum latitude for spatial filtering of PEP data (default is 48.1)
species_name	Character name of the species to be extracted (default is "Triticum aestivum")
functional_group	Character. Optional functional group name to filter by (e.g., "C4_summer_cereals"). If provided, overwrites species_name filtering.
year_min	Minimum year to include in all outputs (default is 1961)
pep_for_giss	Selects which PEP data subset to use for merging with GISS data. Either "near" (box-filtered) or "aggregated" (global). Ignored if giss is NULL.
phase	Integer or vector of phase_id(s) to extract (default is 60 = Heading). Named phase mappings are applied.

### Details

The function aggregates PEP observations by species, year, and phase, calculates mean day-of-year (DOY), applies spatial filtering if needed, and attaches GISS climate anomalies. It also maps phase\_ids to descriptive names and warns if expected phenological phase are missing in the selected data.

If GISS data is provided, temperature anomalies are merged with the selected PEP time series to support phenology-climate analysis.

### Value

A named list with the following elements:

- pep\_agg Global (non-spatially filtered) PEP725 time series for the selected phase(s)
- pep CGI Spatially filtered PEP725 time series based on lat/lon bounding box
- giss Processed GISS global temperature anomaly data (if provided)

pep\_giss Merged data frame of PEP phenology and GISS anomalies (if provided)  
species Character species name used for filtering  
functional\_group Functional group if specified, otherwise NA  
phase Integer phase ID(s) used for filtering

**Author(s)**

Matthias Templ

**See Also**

[pheno\\_plot](#), [pep\\_download](#)

**Examples**

```
pep <- pep_download()

out <- pheno_regional(pep, species_name = "Triticum aestivum", phase = 60)
str(out)
```

---

pheno\_regional\_hh      *Compile Regional Phenological Time Series for Heading and Harvest*

---

**Description**

Generates harmonized phenological time series for heading (BBCH 60) and harvest (BBCH 100) phases of a specified species and spatial region. Optionally integrates GISS climate data for climate-sensitivity studies.

**Usage**

```
pheno_regional_hh(  
  pep,  
  giss = NULL,  
  lon_min = 4.2,  
  lon_max = 8.1,  
  lat_min = 44.7,  
  lat_max = 48.1,  
  species_name = "Triticum aestivum",  
  year_min = 1961,  
  pep_for_giss = c("aggregated", "near")  
)
```

**Arguments**

pep	A data.table containing PEP725 phenological data.
giss	Optional. A data.table with GISS global temperature anomalies (year, dT, dT_sm). If NULL, GISS-related outputs are omitted.
lon_min	Minimum longitude for bounding box filter (default is 4.2)
lon_max	Maximum longitude for bounding box filter (default is 8.1)
lat_min	Minimum latitude for bounding box filter (default is 44.7)
lat_max	Maximum latitude for bounding box filter (default is 48.1)
species_name	Name of the species to filter (default is "Triticum aestivum")
year_min	Minimum year for all time series outputs (default is 1961)
pep_for_giss	Which PEP725 dataset to use for merging with GISS climate anomalies. Either "aggregated" (global) or "near" (spatially filtered). Ignored if giss is NULL.

**Details**

This function filters the PEP725 dataset by species and optionally by spatial box. It aggregates phenological phases (Heading = 60, Harvest = 100) and computes mean day-of-year (DOY) per year. If GISS data is provided, temperature anomalies are merged with the selected PEP time series to support phenology-climate analysis. Warnings are issued if phases are missing for selected periods.

**Value**

A named list with the following elements:

ts_tidy	Long-format time series with consistent phase and source labeling
pep_agg	Aggregated PEP725 time series (global species-level)
pep CGI	Spatially filtered PEP725 time series (based on lat/lon box)
giss	GISS global temperature anomalies (if provided)
pep_giss	PEP time series merged with GISS anomalies (if provided)
species	Species name used for filtering

**Author(s)**

Matthias Templ

**See Also**

[pep\\_import](#), [pheno\\_plot\\_hh](#)

**Examples**

```
pep <- pep_download()

out <- pheno_regional_hh(pep,
  species_name = "Triticum aestivum")
pheno_plot_hh(out)
```

---

pheno\_synchrony      *Analyze Phenological Synchrony Across Stations*

---

### Description

Measures how synchronized phenological events are across stations within regions and whether synchrony is changing over time. Higher synchrony indicates more uniform timing of phenological events.

### Usage

```
pheno_synchrony(
  pep,
  species = NULL,
  phase_id = NULL,
  by = c("country", "year"),
  min_stations = 5,
  compute_trend = TRUE,
  na.rm = TRUE
)
```

### Arguments

pep	A pep object or data.table containing phenological observations with columns year, day, and s_id (station ID).
species	Optional character string to filter by species/genus.
phase_id	Optional integer to filter by a single BBCH phase code.
by	Character vector of column names to group by for synchrony calculation. Default is c("country", "year") for annual synchrony by country.
min_stations	Minimum number of stations required to calculate synchrony. Default is 5. Groups with fewer stations return NA.
compute_trend	Logical. If TRUE (default), computes trend in synchrony over time using robust regression.
na.rm	Logical. Should missing values be removed? Default TRUE.

