

Package ‘photosynthesis’

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Title Tools for Plant Ecophysiology & Modeling

Depends R (>= 4.0.0), ggplot2 (>= 3.3.0), minpack.lm (>= 1.2-1), units (>= 0.6.6)

Imports checkmate (>= 2.0.0), crayon (>= 1.3.4), dplyr (>= 0.8.5), furrr (>= 0.1.0), future (>= 1.17.0), glue (>= 1.4.0), graphics (>= 4.0.0), grDevices (>= 4.0.0), gunit (>= 1.0.0), magrittr (>= 1.5.0), methods (>= 3.5.0), nlme (>= 3.1-147), pkgnet (>= 0.4.1), purrr (>= 0.3.3), rlang (>= 0.4.6), stats (>= 4.0.0), stringr (>= 1.4.0), tealeaves (>= 1.0.5), tidyselect (>= 1.0.0), utils (>= 4.0.0)

Suggests testthat, knitr, rmarkdown, tidyverse

Description Contains modeling and analytical tools for plant ecophysiology.

MODELING: Simulate C3 photosynthesis using the Farquhar, von Caemmerer, Berry (1980) [doi:10.1007/BF00386231](https://doi.org/10.1007/BF00386231) model as described in Buckley and Diaz-Espejo (2015) [doi:10.1111/pce.12459](https://doi.org/10.1111/pce.12459). It uses units to ensure that parameters are properly specified and transformed before calculations.

Temperature response functions get automatically ``baked'' into all parameters based on leaf temperature following Bernacchi et al. (2002) [doi:10.1104/pp.008250](https://doi.org/10.1104/pp.008250). The package includes boundary layer, cuticular, stomatal, and mesophyll conductances to CO₂, which each can vary on the upper and lower portions of the leaf. Use straightforward functions to simulate photosynthesis over environmental gradients such as Photosynthetic Photon Flux Density (PPFD) and leaf temperature, or over trait gradients such as CO₂ conductance or photochemistry.

ANALYTICAL TOOLS: Fit ACi (Farquhar et al. 1980 [doi:10.1007/BF00386231](https://doi.org/10.1007/BF00386231)) and AQ curves (Marshall & Biscoe 1980 [doi:10.1093/jxb/31.1.29](https://doi.org/10.1093/jxb/31.1.29)), temperature responses (Heskel et al. 2016 [doi:10.1073/pnas.1520282113](https://doi.org/10.1073/pnas.1520282113)); Kruse et al. 2008 [doi:10.1111/j.1365-3040.2008.01809.x](https://doi.org/10.1111/j.1365-3040.2008.01809.x), Medlyn et al. 2002 [doi:10.1046/j.1365-3040.2002.00891.x](https://doi.org/10.1046/j.1365-3040.2002.00891.x), Hobbs et al. 2013 [doi:10.1021/cb4005029](https://doi.org/10.1021/cb4005029)), respiration in the light (Kok 1956 [doi:10.1016/0006-3002\(56\)90003-8](https://doi.org/10.1016/0006-3002(56)90003-8)), Walker & Ort 2015 [doi:10.1111/pce.12562](https://doi.org/10.1111/pce.12562), Yin et al. 2009 [doi:10.1111/j.1365-3040.2009.01934.x](https://doi.org/10.1111/j.1365-3040.2009.01934.x), Yin et al. 2011

<[doi:10.1093/jxb/err038](https://doi.org/10.1093/jxb/err038)>), mesophyll conductance (Harley et al. 1992
 <[doi:10.1104/pp.98.4.1429](https://doi.org/10.1104/pp.98.4.1429)>), pressure-volume curves (Koide et al. 2000
 <[doi:10.1007/978-94-009-2221-1_9](https://doi.org/10.1007/978-94-009-2221-1_9)>, Sack et al. 2003
 <[doi:10.1046/j.0016-8025.2003.01058.x](https://doi.org/10.1046/j.0016-8025.2003.01058.x)>, Tyree et al. 1972
 <[doi:10.1093/jxb/23.1.267](https://doi.org/10.1093/jxb/23.1.267)>), hydraulic vulnerability curves (Ogle et al. 2009
 <[doi:10.1111/j.1469-8137.2008.02760.x](https://doi.org/10.1111/j.1469-8137.2008.02760.x)>, Pammeter et al. 1998
 <[doi:10.1093/treephys/18.8-9.589](https://doi.org/10.1093/treephys/18.8-9.589)>), and tools for running sensitivity
 analyses particularly for variables with uncertainty (e.g. g_mc, gamma_star,
 R_d).

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Encoding UTF-8

RoxygenNote 7.1.1

VignetteBuilder knitr

URL <https://github.com/cdmuir/photosynthesis>

BugReports <https://github.com/cdmuir/photosynthesis/issues>

NeedsCompilation no

Author Joseph Stinziano [aut] (<<https://orcid.org/0000-0002-7628-4201>>),
 Cassaundra Roback [aut],
 Demi Sargent [aut],
 Bridget Murphy [aut],
 Patrick Hudson [aut, dtc],
 Chris Muir [aut, cre] (<<https://orcid.org/0000-0003-2555-3878>>)

Maintainer Chris Muir <cdmuir@hawaii.edu>

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R topics documented:

photosynthesis-package	3
analyze_sensitivity	3
aq_response	6
A_supply	6
bake	8
baked-class	9
bake_par	10
calculate_jmax	10
check_dependencies	11
CO2_conductance	11
compile_data	13
compute_sensitivity	14
constants	16
enviro_par	17
fit_aci_response	17
fit_aq_response	21

fit_gs_model	23
fit_g_mc_variableJ	26
fit_hydra_vuln_curve	28
fit_many	31
fit_PV_curve	32
fit_r_light_kok	35
fit_t_response	38
FvCB	42
gs_mod_ballberry	44
gw2gc	45
J	46
leaf_par	47
make_parameters	47
parameter_names	50
photosynthesis	51
ppm2pa	54
print_graphs	55
read_li6800	57
t_response_arrhenius	57

Index**60**

photosynthesis-package
photosynthesis package

Description

Modeling and analytical tools for plant ecophysiology

Details

See the README on [GitHub](#)

analyze_sensitivity *Running 2-parameter sensitivity analyses*

Description

Running 2-parameter sensitivity analyses

Usage

```
analyze_sensitivity(
  data,
  funct,
  test1 = NA,
  values1,
  test2 = NA,
  values2,
  element_out = 1,
  ...
)
```

Arguments

<code>data</code>	Dataframe
<code>funct</code>	Function to use - do not use parentheses
<code>test1</code>	Input parameter to vary and test
<code>values1</code>	Values of test1 to use
<code>test2</code>	Input parameter to vary and test
<code>values2</code>	Values of test2 to use
<code>element_out</code>	List element to compile
<code>...</code>	Additional arguments required for the function

Value

`analyze_sensitivity` runs a 2-parameter sensitivity analysis. Note that any parameter value combinations that break the input function WILL break this function. For 1-parameter sensitivity analysis, use `test1` only.

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Define a grouping factor based on light intensity to split the ACi
# curves
data$Q_2 <- as.factor((round(data$Qin, digits = 0)))

# Convert leaf temperature to K
data$T_leaf <- data$Tleaf + 273.15

# Run a sensitivity analysis on gamma_star and mesophyll conductance
```

```

# at 25 Celsius for one individual curve
pars <- analyze_sensitivity(
  data = data[data$Q_2 == 1500, ],
  funct = fit_aci_response,
  varnames = list(
    A_net = "A",
    T_leaf = "T_leaf",
    C_i = "Ci",
    PPFD = "Qin"
  ),
  useg_mct = TRUE,
  test1 = "gamma_star25",
  element_out = 1,
  test2 = "g_mc25",
  fitTPU = TRUE,
  Ea_gamma_star = 0,
  Ea_g_mc = 0,
  values1 = seq(
    from = 20,
    to = 40,
    by = 2
  ),
  values2 = seq(
    from = 0.5,
    to = 2,
    by = 0.1
  )
)

# Graph V_cmax
ggplot(pars, aes(x = gamma_star25, y = g_mc25, z = V_cmax)) +
  geom_tile(aes(fill = V_cmax)) +
  labs(
    x = expression(Gamma * "*"[25] ~ "(" * mu * mol ~ mol^
      {
        -1
      } * ")"),
    y = expression(g[m][25] ~ "(" * mu * mol ~ m^{
      -2
    } ~ s^{
      -1
    } ~ Pa^
      {
        -1
      } * ")")
  ) +
  scale_fill_distiller(palette = "Greys") +
  geom_contour(colour = "Black", size = 1) +
  theme_bw()

```

aq_response*Non-rectangular hyperbolic model of light responses*

Description

Non-rectangular hyperbolic model of light responses

Usage

```
aq_response(k_sat, phi_J, Q_abs, theta_J)
```

Arguments

k_sat	Light saturated rate of process k
phi_J	Quantum efficiency of process k
Q_abs	Absorbed light intensity (umol m ⁻² s ⁻¹)
theta_J	Curvature of the light response

Value

aq_response is used to describe the response of a process to absorbed light intensity. Assumes that input is absorbed light. Note that if absorbed light is not used, then the meaning of phi_J becomes unclear. This function is designed to be used with fit_aq_response, however it could easily be fed into a different fitting approach (e.g. Bayesian approaches). Originally from Marshall et al. 1980.

References

Marshall B, Biscoe P. 1980. A model for C3 leaves describing the dependence of net photosynthesis on irradiance. J Ex Bot 31:29-39

A_supply*CO₂ supply and demand function (mol / m² s)*

Description

This function is not intended to be called by users directly.

Usage

```
A_supply(C_chl, pars, unitless = FALSE)  
A_demand(C_chl, pars, unitless = FALSE)
```

Arguments

C_chl	Chloroplastic CO2 concentration in Pa of class units
pars	Concatenated parameters (leaf_par, enviro_par, and constants)
unitless	Logical. Should units be set? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.

Details**Supply function:**

$$A = g_{tc}(C_{air} - C_{chl})$$

Demand function:

$$A = (1 - \Gamma * /C_{chl})\min(W_{carbox}, W_{regen}, W_{tpu}) - R_d$$

Symbol	R	Description	Units	Default
A	A	photosynthetic rate	$\mu\text{mol CO}_2 / (\text{m}^2 \text{s})$	calculated
g_tc	g_tc	total conductance to CO2	$\mu\text{mol CO}_2 / (\text{m}^2 \text{s Pa})$	calculated
C_air	C_air	atmospheric CO2 concentration	Pa	41
C_chl	C_chl	chloroplastic CO2 concentration	Pa	calculated
R_d	R_d	nonphotorespiratory CO2 release	$\mu\text{mol CO}_2 / (\text{m}^2 \text{s})$	2
$\Gamma*$	gamma_star	chloroplastic CO2 compensation point	Pa	3.743

Value

Value in mol / (m² s) of class units

Examples

```
bake_par <- make_bakepar()
constants <- make_constants(use_tealeaves = FALSE)
enviro_par <- make_enviropar(use_tealeaves = FALSE)
leaf_par <- make_leafpar(use_tealeaves = FALSE)
leaf_par <- bake(leaf_par, bake_par, constants)
# Or bake with piping (need library(magrittr))
# leaf_par %>% bake(bake_par, constants)
enviro_par$T_air <- leaf_par$T_leaf

pars <- c(leaf_par, enviro_par, constants)
C_chl <- set_units(35, "Pa")

A_supply(C_chl, pars)

A_demand(C_chl, pars)
```

bake

Leaf parameter temperature responses

Description

'bake' leaf parameters using temperature response functions

Constructor function for baked class. This will also inherit class [leaf_par](#). This function ensures that temperature is "baked in" to leaf parameter calculations T_leaf using temperature response functions detailed below.

Usage