### Details

Synchrony measures how similar phenological timing is across different stations within the same region and year. This is important for:

- Understanding spatial coherence of phenological signals
- Detecting changes in spatial variability over time
- Assessing network representativeness

**Value**

A pheno\_synchrony object (list) containing:

**data** A data.table with synchrony metrics per group: n\_stations, mean\_doy, sd\_doy, cv\_pct, and quality indicators

**trend** If compute\_trend = TRUE, a data.table with trend analysis results per region (slope, p-value, direction)

**overall** Summary statistics across all groups

**Synchrony Metrics**

**sd\_doy** Standard deviation across stations - lower values indicate higher synchrony

**cv\_pct** Coefficient of variation (SD/mean \* 100) - relative measure that allows comparison across phases with different mean timing

**range\_doy** Range of DOY values across stations

**Trend Interpretation**

- Negative trend in SD: Increasing synchrony over time
- Positive trend in SD: Decreasing synchrony (more variable)

**Author(s)**

Matthias Templ

**References**

Rosenzweig, C. et al. (2008). Attributing physical and biological impacts to anthropogenic climate change. *Nature*, 453, 353–357. doi:10.1038/nature06937

**See Also**

[pheno\\_normals](#) for climatological statistics, [pheno\\_anomaly](#) for anomaly detection

**Examples**

```
pep <- pep_download()

# Subset to two countries for speed
pep_subset <- pep[country %in% c("Switzerland", "Austria")]

# Calculate synchrony for apple flowering by country and year
sync <- pheno_synchrony(pep_subset,
                        species = "Malus",
                        phase_id = 60)

print(sync)

# Get trend results (robust regression per country)
sync$trend
```

```

# Synchrony without trend analysis (faster)
sync_simple <- pheno_synchrony(pep_subset,
                             species = "Malus",
                             phase_id = 60,
                             compute_trend = FALSE)

sync_simple

# Custom grouping variables
sync_detailed <- pheno_synchrony(pep_subset,
                                 species = "Malus",
                                 phase_id = 60,
                                 by = c("country", "year"))

sync_detailed

```

---

pheno\_trend\_turning     *Detect Trend Turning Points in Phenological Time Series*

---

## Description

Applies a sequential Mann-Kendall test to detect approximate trend turning points in phenological time series. This identifies years where trends may have changed direction (e.g., when spring advancement accelerated or reversed).

## Usage

```
pheno_trend_turning(pep, by = NULL, min_years = 10, aggregate = TRUE)
```

## Arguments

pep	A pep object, data.table, or numeric vector of DOY values. If a data.frame/data.table, must contain year and day columns.
by	Character vector of column names to group by before analysis. Default is NULL (analyze all data as one series). Common choices: c("genus", "species", "phase_id") or c("country", "phase_id").
min_years	Integer. Minimum number of years required for analysis. Default is 10. Shorter series may produce unreliable results.
aggregate	Logical. If TRUE (default), aggregates multiple observations per year (e.g., from multiple stations) using the median. If FALSE, expects exactly one value per year.

## Details

The sequential Mann-Kendall test calculates two series:

- **Progressive:** Kendall's tau computed from the start forward

- **Retrograde:** Kendall's tau computed from the end backward

Points where these series cross indicate potential trend turning points. When either series exceeds confidence thresholds ( $t_{\text{tau}} > 1.96$  for 95%,  $t_{\text{tau}} > 2.58$  for 99%) before and after a crossing, the turning point is considered statistically significant.

### Value

An object of class `pheno_turning` containing:

**results** Data.table with columns: year, day (or median\_day), tau\_prog (progressive tau), tau\_retr (retrograde tau), is\_turning (logical indicating turning points)

**turning\_points** Years identified as potential turning points

**n\_years** Number of years in the series

**by** Grouping variables used (if any)

**group\_results** If by is specified, list of results per group

### Interpretation

- Positive tau indicates an increasing trend (later DOY = delayed phenology)
- Negative tau indicates a decreasing trend (earlier DOY = advanced phenology)
- Crossing points suggest the trend direction may have changed

### Author(s)

Matthias Templ

### References

Sneyers R (1990). On statistical analysis of series of observations. Technical Note No 143. World Meteorological Organization.

### See Also

[pheno\\_gradient](#) for spatial trend analysis, [pheno\\_normals](#) for baseline calculations

### Examples

```
# Simple vector input (fast)
doy_series <- c(120, 118, 122, 115, 110, 108, 112, 105, 102, 100)
turning <- pheno_trend_turning(doy_series)
print(turning)

# Using pep_seed data (no grouping for speed)
data(pep_seed)
vine <- pep_seed[pep_seed$species == "Vitis vinifera" &
                 pep_seed$phase_id == 65, ]
if (nrow(vine) > 0) {
  turning <- pheno_trend_turning(vine)
}
```

---

plot.pep *Plot Method for PEP725 Data*

---

**Description**

Provides default visualizations for PEP725 phenological data.