```
bake(leaf_par, bake_par, constants, assert_units = TRUE)

temp_resp1(par25, E_a, R, T_leaf, T_ref, unitless)

temp_resp2(par25, D_s, E_a, E_d, R, T_leaf, T_ref, unitless)
```

Arguments

<code>leaf_par</code>	A list of leaf parameters inheriting class leaf_par . This can be generated using the make_leafpar function.
<code>bake_par</code>	A list of temperature response parameters inheriting class bake_par . This can be generated using the make_bakepar function.
<code>constants</code>	A list of physical constants inheriting class constants . This can be generated using the make_constants function.
<code>assert_units</code>	Logical. Should parameter units be checked? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.
<code>par25</code>	Parameter value at 25 °C of class units.
<code>E_a</code>	Empirical temperature response value in J/mol of class units.
<code>R</code>	Ideal gas constant in J / (mol K) of class units. See make_constants .
<code>T_leaf</code>	Leaf temperature in K of class units. Will be converted to °C.
<code>T_ref</code>	Reference temperature in K of class units.
<code>unitless</code>	Logical. Should units be set? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.
<code>D_s</code>	Empirical temperature response value in J / (mol K) of class units.
<code>E_d</code>	Empirical temperature response value in J/mol of class units.

Details

Several leaf parameters ([leaf_par](#)) are temperature sensitive. Temperature-sensitive parameters are input at a reference temperature of 25 °C. These parameters are provided as par_name25 and then "baked" using the appropriate temperature response function and parameters in [bake_par](#). The "baked" parameter will have the name without "25" appended (par_name). E.g. V_cmax25 becomes V_cmax.

Temperature response functions following Buckley and Diaz-Espejo (2015)

Temperature response function 1 ([temp_response1](#)):

$$\text{par}(T_{\text{leaf}}) = \text{par25} \exp(E_a/(RT_{\text{ref}})(T_{\text{leaf}} - 25)/(T_{\text{leaf}} + 273.15))$$

T_{ref} is the reference temperature in K

T_{leaf} is the leaf temperature in °C

Temperature response function 2 ([temp_response2](#)) is the above equation multiplied by:

$$(1 + \exp((D_s/R - E_d/(RT_{\text{ref}}))))/(1 + \exp((D_s/R) - (E_d/(R(T_{\text{leaf}} + 273.15)))))$$

Function 1 increases exponentially with temperature; Function 2 peaks a particular temperature.

References

Buckley TN, Diaz-Espejo A. 2015. Partitioning changes in photosynthetic rate into contributions from different variables. Plant, Cell and Environment 38: 1200-1211.

Examples

```
bake_par <- make_bakepar()
constants <- make_constants(use_tealeaves = FALSE)
leaf_par <- make_leafpar(
  replace = list(T_leaf = set_units(293.15, "K")),
  use_tealeaves = FALSE
)
baked_leafpar <- bake(leaf_par, bake_par, constants)

baked_leafpar$V_cmax25
baked_leafpar$V_cmax
```

Description

See [bake](#)

bake_par*S3 class bake_par***Description**

Constructor function for **bake_par** class. This function ensures that leaf temperature gets properly "baked" into leaf parameters.

Usage

```
bake_par(.x)
```

Arguments

.x	A list to be constructed into bake_par .
----	---

calculate_jmax*Inverse non-rectangular hyperbola for J_max calculation***Description**

Inverse non-rectangular hyperbola for J_max calculation

Usage

```
calculate_jmax(PPFD, alpha, J, theta_J)
```

```
calculate_j(PPFD, alpha, J_max, theta_J)
```

Arguments

PPFD	light intensity in umol m-2 s-1
alpha	initial slope of the light response
J	electron transport rate in umol m-2 s-1
theta_J	curvature of the light response
J_max	maximum rate of electron transport in umol m-2 s-1

Value

`calculate_jmax` calculates J_max given PPFD and J. It is necessary for the electron transport component of the `fit_aci_response` function.

`calculate_j` provides a model of the light response of J. It is necessary for fitting the electron transport component of the photosynthetic CO₂ response curves in `fit_aci_response`.

check_dependencies	<i>Checking dependencies of photosynthesis</i>
--------------------	--

Description

Checking dependencies of photosynthesis

Usage

```
check_dependencies()
```

Value

check_dependencies returns an html report that shows dependencies between functions and other packages. Does not take any arguments to run. It is useful for determining which functions need to be changed together.

Examples

```
check_dependencies()
```

CO2_conductance	<i>Conductance to CO₂ (umol / (m² s Pa))</i>
-----------------	--

Description

Conductance to CO₂ (umol / (m² s Pa))

- g_tc: total conductance to CO₂
- g_uc: cuticular conductance to CO₂
- g_bc: boundary layer conductance to CO₂
- g_mc: mesophyll conductance to CO₂
- g_sc: stomatal conductance to CO₂

Usage

```
.get_gtc(pars, unitless)  
  
.get_guc(pars, surface, unitless)  
  
.get_gbc(pars, surface, unitless)  
  
.get_gmc(pars, surface, unitless)  
  
.get_gsc(pars, surface, unitless)
```

Arguments

pars	Concatenated parameters (leaf_par, enviro_par, and constants)
unitless	Logical. Should units be set? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.
surface	Leaf surface (lower or upper)

Details

Total conductance to CO₂ is the sum of parallel conductances on the lower ($g_{c,\text{lower}}$) and upper ($g_{c,\text{upper}}$) leaf portions:

$$g_{c,\text{total}} = g_{c,\text{lower}} + g_{c,\text{upper}}$$

Each partial conductance consists of two parallel conductances, the cuticular conductance ($g_{u,c}$) and the in-series conductances through mesophyll ($g_{m,c}$), stomata ($g_{s,c}$), and boundary layer ($g_{b,c}$). To simplify the formula, I use substitute resistance where $r_x = 1/g_x$. For surface i :

$$g_{c,i} = g_{u,i} + (1/(r_{m,i} + r_{s,i} + r_{b,i}))$$

The cuticular, stomatal, and mesophyll conductances can be the same or different for upper and lower. The partitioning factors (k_x) divide the conductance between surfaces while keeping the total conductance constant:

$$g_{x,\text{lower}} = g_x(1/(1 + k_x))$$

$$g_{x,\text{upper}} = g_x(k_x/(1 + k_x))$$

$$g_x = g_{x,\text{lower}} + g_{x,\text{upper}}$$

How the partitioning factors work:

k_x	description
0	all conductance on lower surface/portion
0.5	2/3 conductance on lower surface
1	conductance evenly divided between surfaces/portions
2	2/3 conductance on upper surface
Inf	all conductance on upper surface/portion

The boundary layer conductances for each are calculated on the basis of mass and heat transfer (see [.get_gbc](#)).

Symbol	R	Description	Units	Default
g_{mc}	g_{mc}	mesophyll conductance to CO ₂ (T_leaf)	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s Pa})$	calculated
g_{sc}	g_{sc}	stomatal conductance to CO ₂	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s Pa})$	4
g_{uc}	g_{uc}	cuticular conductance to CO ₂	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s Pa})$	0.1
k_{mc}	k_{mc}	partition of g_{mc} to lower mesophyll	none	1

k_{sc}	k_{sc}	partition of g_{sc} to lower surface	none	1
k_{uc}	k_{uc}	partition of g_{uc} to lower surface	none	1

compile_data	<i>Compiling outputs from lists</i>
--------------	-------------------------------------

Description

Compiling outputs from lists

Usage

```
compile_data(data, output_type = "list", list_element)
```

Arguments

data	List of elements
output_type	Type of desired output. For graphs or models, use "list", for parameters, use "dataframe".
list_element	Which elements of the sublists do you wish to compile?

Value

compile_data converts the outputs of fit_many into a form more readily usable for analysis. Can be used to create dataframe of all fitted parameters, a list of model outputs, a list of graphs for plotting. This function is NOT restricted to compiling outputs from plantecophys tools but could be used to compile elements from ANY list of lists.

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Define a grouping factor based on light intensity to split the ACi
# curves
data$Q_2 <- as.factor((round(data$Qin, digits = 0)))

# Convert leaf temperature to K
data$T_leaf <- data$Tleaf + 273.15

# Fit many curves
```

```

fits <- fit_many(
  data = data,
  varnames = list(
    A_net = "A",
    T_leaf = "T_leaf",
    C_i = "Ci",
    PPFD = "Qin"
  ),
  funct = fit_aci_response,
  group = "Q_2"
)

# Compile graphs into a list for plotting
fits_graphs <- compile_data(fits,
  list_element = 2
)

# Plot one graph from the compiled list
plot(fits_graphs[[1]])

```

compute_sensitivity *Computing measures of sensitivity*

Description

Computing measures of sensitivity

Usage

```

compute_sensitivity(
  data,
  varnames = list(Par = "Par", test1 = "test1", test2 = "test2"),
  test1_ref,
  test2_ref
)

```

Arguments

data	Dataframe with output from sensitivity_analysis()
varnames	Variable names
test1_ref	Reference value for parameter
test2_ref	Reference value for parameter

Value

compute_sensitivity calculates two sets of sensitivity measures: parameter effect (Bauerle et al., 2014), and control coefficient (Capaldo & Pandis, 1997). This function is useful in determining how much a given input (assumed or otherwise) can affect the model output and conclusions. Particularly useful if a given parameter is unknown during a fitting or modeling process.

References

- Bauerle WL, Daniels AB, Barnard DM. 2014. Carbon and water flux responses to physiology by environment interactions: a sensitivity analysis of variation in climate on photosynthetic and stomatal parameters. Climate Dynamics 42: 2539-2554.
- Capaldo KP, Pandis SN 1997. Dimethylsulfide chemistry in the remote marine atmosphere: evaluation and sensitivity analysis of available mechanisms. J Geophys Res 102:23251-23267

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Define a grouping factor based on light intensity to split the ACi
# curves
data$Q_2 <- as.factor((round(data$Qin, digits = 0)))

# Convert leaf temperature to K
data$T_leaf <- data$Tleaf + 273.15

# Run a sensitivity analysis on gamma_star and mesophyll conductance
# at 25 Celsius for one individual curve
pars <- analyze_sensitivity(
  data = data[data$Q_2 == 1500, ],
  funct = fit_aci_response,
  varnames = list(
    A_net = "A",
    T_leaf = "T_leaf",
    C_i = "Ci",
    PPFD = "Qin"
  ),
  useg_mct = TRUE,
  test1 = "gamma_star25",
  element_out = 1,
  test2 = "g_mc25",
  fitTPU = TRUE,
  Ea_gamma_star = 0,
  Ea_g_mc = 0,
  values1 = seq(
    from = 20,
    to = 60,
    by = 2
  ),
  values2 = seq(
    from = 0.2,
    to = 2,
    by = 0.1
)
)
```

```

        by = 0.1
    )
)
# Compute measures of sensitivity
par2 <- compute_sensitivity(
    data = pars,
    varnames = list(
        Par = "V_cmax",
        test1 = "gamma_star25",
        test2 = "g_mc25"
    ),
    test1_ref = 42,
    test2_ref = 1
)
# Plot control coefficients
ggplot(par2, aes(y = CE_gamma_star25, x = CE_g_mc25, colour = V_cmax)) +
    geom_point() +
    theme_bw()
# Note that in this case a missing point appears due to an infinity

```

constants*S3 class constants***Description**

Constructor function for **constants** class. This function ensures that physical constant inputs are properly formatted.

Usage

```
constants(.x, use_tealeaves)
```

Arguments

- .x A list to be constructed into **constants**.
- use_tealeaves Logical. Should leaf energy balance be used to calculate leaf temperature (T_{leaf})? If TRUE, [tleaf](#) calculates T_{leaf} . If FALSE, user-defined T_{leaf} is used. Additional parameters and constants are required, see [make_parameters](#).

enviro_par*S3 class enviro_par*

Description

Constructor function for enviro_par class. This function ensures that environmental parameter inputs are properly formatted.

Usage