**Usage**

```
## S3 method for class 'pep'  
plot(x, type = c("map", "timeseries", "histogram"), ...)
```

**Arguments**

x	A pep object.
type	Character. Type of plot: "map" for station locations, "timeseries" for temporal trends, "histogram" for DOY distribution. Default is "map".
...	Additional arguments passed to the underlying plot function.

**Value**

A ggplot object (invisibly).

**Author(s)**

Matthias Templ

**Examples**

```
pep <- pep_download()  
plot(pep)  
plot(pep, type = "timeseries")  
plot(pep, type = "histogram")
```

---

plot.pep\_completeness *Plot Method for Completeness Assessment*

---

**Description**

Creates visualizations of data completeness, including heatmaps and bar charts.

**Usage**

```
## S3 method for class 'pep_completeness'  
plot(x, type = c("heatmap", "bar", "timeline"), top = 20, ...)
```

**Arguments**

x	A pep_completeness object
type	Character. Type of plot: "heatmap" for completeness heatmap (requires 2 grouping variables), "bar" for observation counts, "timeline" for year coverage. Default is "heatmap".
top	Integer. Number of top groups to display. Default is 20.
...	Additional arguments (unused)

**Value**

A ggplot object (invisibly)

**Author(s)**

Matthias Templ

**Examples**

```

pep <- pep_download()
pep_ch <- pep[country == "Switzerland"]
comp <- pep_completeness(pep_ch, by = c("genus", "phase_id"))
plot(comp, type = "heatmap")
plot(comp, type = "bar", top = 15)

```

---

plot.pep\_coverage      *Plot Method for PEP Coverage*

---

**Description**

Plot Method for PEP Coverage

**Usage**

```

## S3 method for class 'pep_coverage'
plot(x, ...)

```

**Arguments**

x	A pep_coverage object.
...	Additional arguments (unused).

**Value**

A ggplot object (or list of ggplot objects).

**Author(s)**

Matthias Templ

---

`plot.pep_outliers`      *Plot Method for Outlier Detection Results*

---

**Description**

Plot Method for Outlier Detection Results

**Usage**

```
## S3 method for class 'pep_outliers'  
plot(x, type = c("histogram", "scatter", "timeline"), ...)
```

**Arguments**

<code>x</code>	A <code>pep_outliers</code> object
<code>type</code>	Character. Plot type: "histogram" for deviation distribution, "scatter" for DOY vs expected, "timeline" for outliers over time.
<code>...</code>	Additional arguments

**Value**

A ggplot object (invisibly)

---

`plot.pep_quality`      *Plot Method for Data Quality Assessment*

---

**Description**

Creates visualizations of phenological data quality metrics, including grade distribution and a map of station quality.

**Usage**

```
## S3 method for class 'pep_quality'  
plot(  
  x,  
  which = c("overview", "grades", "map"),  
  pep = NULL,  
  show_grades = c("A", "B", "C", "D"),  
  alpha = 0.6,  
  title = NULL,  
  ...  
)
```

**Arguments**

x	A pep_quality object.
which	Character. Type of plot to produce: <ul style="list-style-type: none"> <li>• "overview" (default): Two-panel overview with grade distribution and station quality map (requires pep argument)</li> <li>• "grades": Bar chart of quality grade distribution only</li> <li>• "map": Map of station locations colored by quality grade (requires pep argument for coordinates)</li> </ul>
pep	Optional pep object providing station coordinates (lon, lat). Required for which = "map" or which = "overview".
show_grades	Character vector of grades to display on the map. Default is c("A", "B", "C", "D") (all grades). Use e.g. show_grades = "D" to show only poor-quality stations.
alpha	Numeric. Transparency of points on the map (0-1). Default is 0.6.
title	Optional character string for the plot title.
...	Additional arguments (unused).

**Details**

For the map visualization, the function:

- Aggregates quality to one grade per station (worst grade if multiple phases)
- Uses country borders from the `rnatrualearth` package
- Uses colorblind-friendly colors (blue=A, cyan=B, orange=C, vermillion=D)

**Value**

A `ggplot` object (invisibly).

**Author(s)**

Matthias Templ

**Examples**

```
pep <- pep_download()

# Assess quality for Swiss stations
pep_ch <- pep[country == "Switzerland"]
quality <- pep_quality(pep_ch, by = c("s_id", "phase_id"))

# Grade distribution only (no pep data needed)
plot(quality, which = "grades")

# Map of station quality (requires pep for coordinates)
plot(quality, which = "map", pep = pep_ch)
```

```
# Map showing only poor-quality stations
plot(quality, which = "map", pep = pep_ch, show_grades = "D")

# Map showing problematic stations (C and D grades)
plot(quality, which = "map", pep = pep_ch, show_grades = c("C", "D"))

# Overview: grades + map
plot(quality, pep = pep_ch)
```

---

plot.pheno\_anomaly      *Plot Method for Phenological Anomalies*

---

## Description

Visualise phenological anomalies as a colour-coded timeline or as a histogram of anomaly magnitudes.