```
enviro_par(.x, use_tealeaves)
```

Arguments

- .x A list to be constructed into **enviro_par**.
 - use_tealeaves Logical. Should leaf energy balance be used to calculate leaf temperature (T_leaf)? If TRUE, `tleaf` calculates T_leaf. If FALSE, user-defined T_leaf is used. Additional parameters and constants are required, see [make_parameters](#).
-

fit_aci_response

Fitting ACi curves

Description

Fitting ACi curves

Usage

```
fit_aci_response(
  data,
  varnames = list(A_net = "A_net", T_leaf = "T_leaf", C_i = "C_i", PPFD = "PPFD", g_mc
    = "g_mc"),
  P = 100,
  fitTPU = TRUE,
  alpha_g = 0,
  R_d_meas = NULL,
  useR_d = FALSE,
  useg_mc = FALSE,
  useg_mct = FALSE,
  usegamma_star = FALSE,
  useK_M = FALSE,
  useK_C_K_0 = FALSE,
  alpha = 0.24,
  theta_J = 0.85,
  gamma_star25 = 42.75,
```

```

Ea_gamma_star = 37830,
K_M25 = 718.4,
Ea_K_M = 65508.28,
g_mc25 = 0.08701,
Ea_g_mc = 0,
K_C25 = NULL,
Ea_K_C = NULL,
K_O25 = NULL,
Ea_K_O = NULL,
Oconc = 21,
gamma_star_set = NULL,
K_M_set = NULL,
...
)

```

Arguments

data	Dataframe for A-Ci curve fitting
varnames	List of variable names. varnames = list(A_net = "A_net", T_leaf = "T_leaf", C_i = "C_i", PPFD = "PPFD", g_mc = "g_mc"), where A_net is net CO2 assimilation, T_leaf is leaf temperature in Celsius, C_i is intercellular CO2 concentration in umol/mol, PPFD is incident irradiance in umol m-2 s-1 (note that it is ASSUMED to be absorbed irradiance, so be sure to adjust according to light absorbance and PSI/PSII partitioning accordingly OR interpret the resultant values of J and J_max with caution), g_mc is mesophyll conductance to CO2 in mol m-2 s-1 Pa-1.
P	Atmospheric pressure in kPa
fitTPU	Should triose phosphate utilization (V_TPU) be fit?
alpha_g	Fraction of respiratory glycolate carbon that is not returned to the chloroplast (von Caemmerer, 2000). If ACi curves show high-CO2 decline, then this value should be > 0.
R_d_meas	Measured value of respiratory CO2 efflux in umol m-2 s-1. Input value should be positive to work as expected with the equations.
useR_d	Use a measured value of R_d? Set to TRUE if using R_d_meas.
useg_mc	Use mesophyll conductance? Set to TRUE if specifying g_mc in varnames above.
useg_mct	Use mesophyll conductance temperature response? Set to TRUE if using a temperature response of mesophyll conductance.
usegamma_star	Specify gamma_star value? If FALSE, uses a temperature response function with Nicotiana tabacum defaults from Bernacchi et al. 2001.
useK_M	Specify K_M? If FALSE, uses an Arrhenius temperature response function with Nicotiana tabacum defaults from Bernacchi et al. 2001.
useK_C_K_O	Use individual carboxylation/oxygenation constants for rubisco? If TRUE, need to specify values at 25C and activation energy for the Arrhenius temperature response function.

alpha	Quantum yield of CO2 assimilation
theta_J	Curvature of the photosynthetic light response curve
gamma_star25	gamma_star at 25C in umol mol-1
Ea_gamma_star	Activation energy of gamma_star in J mol-1
K_M25	Michaelis-Menten constant for rubisco at 25C
Ea_K_M	Activation energy for K_M in J mol-1
g_mc25	Mesophyll conductance at 25C in mol m-2 s-1
Ea_g_mc	Activation energy of g_mc in J mol-1
K_C25	Michaelis-Menten constant for rubisco carboxylation at 25C
Ea_K_C	Activation energy for K_C in J mol-1
K_025	Michaelis-Menten constant for rubisco oxygenation at 25C
Ea_K_O	Activation energy for K_O in J mol-2
Oconc	O2 concentration in intracellular O2 when using K_C_K_O
gamma_star_set	Value of gamma_star to use (in ppm) if usegamma_star = TRUE
K_M_set	Value of K_M to use if useK_M = TRUE
...	Other arguments to pass on

Value

fit_aci_response fits ACi curves using an approach similar to Gu et al. 2010. Iterates all possible C_i transition points and checks for inadmissible curve fits. If no curves are admissible (either due to poor data or poor assumed parameters), the output will include a dataframe of NA values. Default parameters are all from Bernacchi et al. 2001, 2002.

References

- Bernacchi CJ, Singsaas EL, Pimentel C, Portis AR, Long SP. 2001. Improved temperature response functions for models of rubisco-limited photosynthesis. *Plant Cell Environment* 24:253-259.
- Bernacchi CJ, Portis AR, Nakano H, von Caemmerer S, Long SP. 2002. Temperature response of mesophyll conductance. Implications for the determination of rubisco enzyme kinetics and for limitations to photosynthesis in vivo. *Plant Physiology* 130:1992-1998.
- Gu L, Pallardy SG, Tu K, Law BE, Wullschleger SD. 2010. Reliable estimation of biochemical parameters from C3 leaf photosynthesis-intercellular carbon dioxide response curves. *Plant Cell Environment* 33:1852-1874.
- von Caemmerer S. 2000. Biochemical models of leaf photosynthesis. CSIRO Publishing, Collingwood.

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
```

```

data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
                           package = "photosynthesis"
                           )))

# Define a grouping factor based on light intensity to split the ACi
# curves
data$Q_2 <- as.factor((round(data$Qin, digits = 0)))

# Convert leaf temperature to K
data$T_leaf <- data$Tleaf + 273.15

# Fit ACi curve. Note that we are subsetting the dataframe
# here to fit for a single value of Q_2
fit <- fit_aci_response(data[data$Q_2 == 1500, ],
                        varnames = list(
                          A_net = "A",
                          T_leaf = "T_leaf",
                          C_i = "Ci",
                          PPFD = "Qin"
                        )
)
# View fitted parameters
fit[[1]]

# View graph
fit[[2]]

# View data with modelled parameters attached
fit[[3]]

# Fit many curves
fits <- fit_many(
  data = data,
  varnames = list(
    A_net = "A",
    T_leaf = "T_leaf",
    C_i = "Ci",
    PPFD = "Qin"
  ),
  funct = fit_aci_response,
  group = "Q_2"
)
# Print the parameters
# First set of double parentheses selects an individual group value
# Second set selects an element of the sublist
fits[[3]][[1]]

# Print the graph
fits[[3]][[2]]

# Compile graphs into a list for plotting

```

```

fits_graphs <- compile_data(fits,
  list_element = 2
)

# Compile parameters into dataframe for analysis
fits_pars <- compile_data(fits,
  output_type = "dataframe",
  list_element = 1
)

```

fit_aq_response *Fitting light responses of net CO₂ assimilation*

Description

Fitting light responses of net CO₂ assimilation

Usage

```

fit_aq_response(
  data,
  varnames = list(A_net = "A_net", PPFD = "PPFD"),
  usealpha_Q = FALSE,
  alpha_Q = 0.84,
  title = NULL
)

```

Arguments

data	Dataframe containing CO ₂ assimilation light response
varnames	Variable names where varnames = list(A_net = "A_net", PPFD = "PPFD"). A_net is net CO ₂ assimilation in umol m ⁻² s ⁻¹ , PPFD is incident irradiance. PPFD can be corrected for light absorbance by using useapha_Q and setting alpha_Q.
usealpha_Q	Correct light intensity for absorbance? Default is FALSE.
alpha_Q	Absorbance of incident light. Default value is 0.84.
title	Title for graph

Value

fit_aq_response fits the light response of net CO₂ assimilation. Output is a dataframe containing light saturated net CO₂ assimilation, quantum yield of CO₂ assimilation (ϕ_J), curvature of the light response (θ_J), respiration (R_d), light compensation point (LCP), and residual sum of squares (resid_SS). Note that R_d fitted in this way is essentially the same as the Kok method, and represents a respiration value in the light that may not be accurate. R_d output should thus be interpreted more as a residual parameter to ensure an accurate fit of the light response parameters. Model originally from Marshall et al. 1980.

References

Marshall B, Biscoe P. 1980. A model for C3 leaves describing the dependence of net photosynthesis on irradiance. J Ex Bot 31:29-39

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Fit many AQ curves
# Set your grouping variable
# Here we are grouping by CO2_s and individual
data$C_s <- (round(data$CO2_s, digits = 0))

# For this example we need to round sequentially due to CO2_s setpoints
data$C_s <- as.factor(round(data$C_s, digits = -1))

# To fit one AQ curve
fit <- fit_aq_response(data[data$C_s == 600, ],
  varnames = list(
    A_net = "A",
    PPFD = "Qin"
  )
)

# Print model summary
summary(fit[[1]])

# Print fitted parameters
fit[[2]]

# Print graph
fit[[3]]

# Fit many curves
fits <- fit_many(
  data = data,
  varnames = list(
    A_net = "A",
    PPFD = "Qin",
    group = "C_s"
  ),
  funct = fit_aq_response,
  group = "C_s"
)
```

```

# Look at model summary for a given fit
# First set of double parentheses selects an individual group value
# Second set selects an element of the sublist
summary(fits[[3]][[1]])

# Print the parameters
fits[[3]][[2]]

# Print the graph
fits[[3]][[3]]

# Compile graphs into a list for plotting
fits_graphs <- compile_data(fits,
  list_element = 3
)

# Compile parameters into dataframe for analysis
fits_pars <- compile_data(fits,
  output_type = "dataframe",
  list_element = 2
)

```

fit_gs_model*Fitting stomatal conductance models***Description**

Fitting stomatal conductance models

Usage

```

fit_gs_model(
  data,
  varnames = list(A_net = "A_net", C_air = "C_air", g_sw = "g_sw", RH = "RH", VPD =
    "VPD"),
  model = c("BallBerry", "Leuning", "Medlyn_partial", "Medlyn_full"),
  D0 = 3,
  ...
)

```

Arguments

data Dataframe

varnames Variable names

For the Ball-Berry model: varnames = list(A_net = "A_net", C_air = "C_air", g_sw = "g_sw", RH = "RH") where A_net is net CO₂ assimilation, C_air is CO₂ concentration at the leaf surface in umol mol⁻¹, g_sw is stomatal conductance to H₂O, and RH is relative humidity as a proportion.

For the Leuning model: varnames = list(A_net = "A_net", C_air = "C_air", g_sw = "g_sw", VPD = "VPD") where A_net is net CO₂ assimilation, C_air is CO₂ concentration at the leaf surface in umol mol⁻¹, g_sw is stomatal conductance to H₂O, and VPD is leaf to air vapor pressure deficit in kPa.

For the Medlyn et al. 2011 models: varnames = list(A_net = "A_net", C_air = "C_air", g_sw = "g_sw", VPD = "VPD") where A_net is net CO₂ assimilation, C_air is CO₂ concentration at the leaf surface in umol mol⁻¹, g_sw is stomatal conductance to H₂O, and VPD is leaf to air vapor pressure deficit in kPa.

model	Which model(s) to fit? Defaults to all models. Available options are "Ball-Berry", "Leuning", "Medlyn_partial", and "Medlyn_full", from Ball et al. (1987), Leuning (1995), and Medlyn et al. (2011).
D0	Vapor pressure sensitivity of stomata (Leuning 1995)
...	Arguments to pass on to the nlsLM() function for the Medlyn models.

Value

fit_gs_model fits one or more stomatal conductance models to the data. The top level of the output list is named after the fitted model, while the second level contains the Model, Parameters, and Graph, in that order.

References

- Ball JT, Woodrow IE, Berry JA. 1987. A model predicting stomatal conductance and its contribution to the control of photosynthesis under different environmental conditions, in Progress in Photosynthesis Research, Proceedings of the VII International Congress on Photosynthesis, vol. 4, edited by I. Biggins, pp. 221–224, Martinus Nijhoff, Dordrecht, Netherlands.
- Leuning R. 1995. A critical appraisal of a coupled stomatal- photosynthesis model for C3 plants. Plant Cell Environ 18:339-357
- Medlyn BE, Duursma RA, Eamus D, Ellsworth DS, Prentice IC, Barton CVM, Crous KY, Angelis PD, Freeman M, Wingate L. 2011. Reconciling the optimal and empirical approaches to modeling stomatal conductance. Glob Chang Biol 17:2134-2144

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Convert RH to a proportion
data$RH <- data$RHcham / 100

# Fit stomatal conductance models
# Can specify a single model, or all as below
fits <- fit_gs_model(
```

```

data = data,
varnames = list(
  A_net = "A",
  C_air = "Ca",
  g_sw = "gsw",
  RH = "RH",
  VPD = "VPDleaf"
),
model = c(
  "BallBerry",
  "Leuning",
  "Medlyn_partial",
  "Medlyn_full"
),
D0 = 3
)

# Look at BallBerry model summary:
summary(fits[["BallBerry"]][["Model"]])

# Look at BallBerry parameters
fits[["BallBerry"]][["Parameters"]]

# Look at BallBerry plot
fits[["BallBerry"]][["Graph"]]

# Fit many g_sw models
# Set your grouping variable
# Here we are grouping by Qin and individual
data$Q_2 <- as.factor((round(data$Qin, digits = 0)))

fits <- fit_many(data,
  varnames = list(
    A_net = "A",
    C_air = "Ca",
    g_sw = "gsw",
    RH = "RH",
    VPD = "VPDleaf"
),
  funct = fit_gs_model,
  group = "Q_2"
)

# Look at the Medlyn_partial outputs at 750 PAR
# Model summary
summary(fits[["750"]][["Medlyn_partial"]][["Model"]])

# Model parameters
fits[["750"]][["Medlyn_partial"]][["Parameters"]]

# Graph
fits[["750"]][["Medlyn_partial"]][["Graph"]]

```

```

# Compile parameter outputs for BallBerry model
# Note that it's the first element for each PAR value
# First compile list of BallBerry fits
bbmods <- compile_data(
  data = fits,
  output_type = "list",
  list_element = 1
)
# Now compile the parameters (2nd element) into a dataframe
bbpars <- compile_data(
  data = bbmods,
  output_type = "dataframe",
  list_element = 2
)

# Convert group variable back to numeric
bbpars$ID <- as.numeric(bbpars$ID)

# Take quick look at light response of intercept parameters
plot(g0 ~ ID, bbpars)

# Compile graphs
graphs <- compile_data(
  data = bbmods,
  output_type = "list",
  list_element = 3
)

# Look at 3rd graph
graphs[[3]]

```

fit_g_mc_variableJ *Fitting mesophyll conductance with the variable J method*

Description

Fitting mesophyll conductance with the variable J method

Usage

```

fit_g_mc_variableJ(
  data,
  varnames = list(A_net = "A_net", J_etra = "J_etra", C_i = "C_i", PPFD = "PPFD",
    phi_PSII = "phi_PSII"),
  usealpha_Q = FALSE,
  alpha_Q = 0.84,
  beta_Q = 0.5,
  gamma_star,

```

```
R_d,
P = 100
)
```

Arguments

data	Dataframe
varnames	Variable names to fit g_mc. varnames = list(A_net = "A_net", J_etr = "J_etr", C_i = "C_i", PPFD = "PPFD", phi_PSII = "phi_PSII"), where A_net is net CO2 assimilation in umol m-2 s-1, J_etr is linear electron transport flux in umol m-2 s-1, C_i is intercellular CO2 concentration in umol mol-1, PPFD is incident irradiance in umol m-2 s-1, phi_PSII is the operating efficiency of photosystem II.
usealpha_Q	Recalculate electron transport with new absorbance value?
alpha_Q	Absorbance of photosynthetically active radiation
beta_Q	Partitioning of absorbed light energy between PSI and PSII
gamma_star	Photorespiratory CO2 compensation point in umol mol-1
R_d	Respiration rate in umol m-2 s-1
P	Atmospheric pressure in kPa

Value

fit_g_mc_variableJ fits mesophyll conductance according to Harley et al. 1992. It also tests the reliability of the calculation and calculates a mean with only reliable values. Note that the output is in units of umol m-2 s-1 Pa-1.

References

Harley PC, Loreto F, Di Marco G, Sharkey TD. 1992. Theoretical considerations when estimating mesophyll conductance to CO2 flux by analysis of the response of photosynthesis to CO2. Plant Physiol 98:1429 - 1436.

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Note: there will be issues here if the alpha value used
# for calculating ETR is off, if gamma_star is incorrect,
# if R_d is incorrect.
data <- fit_g_mc_variableJ(data,
  varnames = list(
```

```

A_net = "A",
J_etr = "ETR",
C_i = "Ci",
PPFD = "Qin",
phi_PSII = "PhiPS2"
),
gamma_star = 46,
R_d = 0.153,
usealpha_Q = TRUE,
alpha_Q = 0.84,
beta_Q = 0.5,
P = 84
)

# Note that many g_mc values from this method can be unreliable
ggplot(data, aes(x = CO2_s, y = g_mc, colour = reliable)) +
  labs(
    x = expression(CO[2] ~ "(" * mu * mol ~ mol^
      {
        -1
      } * ")"),
    y = expression(g[m] ~ "(mol" ~ m^{-
      2
    } ~ s^{-
      1
    } ~ Pa^
      {
        -1
      } * ")")
  ) +
  geom_point(size = 2) +
  theme_bw() +
  theme(legend.position = "bottom")

# Plot QAQC graph according to Harley et al. 1992
ggplot(data, aes(x = CO2_s, y = dCcdA, colour = reliable)) +
  labs(
    x = expression(CO[2] ~ "(" * mu * mol ~ mol^
      {
        -1
      } * ")"),
    y = expression(delta * C[chl] * "/" * delta * A)
  ) +
  geom_hline(yintercept = 10) +
  geom_point(size = 2) +
  theme_bw() +
  theme(legend.position = "bottom")

```

Description

Fitting hydraulic vulnerability curves

Usage

```
fit_hydra_vuln_curve(
  data,
  varnames = list(psi = "psi", PLC = "PLC"),
  start_weibull = list(a = 2, b = 2),
  title = NULL
)
```

Arguments

data	Dataframe
varnames	List of variable names. varnames = list(psi = "psi", PLC = "PLC") where psi is water potential in MPa, and PLC is percent loss conductivity.
start_weibull	starting values for the nls fitting routine for the Weibull curve
title	Title for the output graph

Value

fit_hydra_vuln_curve fits a sigmoidal function (Pammenter & Van der Willigen, 1998) linearized according to Ogle et al. (2009). Output is a list containing the sigmoidal model in element 1 and Weibull model in element 4, the fit parameters with 95 element 2, and hydraulic parameters in element 3 (including P25, P50, P88, P95, S50, Pe, Pmax, DSI). Px (25 to 95): water potential at which x conductivity is lost. S50: slope at 50 entry point. Pmax: hydraulic failure threshold. DSI: drought stress interval. Element 5 is a graph showing the fit, P50, Pe, and Pmax.

References

- Ogle K, Barber JJ, Willson C, Thompson B. 2009. Hierarchical statistical modeling of xylem vulnerability to cavitation. *New Phytologist* 182:541-554
- Pammenter NW, Van der Willigen CV. 1998. A mathematical and statistical analysis of the curves illustrating vulnerability of xylem to cavitation. *Tree Physiology* 18:589-593

Examples

```
# Read in data
data <- read.csv(system.file("extdata", "hydraulic_vulnerability.csv",
  package = "photosynthesis"
))

# Fit hydraulic vulnerability curve
fit <- fit_hydra_vuln_curve(data[data$Tree == 4 & data$Plot == "Control", ],
  varnames = list(
    psi = "P",
    PLC = "PLC"
```

```

),
title = "Control 4"
)

# Return Sigmoidal model summary
summary(fit[[1]])

# Return Weibull model summary
summary(fit[[4]])

# Return model parameters with 95\% confidence intervals
fit[[2]]

# Return hydraulic parameters
fit[[3]]

# Return graph
fit[[5]]

# Fit many curves
fits <- fit_many(
  data = data,
  varnames = list(
    psi = "P",
    PLC = "PLC"
  ),
  group = "Tree",
  funct = fit_hydra_vuln_curve
)

# To select individuals from the many fits
# Return model summary
summary(fits[[1]][[1]]) # Returns model summary

# Return sigmoidal model output
fits[[1]][[2]]

# Return hydraulic parameters
fits[[1]][[3]]

# Return graph
fits[[1]][[5]]

# Compile parameter outputs
pars <- compile_data(
  data = fits,
  output_type = "dataframe",
  list_element = 3
)

# Compile graphs
graphs <- compile_data(
  data = fits,

```

```
    output_type = "list",
    list_element = 5
)
```

fit_many*Fitting many functions across groups*

Description

Fitting many functions across groups

Usage

```
fit_many(data, funct, group, ...)
```

Arguments

data	Dataframe
funct	Function to fit
group	Grouping variables
...	Arguments for the function to fit. Use ?functionname to read the help file on available arguments for a given function.

Value

fit_many fits a function across every instance of a grouping variable.

Examples

```
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Define a grouping factor based on light intensity to split the ACi
# curves
data$Q_2 <- as.factor((round(data$Qin, digits = 0)))

# Convert leaf temperature to K
data$T_leaf <- data$Tleaf + 273.15

# Fit many curves
fits <- fit_many(
```

```

data = data,
varnames = list(
  A_net = "A",
  T_leaf = "T_leaf",
  C_i = "Ci",
  PPFD = "Qin"
),
funct = fit_aci_response,
group = "Q_2"
)

# Print the parameters
# First set of double parentheses selects an individual group value
# Second set selects an element of the sublist
fits[[3]][[1]]

# Print the graph
fits[[3]][[2]]

# Compile graphs into a list for plotting
fits_graphs <- compile_data(fits,
  list_element = 2
)

# Compile parameters into dataframe for analysis
fits_pars <- compile_data(fits,
  output_type = "dataframe",
  list_element = 1
)

```

fit_PV_curve*Fitting pressure-volume curves***Description**

Fitting pressure-volume curves

Usage