## Usage

```
## S3 method for class 'pheno_anomaly'
plot(x, which = c("timeline", "histogram"), ...)
```

## Arguments

x	A pheno_anomaly object returned by <a href="#">pheno_anomaly</a> .
which	Character string selecting the plot type: "timeline" (default) shows anomaly_days per year as bars coloured by direction with extreme events marked; "histogram" shows the distribution of anomaly values.
...	Additional arguments (unused).

## Value

A ggplot object (returned invisibly).

## Author(s)

Matthias Templ

## See Also

[pheno\\_anomaly](#), [pheno\\_normals](#)

## Examples

```
pep <- pep_download()
vine_ch <- pep[country == "Switzerland" & species == "Vitis vinifera"]

if (nrow(vine_ch) > 0) {
  a <- pheno_anomaly(vine_ch,
                    baseline_period = 1961:1990,
                    phase_id = 65,
                    by = "phase_id")

  plot(a)
  plot(a, which = "histogram")
}
```

---

plot.pheno\_combined *Plot Method for Combined Time Series*

---

## Description

Plot Method for Combined Time Series

## Usage

```
## S3 method for class 'pheno_combined'
plot(x, type = c("series", "stations", "residuals"), group = NULL, ...)
```

## Arguments

x	A pheno_combined object
type	Character. "series" for combined time series, "stations" for station effects, "residuals" for residual distribution.
group	For grouped analysis, which group to plot.
...	Additional arguments

## Value

A ggplot object (invisibly)

---

plot.pheno\_gradient *Plot Method for Phenological Gradient Analysis*

---

**Description**

Plot Method for Phenological Gradient Analysis

**Usage**

```
## S3 method for class 'pheno_gradient'  
plot(x, ...)
```

**Arguments**

x                    A pheno\_gradient object  
...                   Additional arguments passed to ggplot

**Value**

A ggplot object

**Author(s)**

Matthias Templ

---

plot.pheno\_normals *Plot Method for Phenological Normals*

---

**Description**

Visualise phenological normals as a dot plot with interquartile-range bars or as a bar chart.

**Usage**

```
## S3 method for class 'pheno_normals'  
plot(x, which = c("dotplot", "bar"), ...)
```

**Arguments**

x                    A pheno\_normals object returned by [pheno\\_normals](#).  
which                Character string selecting the plot type: "dotplot" (default) shows median\_doy with Q25–Q75 range via geom\_pointrange; "bar" shows median\_doy as columns with error bars.  
...                   Additional arguments (unused).

**Value**

A ggplot object (returned invisibly).

**Author(s)**

Matthias Templ

**See Also**

[pheno\\_normals](#), [pheno\\_anomaly](#)

**Examples**

```
pep <- pep_download()
pep_ch <- pep[country == "Switzerland"]

# Normals by phase for apple
n <- pheno_normals(pep_ch, species = "Malus", by = "phase_id")
plot(n)
plot(n, which = "bar")
```

---

plot.pheno\_pls

*Plot Method for PLS Phenology Results*


---

**Description**

Plot Method for PLS Phenology Results

**Usage**

```
## S3 method for class 'pheno_pls'
plot(x, type = c("vip", "coef", "both"), vip_threshold = 0.8, ...)
```

**Arguments**

x	A pheno_pls object
type	Character. Plot type: "vip" for VIP scores (default), "coef" for coefficients, "both" for combined plot.
vip_threshold	Numeric. Threshold line for VIP plot. Default 0.8.
...	Additional arguments passed to plotting functions

**Value**

A ggplot object (invisibly)

---

plot.pheno\_synchrony *Plot Method for Phenological Synchrony Analysis*

---

**Description**

Creates a time series plot of synchrony (SD) over years.

**Usage**

```
## S3 method for class 'pheno_synchrony'  
plot(x, region = NULL, ...)
```

**Arguments**

x	A pheno_synchrony object
region	Optional region to plot (for multi-region analyses)
...	Additional arguments passed to ggplot

**Value**

A ggplot object

**Author(s)**

Matthias Templ

---

plot.pheno\_turning *Plot Method for Trend Turning Analysis*

---

**Description**

Plot Method for Trend Turning Analysis

**Usage**

```
## S3 method for class 'pheno_turning'  
plot(x, group = NULL, show_thresholds = TRUE, ...)
```

**Arguments**

x	A pheno_turning object
group	Character. For grouped analysis, which group to plot. If NULL (default), plots the first group or ungrouped results.
show_thresholds	Logical. Show significance threshold lines? Default TRUE.
...	Additional arguments passed to ggplot

**Value**

A ggplot object (invisibly)

---

plot.second\_events      *Plot Method for Second Events Detection*

---

**Description**

Plot Method for Second Events Detection

**Usage**

```
## S3 method for class 'second_events'
plot(
  x,
  type = c("overview", "timeline", "seasonal", "map"),
  scale = c("absolute", "relative"),
  from_year = NULL,
  ...
)
```

**Arguments**

x	A second_events object
type	Character. Plot type: "overview", "timeline", "seasonal", "map"
scale	Character. Scale for y-axis in timeline/overview plots: <b>"absolute"</b> (Default) Show raw counts of second events per year. <b>"relative"</b> Show proportion of second events relative to total observations per year. This accounts for varying data availability over time and reveals whether second events are becoming proportionally more common.
from_year	Integer. If specified, only show data from this year onwards in timeline and overview plots. Useful for focusing on recent trends. Default is NULL (show all years).
...	Additional arguments

**Value**

A ggplot object (invisibly)

---

plot\_phenology\_trends *Plot Robust Phenological Trends*


---

**Description**

This function visualizes long-term phenological trends for one genus or species, using robust MM-regression (`robustbase::lmrob`) applied to mean annual DOYs.