```

fit_PV_curve(
  data,
  varnames = list(psi = "psi", mass = "mass", leaf_mass = "leaf_mass", bag_mass =
    "bag_mass", leaf_area = "leaf_area"),
  title = NULL
)

```

Arguments

data	Dataframe
varnames	Variable names. varnames = list(psi = "psi", mass = "mass", leaf_mass = "leaf_mass", bag_mass = "bag_mass", leaf_area = "leaf_area") where psi is leaf water potential in MPa, mass is the weighed mass of the bag and leaf in g, leaf_mass is the mass of the leaf in g, bag_mass is the mass of the bag in g, and leaf_area is the area of the leaf in cm ² .
title	Graph title

Value

fit_PV_curve fits pressure-volume curve data to determine: SWC: saturated water content per leaf mass (g H₂O g leaf dry mass ⁻¹), PI_o: osmotic potential at full turgor (MPa), psi_TLP: leaf water potential at turgor loss point (TLP) (MPa), RWC_TLP: relative water content at TLP (C_FT: relative capacitance at full turgor (MPa ⁻¹), C_TLP: relative capacitance at TLP (MPa ⁻¹), and C_FTStar: absolute capacitance per leaf area (g m ⁻² MPa ⁻¹). Element 1 of the output list contains the fitted parameters, element 2 contains the water-psi graph, and element 3 contains the 1/psi-100-RWC graph.

References

- Koide RT, Robichaux RH, Morse SR, Smith CM. 2000. Plant water status, hydraulic resistance and capacitance. In: Plant Physiological Ecology: Field Methods and Instrumentation (eds RW Pearcy, JR Ehleringer, HA Mooney, PW Rundel), pp. 161-183. Kluwer, Dordrecht, the Netherlands
- Sack L, Cowan PD, Jaikumar N, Holbrook NM. 2003. The 'hydrology' of leaves: co-ordination of structure and function in temperate woody species. Plant, Cell and Environment, 26, 1343-1356
- Tyree MT, Hammel HT. 1972. Measurement of turgor pressure and water relations of plants by pressure bomb technique. Journal of Experimental Botany, 23, 267

Examples

```
# Read in data
data <- read.csv(system.file("extdata", "PV_curve.csv",
                             package = "photosynthesis"
))

# Fit one PV curve
fit <- fit_PV_curve(data[data$ID == "L2", ],
                     varnames = list(
                       psi = "psi",
                       mass = "mass",
                       leaf_mass = "leaf_mass",
                       bag_mass = "bag_mass",
                       leaf_area = "leaf_area"
                     )
)

# See fitted parameters
```

```

fit[[1]]

# Plot water mass graph
fit[[2]]

# Plot PV Curve
fit[[3]]

# Fit all PV curves in a file
fits <- fit_many(data,
  group = "ID",
  funct = fit_PV_curve,
  varnames = list(
    psi = "psi",
    mass = "mass",
    leaf_mass = "leaf_mass",
    bag_mass = "bag_mass",
    leaf_area = "leaf_area"
  )
)

# See parameters
fits[[1]][[1]]

# See water mass - water potential graph
fits[[1]][[2]]

# See PV curve
fits[[1]][[3]]

# Compile parameter outputs
pars <- compile_data(
  data = fits,
  output_type = "dataframe",
  list_element = 1
)

# Compile the water mass - water potential graphs
graphs1 <- compile_data(
  data = fits,
  output_type = "list",
  list_element = 2
)

# Compile the PV graphs
graphs2 <- compile_data(
  data = fits,
  output_type = "list",
  list_element = 3
)

```

 fit_r_light_kok *Estimating light respiration*

Description

Estimating light respiration

Usage

```
fit_r_light_kok(
  data,
  varnames = list(A_net = "A_net", PPFD = "PPFD"),
  PPFD_lower = 40,
  PPFD_upper = 100
)

fit_r_light_WalkerOrt(
  data,
  varnames = list(A_net = "A_net", C_i = "C_i", PPFD = "PPFD"),
  P = 100,
  C_i_threshold = 300
)

fit_r_light_yin(
  data,
  varnames = list(A_net = "A_net", PPFD = "PPFD", phi_PSII = "phi_PSII"),
  PPFD_lower = 40,
  PPFD_upper = 100
)
```

Arguments

data	Dataframe
varnames	List of variable names
PPFD_lower	Lower light intensity limit for estimating Rlight (Kok & Yin)
PPFD_upper	Upper light intensity limit for estimating Rlight (Kok & Yin)
P	Atmospheric pressure in kPa (Walker & Ort, 2015)
C_i_threshold	Threshold C_i (in umol mol-1) to cut data to linear region for fitting light respiration and gamma_star (Walker & Ort, 2015)

Value

fit_r_light_kok estimates light respiration using the Kok method (Kok, 1956). The Kok method involves looking for a breakpoint in the light response of net CO₂ assimilation at very low light intensities and extrapolating from data above the breakpoint to estimate light respiration as the y-intercept. r_light value should be negative, denoting an efflux of CO₂.

fit_r_light_WalkerOrt estimates light respiration and GammaStar according to Walk & Ort (2015) using a slope- intercept regression method to find the intercept of multiple ACi curves run at multiple light intensities. Output GammaStar and respiration should be negative If output respiration is positive this could indicate issues (i.e. leaks) in the gas exchange measurements. GammaStar is output in umol mol-1, and respiration is output in umol m-2 s-1 of respiratory flux. Output is a list containing the slope intercept regression model, a graph of the fit, and estimates of the coefficients. NOTE: if using C_i, the output value is technically C_istar. You need to use Cc to get GammaStar. Also note, however, that the convention in the field is to completely ignore this note.

fit_r_light_yin estimates light respiration according to the Yin et al. (2009, 2011) modifications of the Kok method. The modification uses fluorescence data to get a better estimate of light respiration. Note that respiration output should be negative here to denote an efflux of CO₂.

References

- Kok B. 1956. On the inhibition of photosynthesis by intense light. *Biochimica et Biophysica Acta* 21: 234–244
- Walker BJ, Ort DR. 2015. Improved method for measuring the apparent CO₂ photocompensation point resolves the impact of multiple internal conductances to CO₂ to net gas exchange. *Plant Cell Environ* 38:2462- 2474
- Yin X, Struik PC, Romero P, Harbinson J, Evers JB, van der Putten PEL, Vos J. 2009. Using combined measurements of gas exchange and chlorophyll fluorescence to estimate parameters of a biochemical C₃ photosynthesis model: a critical appraisal and a new integrated approach applied to leaves in a wheat (*Triticum aestivum*) canopy. *Plant Cell Environ* 32:448-464
- Yin X, Sun Z, Struik PC, Gu J. 2011. Evaluating a new method to estimate the rate of leaf respiration in the light by analysis of combined gas exchange and chlorophyll fluorescence measurements. *Journal of Experimental Botany* 62: 3489–3499

Examples

```
# FITTING KOK METHOD
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
  package = "photosynthesis"
))

# Fit light respiration with Kok method
r_light <- fit_r_light_kok(
  data = data,
  varnames = list(
    A_net = "A",
    PPFD = "Qin"
  ),
  PPFD_lower = 20,
  PPFD_upper = 150
)
```

```
# Return r_light
r_light

# FITTING WALKER-ORT METHOD
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
                           package = "photosynthesis"
))

# Fit the Walker-Ort method for GammaStar and light respiration
walker_ort <- fit_r_light_WalkerOrt(data,
                                      varnames = list(
                                        A_net = "A",
                                        C_i = "Ci",
                                        PPFD = "Qin"
                                      )
)
# Extract model
summary(walker_ort[[1]])

# View graph
walker_ort[[2]]

# View coefficients
walker_ort[[3]]

# FITTING THE YIN METHOD
# Read in your data
# Note that this data is coming from data supplied by the package
# hence the complicated argument in read.csv()
# This dataset is a CO2 by light response curve for a single sunflower
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
                           package = "photosynthesis"
))

# Fit light respiration with Yin method
r_light <- fit_r_light_yin(
  data = data,
  varnames = list(
    A_net = "A",
    PPFD = "Qin",
    phi_PSII = "PhiPS2"
  ),
  PPFD_lower = 20,
  PPFD_upper = 250
)
```

fit_t_response	<i>Fitting temperature responses</i>
-----------------------	--------------------------------------

Description

Fitting temperature responses

Usage

```
fit_t_response(
  data,
  varnames = list(Par = "Par", T_leaf = "T_leaf"),
  model = c("Arrhenius", "Kruse", "Heskel", "Medlyn", "MMRT", "Quadratic", "Topt"),
  start = list(a = 1, b = 1, c = 1, dEa = 1, Ea_ref = 1, Par_ref = 1, Ea = 40000, Par25
    = 50, Hd = 2e+05, dS = 650, dCp = 1, dG = 1, dH = 1),
  setvar = "none",
  hdset = 2e+05,
  dSset = 650,
  title = NULL,
  ...
)
```

Arguments

data	Dataframe with temperature response variables
varnames	Variable names, where Par is the parameter of interest, and T_leaf is the leaf temperature in K.
model	Which temperature response model do you want to use? Defaults to all: Arrhenius, Heskel, Kruse, Medlyn, MMRT, Quadratic, and Topt.
start	List of starting parameters for the nls model fits. a, b, and c are needed for the Heskel model, dEa, Ea_ref, and Par_ref are needed for the Kruse model, Ea, Par25, and Hd are all needed for the Medlyn and Topt models while the Medlyn model also requires dS, and dCP, dG, and dH are all for the MMRT model.
setvar	Which variable to set as constant for the Medlyn model? Defaults to "none", while "Hd" and "dS" options are available.
hdset	Which value should Hd be set to when setvar = "Hd"? Specify in J/mol.
dSset	Which value should dS be set to when setvar = "dS"? Specify in J/mol/K.
title	Title of output graphs
...	Further arguments to pass on to the nlsLM() function

Value

fit_t_response fits one or more temperature response models to a dataset, returning a list of lists. The parent list contains the models, while the child list for each model contains the fitted model in element 1, the coefficients in element 2, and a graph in element 3.

References

- Arrhenius S. 1915. Quantitative laws in biological chemistry. Bell.
- Heskel MA, O'Sullivan OS, Reich PB, Tjoelker MG, Weerasinghe LK, Penillard A, Egerton JJJG, Creek D, Bloomfield KJ, Xiang J, Sinca F, Stangl ZR, la Torre AM, Griffin KL, Huntingford C, Hurry V, Meir P, Turnbull MH, Atkin OK. 2016. Convergence in the temperature response of leaf respiration across biomes and plant functional types. PNAS 113:3832-3837
- Hobbs JK, Jiao W, Easter AD, Parker EJ, Schipper LA, Arcus VL. 2013. Change in heat capacity for enzyme catalysis determines temperature dependence of enzyme catalyzed rates. ACS Chemical Biology 8:2388-2393.
- Kruse J, Adams MA. 2008. Three parameters comprehensively describe the temperature response of respiratory oxygen reduction. Plant Cell Environ 31:954-967
- Liang LL, Arcus VL, Heskel MA, O'Sullivan OS, Weerasinghe LK, Creek D, Egerton JJJG, Tjoelker MG, Atkin OK, Schipper LA. 2018. Macromolecular rate theory (MMRT) provides a thermodynamics rationale to underpin the convergent temperature response in plant leaf respiration. Glob Chang Biol 24:1538-1547
- Medlyn BE, Dreyer E, Ellsworth D, Forstreuter M, Harley PC, Kirschbaum MUF, Le Roux X, Montpied P, Strassemeyer J, Walcroft A, Wang K, Loutstau D. 2002. Temperature response of parameters of a biochemically based model of photosynthesis. II. A review of experimental data. Plant Cell Environ 25:1167-1179

Examples

```
# Read in data
data <- read.csv(system.file("extdata", "A_Ci_T_data.csv",
  package = "photosynthesis"
),
stringsAsFactors = FALSE
)

library(tidyr)

# Round temperatures to group them appropriately
# Use sequential rounding
data$T2 <- round(data$Tleaf, 1)
data$T2 <- round(data$Tleaf, 0)

# Look at unique values to detect rounding issues
unique(data$T2)

# Some still did not round correctly,
# manually correct
for (i in 1:nrow(data)) {
  if (data$T2[i] == 18) {
    data$T2[i] <- 17
  }
  if (data$T2[i] == 23) {
    data$T2[i] <- 22
  }
}
```

```

if (data$T2[i] == 28) {
  data$T2[i] <- 27
}
if (data$T2[i] == 33) {
  data$T2[i] <- 32
}
if (data$T2[i] == 38) {
  data$T2[i] <- 37
}
}

# Make sure it is a character string for grouping
data$T2 <- as.character(data$T2)

# Create grouping variable by ID and measurement temperature
data <- unite(data,
  col = "ID2", c("ID", "T2"),
  sep = "_"
)

# Split by temperature group
data <- split(data, data$ID2)

# Obtain mean temperature for group so temperature
# response fitting is acceptable later, round to
# 2 decimal places
for (i in 1:length(data)) {
  data[[i]]$Curve_Tleaf <- round(mean(data[[i]]$Tleaf), 2)
}

# Convert from list back to dataframe
data <- do.call("rbind", data)

# Parse grouping variable by ID and measurement temperature
data <- separate(data,
  col = "ID2", into = c("ID", "T2"),
  sep = "_"
)

# Make sure number of values matches number of measurement
# temperatures. May vary slightly if plants had slightly
# different leaf temperatures during the measurements
unique(data$Curve_Tleaf)

# Create ID column to curve fit by ID and temperature
data <- unite(data,
  col = "ID2", c("ID", "Curve_Tleaf"),
  sep = "_"
)

# Convert leaf temperature to K
data$T_leaf <- data$Tleaf + 273.15

```

```
# Fit many CO2 response curves
fits2 <- fit_many(
  data = data,
  group = "ID2",
  varnames = list(
    A_net = "A",
    C_i = "Ci",
    T_leaf = "T_leaf",
    PPFD = "Qin",
    g_mc = "g_mc"
  ),
  funct = fit_aci_response,
  alphag = 0
)

# Extract ACi parameters
pars <- compile_data(fits2,
  output_type = "dataframe",
  list_element = 1
)

# Extract ACi graphs
graphs <- compile_data(fits2,
  output_type = "list",
  list_element = 2
)

# Parse the ID variable
pars <- separate(pars, col = "ID", into = c("ID", "Curve_Tleaf"), sep = "_")

# Make sure curve leaf temperature is numeric
pars$Curve_Tleaf <- as.numeric(pars$Curve_Tleaf)
pars$T_leaf <- pars$Curve_Tleaf + 273.15

# Fit all models, set Hd to constant in Medlyn model
out <- fit_t_response(
  data = pars[pars$ID == "S2", ],
  varnames = list(
    Par = "V_cmax",
    T_leaf = "T_leaf"
  ),
  setvar = "Hd",
  hdset = 200000
)

out[["Arrhenius"]][["Graph"]]
out[["Heskell"]][["Graph"]]
out[["Kruse"]][["Graph"]]
out[["Medlyn"]][["Graph"]]
out[["MMRT"]][["Graph"]]
out[["Quadratic"]][["Graph"]]
out[["Topt"]][["Graph"]]
```

FvCB

Farquhar-von Caemmerer-Berry (FvCB) C3 photosynthesis model

Description

Farquhar-von Caemmerer-Berry (FvCB) C3 photosynthesis model
 Rubisco-limited assimilation rate
 RuBP regeneration-limited assimilation rate
 TPU-limited assimilation rate

Usage

```
FvCB(C_chl, pars, unitless = FALSE)

W_carbox(C_chl, pars, unitless = FALSE)

W_regen(C_chl, pars, unitless = FALSE)

W_tpu(C_chl, pars, unitless = FALSE)
```

Arguments

C_chl	Chloroplastic CO ₂ concentration in Pa of class units
pars	Concatenated parameters (leaf_par, enviro_par, and constants)
unitless	Logical. Should units be set? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.

Details

Equations following Buckley and Diaz-Espejo (2015):

Rubisco-limited assimilation rate:

$$W_{\text{carbox}} = V_{c,\text{max}} C_{\text{chl}} / (C_{\text{chl}} + K_m)$$

where:

$$K_m = K_C (1 + O/K_O)$$

RuBP regeneration-limited assimilation rate:

$$W_{\text{regen}} = J C_{\text{chl}} / (4 C_{\text{chl}} + 8 \Gamma *)$$

where J is a function of PPF, obtained by solving the equation:

$$0 = \theta_J J^2 - J(J_{\max} + \phi_J PPF D) + J_{\max} \phi_J PPF D$$

TPU-limited assimilation rate:

$$W_{\text{tpu}} = 3V_{\text{tpu}}C_{\text{chl}}/(C_{\text{chl}} - \Gamma*)$$

Symbol	R	Description	Units	Default
C_{chl}	C_{chl}	chloroplastic CO ₂ concentration	Pa	input
$\Gamma*$	gamma_star	chloroplastic CO ₂ compensation point (T_leaf)	Pa	calculated
J_{\max}	J_max	potential electron transport (T_leaf)	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$	calculated
K_C	K_C	Michaelis constant for carboxylation (T_leaf)	$\mu\text{mol / mol}$	calculated
K_O	K_O	Michaelis constant for oxygenation (T_leaf)	$\mu\text{mol / mol}$	calculated
O	O	atmospheric O ₂ concentration	kPa	21.27565
ϕ_J	phi_J	initial slope of the response of J to PPFD	none	0.331
PPFD	PPFD	photosynthetic photon flux density	$\mu\text{mol quanta / (m}^2 \text{ s)}$	1500
R_d	R_d	nonphotorespiratory CO ₂ release (T_leaf)	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$	calculated
θ_J	theta_J	curvature factor for light-response curve	none	0.825
$V_{c,\max}$	V_cmax	maximum rate of carboxylation (T_leaf)	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$	calculated
V_{tpu}	V_tpu	rate of triose phosphate utilization (T_leaf)	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$	calculated

Value

A list of four values with units $\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$ of class `units`:

- W_{carbox} : Rubisco-limited assimilation rate
- W_{regen} : RuBP regeneration-limited assimilation rate
- W_{tpu} : TPU-limited assimilation rate
- A: minimum of W_{carbox} , W_{regen} , and W_{tpu}

References

- Buckley TN and Diaz-Espejo A. 2015. Partitioning changes in photosynthetic rate into contributions from different variables. *Plant, Cell & Environment* 38: 1200-11.
- Farquhar GD, Caemmerer S, Berry JA. 1980. A biochemical model of photosynthetic CO₂ assimilation in leaves of C3 species. *Planta* 149: 78–90.

Examples

```
bake_par <- make_bakepar()
constants <- make_constants(use_tealeaves = FALSE)
enviro_par <- make_enviropar(use_tealeaves = FALSE)
leaf_par <- make_leafpar(use_tealeaves = FALSE)
leaf_par <- bake(leaf_par, bake_par, constants)
```

```

pars <- c(leaf_par, enviro_par, constants)
C_chl <- set_units(24.28, "Pa")
FvCB(C_chl, pars)

```

gs_mod_ballberry *Stomatal conductance models*

Description

Stomatal conductance models

Usage

```

gs_mod_ballberry(A_net, C_air, RH)

gs_mod_leuning(A_net, C_air, D0, VPD)

gs_mod_opti(g0, g1, VPD, A_net, C_air)

gs_mod_optifull(g0, g1, gk, VPD, A_net, C_air)

```

Arguments

A_net	Net CO2 assimilation in umol m-2 s-1
C_air	CO2 concentration at the leaf surface in umol mol-1
RH	Relative humidity as a proportion
D0	Vapor pressure sensitivity of stomata (Leuning 1995)
VPD	Vapor pressure deficit (kPa)
g0	Optimization model intercept term (Medlyn et al. 2011)
g1	Optimization model slope term (Medlyn et al. 2011)
gk	Optimization model root term (Medlyn et al. 2011)

Value

gs_mod_ballberry is used for fitting the Ball et al. (1987) model of stomatal conductance

gs_mod_leuning is used for fitting the Leuning (1995) model of stomatal conductance

gs_mod_opti fits the optimal stomatal conductance model according to Medlyn et al. 2011

gs_mod_optifull fits the full optimal stomatal conductance model according to Medlyn et al. 2011

References

- Ball JT, Woodrow IE, Berry JA. 1987. A model predicting stomatal conductance and its contribution to the control of photosynthesis under different environmental conditions, in Progress in Photosynthesis Research, Proceedings of the VII International Congress on Photosynthesis, vol. 4, edited by I. Biggins, pp. 221–224, Martinus Nijhoff, Dordrecht, Netherlands.
- Leuning R. 1995. A critical appraisal of a coupled stomatal- photosynthesis model for C3 plants. *Plant Cell Environ* 18:339-357
- Medlyn BE, Duursma RA, Eamus D, Ellsworth DS, Prentice IC, Barton CVM, Crous KY, Angelis PD, Freeman M, Wingate L. 2011. Reconciling the optimal and empirical approaches to modeling stomatal conductance. *Glob Chang Biol* 17:2134-2144

gw2gc

Convert g_c ($\mu\text{mol CO}_2/\text{m}^2/\text{s}/\text{Pa}$) to g_w ($\mu\text{mol H}_2\text{O}/\text{m}^2/\text{s}/\text{Pa}$)

Description

Convert g_c ($\mu\text{mol CO}_2/\text{m}^2/\text{s}/\text{Pa}$) to g_w ($\mu\text{mol H}_2\text{O}/\text{m}^2/\text{s}/\text{Pa}$)
 Convert g_c ($\mu\text{mol CO}_2/\text{m}^2/\text{s}/\text{Pa}$) to g_w ($\mu\text{mol H}_2\text{O}/\text{m}^2/\text{s}/\text{Pa}$)

Usage

```
gw2gc(g_w, D_c, D_w, unitless)
gc2gw(g_c, D_c, D_w, unitless)
```

Arguments

g_w	conductance to water vapor in units ($\mu\text{mol H}_2\text{O}/(\text{m}^2 \text{s Pa})$) of class units.
D_c	diffusion coefficient for CO2 in air in units of m^2/s of call units
D_w	diffusion coefficient for H2O in air in units of m^2/s of call units
unitless	Logical. Should units be set? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.
g_c	conductance to CO2 in units ($\mu\text{mol H}_2\text{O}/(\text{m}^2 \text{s Pa})$) of class units.

Details

Diffusive conductance to CO2 is generally about ~1.6x that of H2O because of the higher molecular weight. To convert, multiply conductance by the ratio of diffusion coefficients:

$$g_c = g_w D_c / D_w$$

$$g_w = g_c D_w / D_c$$

Value

Value with units $\mu\text{mol}/(\text{m}^2 \text{s Pa})$ of class units.

Note

This function will soon be moving to the standalone gunit package.

Examples

```
D_c <- set_units(1.29e-05, "m^2/s")
D_w <- set_units(2.12e-05, "m^2/s")
g_c <- set_units(3, "umol/m^2/s/Pa")
g_w <- gc2gw(g_c, D_c, D_w, unitless = FALSE)
g_w

gw2gc(g_w, D_c, D_w, unitles = FALSE)
```

J

J: Rate of electron transport (umol/m²/s)

Description

Calculate the rate of electron transport as a function of photosynthetic photon flux density (PPFD).

Usage

```
J(pars, unitless = FALSE)
```

Arguments

pars	Concatenated parameters (leaf_par, enviro_par, and constants)
unitless	Logical. Should units be set? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.

Details

J as a function of PPFD is the solution to the quadratic expression:

$$0 = \theta_J J^2 - J(J_{\max} + \phi_J PPF) + J_{\max}\phi_J PPF$$

Symbol	R	Description	Units	Default
J_{\max}	J_{\max}	potential electron transport (T_leaf)	$\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$	calculated
ϕ_J	ϕ_J	initial slope of the response of <i>J</i> to PPF	none	0.331
PPFD	PPFD	photosynthetic photon flux density	$\mu\text{mol quanta} / (\text{m}^2 \text{ s})$	1500
θ_J	θ_J	curvature factor for light-response curve	none	0.825

Value

Value in $\mu\text{mol}/(\text{m}^2 \text{ s})$ of class units

Examples