**Usage**

```
plot_phenology_trends(
  pep,
  genus_name = NULL,
  species_name = NULL,
  subspecies_name = NULL,
  phases = c(65, 87),
  common_stations = TRUE,
  combine_regions = FALSE,
  combine_layout = c("vertical", "horizontal"),
  years = 1961:2024,
  calib_years = 1991:2020,
  pred_years = NULL,
  subregions = c("Austria", "Germany-North", "Germany-South", "Switzerland"),
  giss_data = NULL,
  layout = c("country_phase", "phase_country"),
  title = NULL
)
```

**Arguments**

<code>pep</code>	A PEP725-style data frame with columns <code>year</code> , <code>DOY</code> , <code>phase_id</code> , <code>country</code> , <code>s_id</code> , <code>genus</code> , <code>species</code> .
<code>genus_name</code>	Character string specifying a genus to filter (optional).
<code>species_name</code>	Character string specifying a species to filter (optional). If both <code>genus</code> and <code>species</code> are provided, the <code>species</code> filter is applied last.
<code>subspecies_name</code>	Character string specifying a subspecies to filter (optional). Only one of <code>genus_name</code> , <code>species_name</code> , or <code>subspecies_name</code> can be specified.
<code>phases</code>	Integer vector of phenological phase IDs to include (default: <code>c(65, 87)</code> for flowering and maturity of fruit).
<code>common_stations</code>	Logical. If <code>TRUE</code> (default), only stations with observations for all selected phases are included. If <code>FALSE</code> , all stations with any of the selected phases are included.
<code>combine_regions</code>	Logical. If <code>TRUE</code> , combines all regions into a single panel. Default is <code>FALSE</code> .

combine_layout	Character. Layout for combined regions plot: either "vertical" or "horizontal".
years	Numeric vector of years to include in the analysis. Default is 1961:2024.
calib_years	Numeric vector specifying the calibration window for robust regression (default: 1991:2020).
pred_years	Numeric vector of years for projecting future phenology. Default is 1990:2090.
subregions	Character vector of country names to include (default: DACH region).
giss_data	Deprecated. Previously used for climate data integration; now ignored.
layout	Either "country_phase" (default) or "phase_country" to control facet arrangement.
title	Optional plot title. If NULL, the function generates one from genus and species information.

### Details

The function produces a faceted panel plot showing:

- annual mean DOY and interquartile range,
- robust trend lines for past and future periods,
- CTRL (1991-2020) and PGW (2066-2095) climate windows,
- optional layouts (country x phase or phase x country).

The function performs the following steps:

1. Filter PEP data by species, phases, regions, years.
2. Aggregate DOYs by year-country-phase.
3. Fit robust regression  $DOY \sim year$  over the calibration window.
4. Predict past and future trends.
5. Draw ribbon (IQR), annual means, trend lines, climate windows.
6. Facet by country or by phase.

The robust regression uses `robustbase::lmrob`, which is resistant to outliers and non-normality, and therefore ideal for phenological time series.

### Value

A `ggplot` object.

### Author(s)

Matthias Templ

**Examples**

```
pep <- pep_download()
plot_phenology_trends(
  pep,
  species_name = "Vitis vinifera",
  phases = c(65, 81),
  layout = "country_phase"
)
```

---

print.pep

*Print Method for PEP725 Data*

---

**Description**

Displays a concise summary of the PEP725 phenological dataset.

**Usage**

```
## S3 method for class 'pep'
print(x, n = 5, ...)
```

**Arguments**

x	A pep object.
n	Number of rows to display (default 5).
...	Additional arguments passed to print.

**Value**

Invisibly returns x.

**Author(s)**

Matthias Templ

**Examples**

```
pep <- pep_download()
print(pep)
```

print.pep\_completeness

*Print Method for Completeness Assessment*

---

### Description

Print Method for Completeness Assessment

### Usage

```
## S3 method for class 'pep_completeness'  
print(x, n = 15, ...)
```

### Arguments

x	A pep_completeness object
n	Number of rows to display
...	Additional arguments (unused)

### Value

Invisibly returns x

### Author(s)

Matthias Templ

---

print.pep\_coverage

*Print Method for PEP Coverage*

---

### Description

Print Method for PEP Coverage

### Usage

```
## S3 method for class 'pep_coverage'  
print(x, ...)
```

### Arguments

x	A pep_coverage object.
...	Additional arguments (unused).

**Value**

Invisibly returns x.

**Author(s)**

Matthias Templ

---

*print.pep\_outliers*      *Print Method for Outlier Detection Results*

---

**Description**

Print Method for Outlier Detection Results

**Usage**

```
## S3 method for class 'pep_outliers'  
print(x, ...)
```

**Arguments**

x	A pep_outliers object
...	Additional arguments (unused)

**Value**

Invisibly returns x

---

*print.pep\_quality*      *Print Method for Data Quality Assessment*

---

**Description**

Print Method for Data Quality Assessment

**Usage**

```
## S3 method for class 'pep_quality'  
print(x, n = 10, ...)
```

**Arguments**

x	A pep_quality object
n	Number of rows to display
...	Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

---

`print.phase_check`      *Print Method for Phase Check Results*

---

**Description**

Print Method for Phase Check Results

**Usage**

```
## S3 method for class 'phase_check'  
print(x, ...)
```

**Arguments**

<code>x</code>	A <code>phase_check</code> object
<code>...</code>	Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

---

`print.phase_check_multi`  
*Print Method for Multi-Species Phase Check*

---

**Description**

Print Method for Multi-Species Phase Check

**Usage**

```
## S3 method for class 'phase_check_multi'  
print(x, ...)
```

**Arguments**

x                    A phase\_check\_multi object  
...                   Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

---

*print.pheno\_anomaly*    *Print Method for Phenological Anomalies*

---

**Description**

Print Method for Phenological Anomalies

**Usage**

```
## S3 method for class 'pheno_anomaly'  
print(x, n = 10, ...)
```

**Arguments**

x                    A pheno\_anomaly object  
n                    Number of rows to display  
...                   Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

print.pheno\_combined *Print Method for Combined Time Series*

---

**Description**

Print Method for Combined Time Series

**Usage**

```
## S3 method for class 'pheno_combined'  
print(x, ...)
```

**Arguments**

x	A pheno_combined object
...	Additional arguments (unused)

**Value**

Invisibly returns x

---

print.pheno\_gradient *Print Method for Phenological Gradient Analysis*

---

**Description**

Print Method for Phenological Gradient Analysis

**Usage**

```
## S3 method for class 'pheno_gradient'  
print(x, ...)
```

**Arguments**

x	A pheno_gradient object
...	Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

---

print.pheno\_normals     *Print Method for Phenological Normals*

---

**Description**

Print Method for Phenological Normals

**Usage**

```
## S3 method for class 'pheno_normals'  
print(x, n = 10, ...)
```

**Arguments**

x	A pheno_normals object
n	Number of rows to display
...	Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

---

print.pheno\_pls     *Print Method for PLS Phenology Results*

---

**Description**

Print Method for PLS Phenology Results

**Usage**

```
## S3 method for class 'pheno_pls'  
print(x, ...)
```

**Arguments**

x	A pheno_pls object
...	Additional arguments (unused)

**Value**

Invisibly returns x

---

print.pheno\_synchrony *Print Method for Phenological Synchrony Analysis*

---

**Description**

Print Method for Phenological Synchrony Analysis

**Usage**

```
## S3 method for class 'pheno_synchrony'  
print(x, n = 10, ...)
```

**Arguments**

x	A pheno_synchrony object
n	Number of rows to display
...	Additional arguments (unused)

**Value**

Invisibly returns x

**Author(s)**

Matthias Templ

---

print.pheno\_turning *Print Method for Trend Turning Analysis*

---

**Description**

Print Method for Trend Turning Analysis

**Usage**

```
## S3 method for class 'pheno_turning'  
print(x, ...)
```

**Arguments**

x	A pheno_turning object
...	Additional arguments (unused)

**Value**

Invisibly returns x

---

```
print.second_events Print Method for Second Events Detection
```

---

**Description**

Print Method for Second Events Detection

**Usage**

```
## S3 method for class 'second_events'
print(x, ...)
```

**Arguments**

x	A second_events object
...	Additional arguments (unused)

**Value**

Invisibly returns x

---

```
select_phase Select and Label Phenophases from PEP Time Series
```

---

**Description**

Extracts a subset of phenological observations from a processed PEP time series (e.g., output from species-level aggregation), filters by species, year, and phenophase ID, and attaches meaningful phase labels based on standard BBCH codes.

**Usage**

```
select_phase(dt, label, sp, yrmin, phases = c(60, 100))
```

**Arguments**

dt	A data.table containing columns species, year, phase_id, and mean_day. Typically produced by aggregating PEP725 data (e.g., via <a href="#">pheno_regional</a> ).
label	A character string used as identifier for source and site columns in the output.
sp	Species name (as character string) to filter, e.g., "Triticum aestivum".
yrmin	Minimum year (inclusive) for filtering.
phases	Integer vector of phase IDs to include (default: c(60, 100) corresponding to "Heading" and "Harvest").

## Details

This function uses an internal lookup table to map numeric `phase_id` codes to descriptive BBCH phase names (e.g., 60 = "Heading", 100 = "Harvest"). Unmapped phase IDs will trigger a warning. If expected phases are missing from the result, a separate warning is issued.