```
library(magrittr)
library(photosynthesis)

bake_par <- make_bakepar()
constants <- make_constants(use_tealeaves = FALSE)
enviro_par <- make_enviropar(use_tealeaves = FALSE)
leaf_par <- make_leafpar(use_tealeaves = FALSE)
enviro_par$T_air <- leaf_par$T_leaf
leaf_par %>% bake(bake_par, constants)

pars <- c(leaf_par, enviro_par, constants)
J(pars, FALSE)
```

leaf_par

S3 class leaf_par

Description

Constructor function for leaf_par class. This function ensures that leaf parameter inputs are properly formatted.

Usage

```
leaf_par(.x, use_tealeaves)
```

Arguments

- .x A list to be constructed into **leaf_par**.
- use_tealeaves Logical. Should leaf energy balance be used to calculate leaf temperature (T_leaf)? If TRUE, **tleaf** calculates T_leaf. If FALSE, user-defined T_leaf is used. Additional parameters and constants are required, see [make_parameters](#).

make_parameters

Make lists of parameters for photosynthesis

Description

Make lists of parameters for photosynthesis

[make_leafpar](#)
[make_enviropar](#)
[make_bakepar](#)
[make_constants](#)

Usage

```
make_leafpar(replace = NULL, use_tealeaves)

make_enviropar(replace = NULL, use_tealeaves)

make_bakepar(replace = NULL)

make_constants(replace = NULL, use_tealeaves)
```

Arguments

<code>replace</code>	A named list of parameters to replace defaults. If <code>NULL</code> , defaults will be used.
<code>use_tealeaves</code>	Logical. Should leaf energy balance be used to calculate leaf temperature (<code>T_leaf</code>)? If <code>TRUE</code> , <code>tleaf</code> calculates <code>T_leaf</code> . If <code>FALSE</code> , user-defined <code>T_leaf</code> is used. Additional parameters and constants are required, see make_parameters .

Details**Constants:**

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
$D_{c,0}$	<code>D_c0</code>	diffusion coefficient for CO ₂ in air at 0 °C	m ² / s	1.29e-5
$D_{h,0}$	<code>D_h0</code>	diffusion coefficient for heat in air at 0 °C	m ² / s	1.90e-5
$D_{m,0}$	<code>D_m0</code>	diffusion coefficient for momentum in air at 0 °C	m ² / s	1.33e-5
$D_{w,0}$	<code>D_w0</code>	diffusion coefficient for water vapor in air at 0 °C	m ² / s	2.12e-5
ϵ	<code>epsilon</code>	ratio of water to air molar masses	none	0.622
G	<code>G</code>	gravitational acceleration	m / s ²	9.8
eT	<code>eT</code>	exponent for temperature dependence of diffusion	none	1.75
R	<code>R</code>	ideal gas constant	J / (mol K)	8.3144598
σ	<code>s</code>	Stephan-Boltzmann constant	W / (m ² K ⁴)	5.67e-08
Sh	<code>Sh</code>	Sherwood number	none	calculated

Baking (i.e. temperature response) parameters:

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
$D_{s,gmc}$	<code>Ds_gmc</code>	empirical temperature response parameter	J / (mol K)	487.29
$D_{s,Jmax}$	<code>Ds_Jmax</code>	empirical temperature response parameter	J / (mol K)	388.04
E_{a,Γ^*}	<code>Ea_gammastar</code>	empirical temperature response parameter	J / mol	24459.97
$E_{a,gmc}$	<code>Ea_gmc</code>	empirical temperature response parameter	J / mol	68901.56
$E_{a,Jmax}$	<code>Ea_Jmax</code>	empirical temperature response parameter	J / mol	56095.18
$E_{a,KC}$	<code>Ea_KC</code>	empirical temperature response parameter	J / mol	80989.78
$E_{a,KO}$	<code>Ea_KO</code>	empirical temperature response parameter	J / mol	23719.97
$E_{a,Rd}$	<code>Ea_Rd</code>	empirical temperature response parameter	J / mol	40446.75
$E_{a,Vcmax}$	<code>Ea_Vcmax</code>	empirical temperature response parameter	J / mol	52245.78
$E_{d,gmc}$	<code>Ed_gmc</code>	empirical temperature response parameter	J / mol	148788.56
$E_{d,Jmax}$	<code>Ed_Jmax</code>	empirical temperature response parameter	J / mol	121244.79

Environment parameters:

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
C_{air}	C_{air}	atmospheric CO ₂ concentration	Pa	41
O	O	atmospheric O ₂ concentration	kPa	21.27565
P	P	atmospheric pressure	kPa	101.3246
PPFD	PPFD	photosynthetic photon flux density	umol quanta / (m ² s)	1500
RH	RH	relative humidity	none	0.50
u	wind	windspeed	m / s	2

Leaf parameters:

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
d	leafsize	leaf characteristic dimension	m	0.1
Γ^*	gamma_star	chloroplastic CO ₂ compensation point (T_leaf)	Pa	calculated
Γ^{*25}	gamma_star25	chloroplastic CO ₂ compensation point (25 °C)	Pa	3.743
g_{mc}	g_{mc}	mesophyll conductance to CO ₂ (T_leaf)	μmol CO ₂ / (m ² s Pa)	calculated
g_{mc}	$g_{\text{mc}25}$	mesophyll conductance to CO ₂ (25 °C)	μmol CO ₂ / (m ² s Pa)	4
g_{sc}	g_{sc}	stomatal conductance to CO ₂	μmol CO ₂ / (m ² s Pa)	4
g_{uc}	g_{uc}	cuticular conductance to CO ₂	μmol CO ₂ / (m ² s Pa)	0.1
$J_{\text{max}25}$	$J_{\text{max}25}$	potential electron transport (25 °C)	μmol CO ₂ / (m ² s)	200
J_{max}	J_{max}	potential electron transport (T_leaf)	μmol CO ₂ / (m ² s)	calculated
k_{mc}	k_{mc}	partition of g_{mc} to lower mesophyll	none	1
k_{sc}	k_{sc}	partition of g_{sc} to lower surface	none	1
k_{uc}	k_{uc}	partition of g_{uc} to lower surface	none	1
$K_{\text{C}25}$	$K_{\text{C}25}$	Michaelis constant for carboxylation (25 °C)	μmol / mol	268.3
K_{C}	K_{C}	Michaelis constant for carboxylation (T_leaf)	μmol / mol	calculated
$K_{\text{O}25}$	$K_{\text{O}25}$	Michaelis constant for oxygenation (25 °C)	μmol / mol	165084.2
K_{O}	K_{O}	Michaelis constant for oxygenation (T_leaf)	μmol / mol	calculated
ϕ_J	phi_J	initial slope of the response of J to PPFD	none	0.331
$R_{\text{d}25}$	$R_{\text{d}25}$	nonphotorespiratory CO ₂ release (25 °C)	μmol CO ₂ / (m ² s)	2
R_{d}	R_{d}	nonphotorespiratory CO ₂ release (T_leaf)	μmol CO ₂ / (m ² s)	calculated
θ_J	theta_J	curvature factor for light-response curve	none	0.825
T_{leaf}	T_{leaf}	leaf temperature	K	298.15
$V_{\text{c},\text{max}25}$	$V_{\text{cmax}25}$	maximum rate of carboxylation (25 °C)	μmol CO ₂ / (m ² s)	150
$V_{\text{c},\text{max}}$	V_{cmax}	maximum rate of carboxylation (T_leaf)	μmol CO ₂ / (m ² s)	calculated
$V_{\text{tpu}25}$	$V_{\text{tpu}25}$	rate of triose phosphate utilization (25 °C)	μmol CO ₂ / (m ² s)	200
V_{tpu}	V_{tpu}	rate of triose phosphate utilisation (T_leaf)	μmol CO ₂ / (m ² s)	calculated

If use_tealeaves = TRUE, additional parameters are:

Constants:

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
c_p	c_p	heat capacity of air	J / (g K)	1.01
R_{air}	R_{air}	specific gas constant for dry air	J / (kg K)	287.058

Environmental parameters:

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
---------------	----------	--------------------	--------------	----------------

E_q	E_{q}	energy per mole quanta	kJ / mol^2	22
f_{PAR}	f_{par}	fraction of incoming shortwave radiation that is photosynthetically active radiation (PAR)	none	0.5
r	r	reflectance for shortwave irradiance (albedo)	none	0.2
T_{air}	T_{air}	air temperature	K	29

Leaf parameters:

<i>Symbol</i>	<i>R</i>	<i>Description</i>	<i>Units</i>	<i>Default</i>
α_l	abs_l	absorbivity of longwave radiation (4 - 80 μm)	none	0.97
α_s	abs_s	absorbivity of shortwave radiation (0.3 - 4 μm)	none	0.50
g_{sw}	g_{sw}	stomatal conductance to H ₂ O	($\mu\text{mol H}_2\text{O}$) / ($\text{m}^2 \text{ s Pa}$)	converted from g_{sc}
g_{uw}	g_{uw}	cuticular conductance to H ₂ O	($\mu\text{mol H}_2\text{O}$) / ($\text{m}^2 \text{ s Pa}$)	converted from g_{uc}
$\text{logit}(sr)$	logit_sr	stomatal ratio (logit transformed)	none	converted from k_{sc}

Value

`make_leafpar`: An object inheriting from class `leaf_par`
`make_enviropar`: An object inheriting from class `enviro_par`
`make_bakepar`: An object inheriting from class `bake_par`
`make_constants`: An object inheriting from class `constants`

References

Buckley TN and Diaz-Espejo A. 2015. Partitioning changes in photosynthetic rate into contributions from different variables. Plant, Cell & Environment 38: 1200-11.

Examples

```
bake_par <- make_bakepar()
constants <- make_constants(use_tealeaves = FALSE)
enviro_par <- make_enviropar(use_tealeaves = FALSE)
leaf_par <- make_leafpar(use_tealeaves = FALSE)

leaf_par <- make_leafpar(
  replace = list(
    g_sc = set_units(3, "umol/m^2/s/Pa"),
    V_cmax25 = set_units(100, "umol/m^2/s")
  ), use_tealeaves = FALSE
)
```

parameter_names *Get vector of parameter names*

Description

Get vector of parameter names

Usage

```
parameter_names(which, use_tealeaves)
```

Arguments

- which A character string indicating which parameter names to retrieve: "leaf", "enviro", "bake", or "constants". Partial matching allowed.
- use_tealeaves Logical. Should leaf energy balance be used to calculate leaf temperature (T_{leaf})? If TRUE, `tleaf` calculates T_{leaf} . If FALSE, user-defined T_{leaf} is used. Additional parameters and constants are required, see [make_parameters](#).

Examples

```
parameter_names("leaf", use_tealeaves = FALSE)
```

photosynthesis	<i>Simulate C3 photosynthesis</i>
----------------	-----------------------------------

Description

- photosynthesis: simulate C3 photosynthesis over multiple parameter sets
photo: simulate C3 photosynthesis over a single parameter set

Usage

```
photosynthesis(  
  leaf_par,  
  enviro_par,  
  bake_par,  
  constants,  
  use_tealeaves,  
  progress = TRUE,  
  quiet = FALSE,  
  assert_units = TRUE,  
  parallel = FALSE  
)  
  
photo(  
  leaf_par,  
  enviro_par,  
  bake_par,  
  constants,  
  use_tealeaves,  
  quiet = FALSE,  
  assert_units = TRUE,
```