The function is particularly useful for visualizing specific growth stages or comparing phenological trends across datasets and locations.

## Value

A `data.table` with columns:

`year` Year of observation

`phase` Mapped BBCH phenological stage name

`DOY` Mean day-of-year for this phase, species, and year

`source` Set to `label`

`site` Set to `label`

## Author(s)

Matthias Templ

## See Also

[pheno\\_regional](#), [pep\\_download](#), [pheno\\_plot](#)

## Examples

```
pep <- pep_download()
agg <- pheno_regional(pep, species_name = "Triticum aestivum", phase = 60)$pep_agg
df <- select_phase(agg, label = "PEP725", sp = "Triticum aestivum", yrmin = 1961)
head(df)
```

## Description

This function produces a complete report for phenological subspecies availability, combining: (1) a tidy summary table, (2) a publication-ready wide table, and (3) a `ggplot2` heatmap showing data presence/absence.

**Usage**

```
subspecies_report(
  pep,
  genus_name = NULL,
  species_name = NULL,
  subspecies_name = NULL,
  phases = c(65, 87),
  years = 1961:2024,
  subregions = c("Austria", "Germany-North", "Germany-South", "Switzerland"),
  include_empty = TRUE,
  metric = "all",
  bbch_names = NULL,
  return_results = TRUE
)
```

**Arguments**

pep	A PEP725-style data frame containing at least: genus, species, subspecies, country, year, phase_id, s_id, and day or DOY.
genus_name	Character (optional). Genus to filter.
species_name	Character (optional). Species to filter.
subspecies_name	Character vector (optional). One or more subspecies.
phases	Integer vector. BBCH phases to evaluate.
years	Numeric vector. Years to include.
subregions	Character vector of region/country names.
include_empty	Logical. If TRUE, include subspecies x country rows even when no observations are present.
metric	Character vector. Metrics to compute. Use "all" for: "n_obs", "n_stations", "first_year", "last_year", "median_doy", "completeness".
bbch_names	Optional named list mapping BBCH phase numbers to labels, e.g. list("65" = "Flowering", "87" = "Maturity").
return_results	Logical. If TRUE, returns a list invisibly.

**Details**

Users may specify a **genus**, **species**, or one or more **subspecies**. The report evaluates BBCH phases, regions, years, completeness, and several optional availability metrics.

**Value**

Invisibly returns a list with:

**summary\_table** Tidy long-format summary

**wide\_table** Publication-ready wide-format summary

**heatmap** A ggplot2 object

**Author(s)**

Matthias Templ

**See Also**[pheno\\_trend\\_turning](#) for trend visualization**Examples**

```

pep <- pep_download()

# Use Alpine subset for faster computation
pep_alpine <- pep[country %in% c("Switzerland", "Austria")]

# Grapevine subspecies (wine varieties) - longest historical records
if (nrow(pep_alpine[species == "Vitis vinifera"]) > 0) {
  subspecies_report(
    pep_alpine,
    subspecies_name = c("Vitis vinifera Riesling",
                       "Vitis vinifera MUELLER THURGAU WEISS"),
    subregions = c("Switzerland", "Austria"),
    metric = "all",
    bbch_names = list("65"="Flowering", "81"="Veraison")
  )

  subspecies_report(
    pep_alpine,
    species_name = "Vitis vinifera",
    subregions = c("Switzerland", "Austria"),
    metric = c("n_obs", "median_doy", "completeness")
  )

  subspecies_report(
    pep_alpine,
    genus_name = "Vitis",
    subregions = c("Switzerland", "Austria"),
    metric = "all"
  )
}

```

---

summarize\_subspecies\_availability

*Summarize Subspecies Phenological Data Availability*


---

**Description**

Computes data availability metrics for specified genus, species, or subspecies across BBCH phases and regions. Metrics include numbers of observations, station counts, first/last observation years, median DOY, and completeness ratios.

**Usage**

```
summarize_subspecies_availability(
  pep,
  genus_name = NULL,
  species_name = NULL,
  subspecies_name = NULL,
  phases = c(65, 87),
  years = 1961:2024,
  subregions = c("Austria", "Germany-North", "Germany-South", "Switzerland"),
  include_empty = FALSE,
  metric = c("n_obs", "n_stations", "first_year", "last_year", "median_doy",
             "completeness")
)
```

**Arguments**

pep	A PEP725-style data frame containing at least: genus, species, subspecies, country, year, phase_id, s_id, and day or DOY.
genus_name	Character (optional). Genus to filter.
species_name	Character (optional). Species to filter.
subspecies_name	Character vector (optional). One or more subspecies.
phases	Integer vector. BBCH phases to evaluate.
years	Numeric vector. Years to include.
subregions	Character vector of region/country names.
include_empty	Logical. If TRUE, include subspecies x country rows even when no observations are present.
metric	Character vector specifying which metrics to compute. Allowed values: "n_obs", "n_stations", "first_year", "last_year", "median_doy", "completeness", or "all".

**Value**

A tibble in long format with one row per subspecies x country.

**Author(s)**

Matthias Templ

**Examples**

```
pep <- pep_download()
pep_alpine <- pep[country %in% c("Switzerland", "Austria")]

# Grapevine species (longest historical records)
if (nrow(pep_alpine[species == "Vitis vinifera"]) > 0) {
  summarize_subspecies_availability(
```

```

    pep_alpine,
    species_name = "Vitis vinifera",
    subregions = c("Switzerland", "Austria"),
    metric = "all"
  )

  summarize_subspecies_availability(
    pep_alpine,
    subspecies_name = c("Vitis vinifera Riesling",
                        "Vitis vinifera MUELLER THURGAU WEISS"),
    subregions = c("Switzerland", "Austria"),
    metric = c("n_obs", "median_doy")
  )
}

```

---

summary.pep

*Summary Method for PEP725 Data*


---

## Description

Provides a detailed phenological summary of the dataset, including breakdowns by species, phase, country, and temporal coverage.

## Usage

```

## S3 method for class 'pep'
summary(object, by = c("species", "phase", "country", "year"), top = 10, ...)

```

## Arguments

object	A pep object.
by	Character. Summarize by "species", "phase", "country", or "year". Default is "species".
top	Integer. Number of top entries to show (default 10).
...	Additional arguments (currently unused).

## Value

A pep\_summary object (list) containing summary tables, printed to console.

## Author(s)

Matthias Templ

**Examples**

```
pep <- pep_download()
summary(pep)
summary(pep, by = "phase")
summary(pep, by = "country")
```

---

summary.pep\_completeness

*Summary Method for Completeness Assessment*

---

**Description**

Summary Method for Completeness Assessment

**Usage**

```
## S3 method for class 'pep_completeness'
summary(object, ...)
```

**Arguments**

object	A pep_completeness object
...	Additional arguments (unused)

**Value**

Invisibly returns a summary list

**Author(s)**

Matthias Templ

---

summary.pep\_outliers *Summary Method for Outlier Detection Results*

---

**Description**

Summary Method for Outlier Detection Results

**Usage**

```
## S3 method for class 'pep_outliers'
summary(object, ...)
```

**Arguments**

object            A pep\_outliers object  
...                Additional arguments (unused)

**Value**

Invisibly returns a summary data.table

---

summary.pep\_quality    *Summary Method for Data Quality Assessment*

---

**Description**

Summary Method for Data Quality Assessment

**Usage**

```
## S3 method for class 'pep_quality'  
summary(object, ...)
```

**Arguments**

object            A pep\_quality object  
...                Additional arguments (unused)

**Value**

Invisibly returns a summary list

**Author(s)**

Matthias Templ

---

summary.pheno\_anomaly    *Summary Method for Phenological Anomalies*

---

**Description**

Summary Method for Phenological Anomalies

**Usage**

```
## S3 method for class 'pheno_anomaly'  
summary(object, ...)
```

**Arguments**

object            A pheno\_anomaly object  
...                Additional arguments (unused)

**Value**

Invisibly returns a summary list

**Author(s)**

Matthias Templ

---

summary.pheno\_normals *Summary Method for Phenological Normals*

---

**Description**

Summary Method for Phenological Normals

**Usage**

```
## S3 method for class 'pheno_normals'  
summary(object, ...)
```

**Arguments**

object            A pheno\_normals object  
...                Additional arguments (unused)

**Value**

Invisibly returns a summary list

**Author(s)**

Matthias Templ

---

summary.pheno\_pls      *Summary Method for PLS Phenology Results*

---

**Description**

Summary Method for PLS Phenology Results

**Usage**

```
## S3 method for class 'pheno_pls'
summary(object, vip_threshold = 0.8, ...)
```

**Arguments**

object	A pheno_pls object
vip_threshold	Numeric. VIP threshold for identifying important periods. Default 0.8.
...	Additional arguments (unused)

**Value**

Invisibly returns a summary data.frame

---

summary.pheno\_synchrony  
*Summary Method for Phenological Synchrony Analysis*

---

**Description**

Summary Method for Phenological Synchrony Analysis

**Usage**

```
## S3 method for class 'pheno_synchrony'
summary(object, ...)
```

**Arguments**

object	A pheno_synchrony object
...	Additional arguments (unused)

**Value**

Invisibly returns a summary list

**Author(s)**

Matthias Templ

---

summary.second\_events *Summary Method for Second Events Detection*

---

**Description**

Summary Method for Second Events Detection

**Usage**

```
## S3 method for class 'second_events'  
summary(object, ...)
```

**Arguments**

object	A second_events object
...	Additional arguments (unused)

**Value**

The summary data.table (invisibly)

---

[.pep *Subset PEP725 Data While Preserving Class*

---

**Description**

Ensures that subsetting operations return a pep object when the result still has the required structure.

**Usage**

```
## S3 method for class 'pep'  
x[...]
```

**Arguments**

x	A pep object.
...	Arguments passed to the data.table subset method.

**Value**

A pep object if structure is preserved, otherwise a data.table.

**Author(s)**

Matthias Templ

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