```

check = TRUE,
prepare_for_tleaf = use_tealeaves
)

```

Arguments

<code>leaf_par</code>	A list of leaf parameters inheriting class <code>leaf_par</code> . This can be generated using the <code>make_leafpar</code> function.
<code>enviro_par</code>	A list of environmental parameters inheriting class <code>enviro_par</code> . This can be generated using the <code>make_enviropar</code> function.
<code>bake_par</code>	A list of temperature response parameters inheriting class <code>bake_par</code> . This can be generated using the <code>make_bakepar</code> function.
<code>constants</code>	A list of physical constants inheriting class <code>constants</code> . This can be generated using the <code>make_constants</code> function.
<code>use_tealeaves</code>	Logical. Should leaf energy balance be used to calculate leaf temperature (<code>T_leaf</code>)? If TRUE, <code>tleaf</code> calculates <code>T_leaf</code> . If FALSE, user-defined <code>T_leaf</code> is used. Additional parameters and constants are required, see make_parameters .
<code>progress</code>	Logical. Should a progress bar be displayed?
<code>quiet</code>	Logical. Should messages be displayed?
<code>assert_units</code>	Logical. Should parameter units be checked? The function is faster when FALSE, but input must be in correct units or else results will be incorrect without any warning.
<code>parallel</code>	Logical. Should parallel processing be used via future_map ?
<code>check</code>	Logical. Should arguments checks be done? This is intended to be disabled when <code>photo</code> is called from <code>photosynthesis</code> . Default is TRUE.
<code>prepare_for_tleaf</code>	Logical. Should arguments additional calculations for <code>tleaf</code> ? This is intended to be disabled when <code>photo</code> is called from <code>photosynthesis</code> . Default is <code>use_tealeaves</code> .

Details

`photo`: This function takes simulates photosynthetic rate using the Farquhar-von Caemmerer-Berry (FvCB) model of C3 photosynthesis for single combined set of leaf parameters (`leaf_par`), environmental parameters (`enviro_par`), and physical constants (`constants`). Leaf parameters are provided at reference temperature (25 °C) and then "baked" to the appropriate leaf temperature using temperature response functions (see `bake`).

`photosynthesis`: This function uses `photo` to simulate photosynthesis over multiple parameter sets that are generated using `cross_df`.

Value

A data.frame with the following units columns

Input:

C_air	atmospheric CO2 concentration (Pa)
g_mc25	mesophyll conductance to CO2 at 25 °C ($\mu\text{mol} / (\text{m}^2 \text{ s Pa})$)
g_sc	stomatal conductance to CO2 ($\mu\text{mol} / (\text{m}^2 \text{ s Pa})$)
g_uc	cuticular conductance to CO2 ($\mu\text{mol} / (\text{m}^2 \text{ s Pa})$)
gamma_star25	chloroplastic CO2 compensation point at 25 °C (Pa)
J_max25	potential electron transport at 25 °C ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
K_C25	Michaelis constant for carboxylation at 25 °C ($\mu\text{mol} / \text{mol}$)
K_O25	Michaelis constant for oxygenation at 25 °C ($\mu\text{mol} / \text{mol}$)
k_mc	partition of g_{mc} to lower mesophyll (unitless)
k_sc	partition of g_{sc} to lower surface (unitless)
k_uc	partition of g_{uc} to lower surface (unitless)
leafsize	leaf characteristic dimension (m)
O	atmospheric O2 concentration (kPa)
P	atmospheric pressure (kPa)
phi_J	initial slope of the response of J to PPFD (unitless)
PPFD	photosynthetic photon flux density ($\mu\text{mol quanta} / (\text{m}^2 \text{ s})$)
R_d25	nonphotorespiratory CO2 release at 25 °C ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
RH	relative humidity (unitless)
theta_J	curvature factor for light-response curve (unitless)
T_air	air temperature (K)
T_leaf	leaf temperature (K)
V_cmax25	maximum rate of carboxylation at 25 °C ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
V_tpu25	rate of triose phosphate utilization at 25 °C ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
wind	wind speed (m / s)

Baked Input:

g_mc	mesophyll conductance to CO2 at T_leaf ($\mu\text{mol} / (\text{m}^2 \text{ s Pa})$)
gamma_star	chloroplastic CO2 compensation point at T_leaf (Pa)
J_max	potential electron transport at T_leaf ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
K_C	Michaelis constant for carboxylation at T_leaf ($\mu\text{mol} / \text{mol}$)
K_O	Michaelis constant for oxygenation at T_leaf ($\mu\text{mol} / \text{mol}$)
R_d	nonphotorespiratory CO2 release at T_leaf ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
V_cmax	maximum rate of carboxylation at T_leaf ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
V_tpu	rate of triose phosphate utilisation at T_leaf ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)

Output:

A	photosynthetic rate at C_chl ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s})$)
C_chl	chloroplastic CO2 concentration where A_supply intersects A_demand (Pa)
g_tc	total conductance to CO2 at T_leaf ($\mu\text{mol CO}_2 / (\text{m}^2 \text{ s Pa})$)
value	A_supply - A_demand ($\mu\text{mol} / (\text{m}^2 \text{ s})$) at C_chl
convergence	convergence code (0 = converged)

Examples

```
# Single parameter set with 'photo'
```

```

bake_par <- make_bakepar()
constants <- make_constants(use_tealeaves = FALSE)
enviro_par <- make_enviropar(use_tealeaves = FALSE)
leaf_par <- make_leafpar(use_tealeaves = FALSE)
photo(leaf_par, enviro_par, bake_par, constants,
      use_tealeaves = FALSE
    )

# Multiple parameter sets with 'photosynthesis'

leaf_par <- make_leafpar(
  replace = list(
    T_leaf = set_units(c(293.14, 298.15), "K")
  ), use_tealeaves = FALSE
)
photosynthesis(leaf_par, enviro_par, bake_par, constants,
      use_tealeaves = FALSE
)

```

ppm2pa

*Convert pressure from PPM to Pascals***Description**

Convert pressure from PPM to Pascals

Usage

```
ppm2pa(ppm, P)
```

Arguments

ppm	Pressure value in umol/mol of class units
P	Atmospheric pressure value in kPa of class units

Details

$$\text{Press}(kPa) = \text{Press}(ppm)P(kPa)$$

$$\text{Press}(Pa) = 1000\text{Press}(kPa)$$

Value

Value in Pa of class units

Examples

```

ppm <- set_units(400, "umol/mol")
P <- set_units(101.325, "kPa")
ppm2pa(ppm, P)

```

print_graphs	<i>Printing graphs to system</i>
--------------	----------------------------------

Description

Printing graphs to system

Usage

```
print_graphs(  
  data,  
  path,  
  output_type = "jpeg",  
  height = 5,  
  width = 5,  
  res = 600,  
  units = "in",  
  pdf_filename,  
  ...  
)
```

Arguments

data	List of graphs
path	File path for printing our graphs. Use "./" to set to current working directory
output_type	Type of output file, jpeg or pdf
height	Height of jpegs
width	Width of jpegs
res	Resolution of jpegs
units	Units of height and width
pdf_filename	Filename for pdf option
...	Further arguments for jpeg() and pdf()

Value

print_graphs creates graph files in current working directory from a list of graphs

Examples

```
# Read in your data  
# Note that this data is coming from data supplied by the package  
# hence the complicated argument in read.csv()  
# This dataset is a CO2 by light response curve for a single sunflower  
data <- read.csv(system.file("extdata", "A_Ci_Q_data_1.csv",
```

```

package = "photosynthesis"
))

# Fit many AQ curves
# Set your grouping variable
# Here we are grouping by CO2_s and individual
data$C_s <- (round(data$CO2_s, digits = 0))

# For this example we need to round sequentially due to CO2_s setpoints
data$C_s <- as.factor(round(data$C_s, digits = -1))

# To fit one AQ curve
fit <- fit_aq_response(data[data$C_s == 600, ],
  varnames = list(
    A_net = "A",
    PPFD = "Qin"
  )
)

# Print model summary
summary(fit[[1]])

# Print fitted parameters
fit[[2]]

# Print graph
fit[[3]]

# Fit many curves
fits <- fit_many(
  data = data,
  varnames = list(
    A_net = "A",
    PPFD = "Qin",
    group = "C_s"
  ),
  funct = fit_aq_response,
  group = "C_s"
)

# Look at model summary for a given fit
# First set of double parentheses selects an individual group value
# Second set selects an element of the sublist
summary(fits[[3]][[1]])

# Print the parameters
fits[[3]][[2]]

# Print the graph
fits[[3]][[3]]

# Compile graphs into a list for plotting
fits_graphs <- compile_data(fits,

```

```
    list_element = 3
)

# Print graphs to pdf
# Uncomment to run
# print_graphs(data = fits_graphs,
#               output_type = "pdf",
#               path = tempdir(),
#               pdf_filename = "mygraphs.pdf")
```

read_li6800*Reading in LiCor files*

Description

Reading in LiCor files

Usage

```
read_li6800(x)
```

Arguments

x File name

Value

Returns a dataframe from raw LiCor files. Current support for LiCor 6800 files only. LiCor 6400 file reading will be supported in a later version.

t_response_arrhenius *Temperature response functions*

Description

Temperature response functions

Usage

```
t_response_arrhenius(T_leaf, Ea)
t_response_arrhenius_kruse(dEa, Ea_ref, Par_ref, T2)
t_response_arrhenius_medlyn(T_leaf, Ea, Hd, dS)
t_response_arrhenius_topt(T_leaf, Ea, Hd, Topt)
```

```

t_response_calc_dS(Ea, Hd, Topt)
t_response_calc_topt(Hd, dS, Ea)
t_response_heskel(T_leaf, a, b, c)
t_response_mmrt(dCp, dG, dH, T_leaf)

```

Arguments

T_leaf	Leaf temperature in K
Ea	Activation energy in J mol-1 (Medlyn et al. 2002)
dEa	Temperature-dependent change in Ea in K^2 (Kruse et al. 2008)
Ea_ref	Activation energy in J mol-1 (Kruse et al. 2008)
Par_ref	Parameter at reference temperature of 25 Celsius (Kruse et al. 2008)
T2	Leaf temperature term (Kruse et al. 2008)
Hd	Deactivation energy in J mol-1 (Medlyn et al. 2002)
dS	Entropy parameter in J mol-1 (Medlyn et al. 2002)
Topt	Optimum temperature of the process in K (Medlyn et al. 2002)
a	Constant to minimize residuals (Heskel et al. 2016)
b	Linear coefficient to minimize residuals (Heskel et al. 2016)
c	Quadratic coefficient to minimize residuals (Heskel et al. 2016)
dCp	Change in heat capacity of the enzyme between the enzyme-substrate #' and enzyme-transition states in J mol-1 K-1 (Hobbs et al. 2013)
dG	Change in Gibbs free energy of the reaction at 25 C in J mol-1 (Hobbs et al. 2013)
dH	Change in enthalpy of the reaction at 25 C in J mol-1 (Hobbs et al. 2013)

Value

t_response_arrhenius calculates the rate of a process based on an Arrhenius-type curve
t_response_arrhenius_kruse fits a peaked Arrhenius response according to Kruse et al. 2008.
t_response_arrhenius_medlyn is a peaked Arrhenius response as found in Medlyn et al. 2002.
t_response_arrhenius_topt is a peaked Arrhenius temperature response function.
t_response_calc_dS calculates dS from the fitted Topt model.
t_response_calc_topt calculates Topt for a process from Arrhenius parameters.
t_response_heskel is a quadratic temperature response according to Heskel et al. 2016.
t_response_mmrt is a macromolecular rate theory temperature response according to Hobbs et al. 2013.

References

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- Heskel et al. 2016. Convergence in the temperature response of leaf respiration across biomes and plant functional types. PNAS 113:3832-3837
- Hobbs et al. 2013. Change in heat capacity for enzyme catalysis determines temperature dependence of enzyme catalyzed rates. ACS Chemical Biology 8:2388-2393
- Kruse J, Adams MA. 2008. Three parameters comprehensively describe the temperature response of respiratory oxygen reduction. Plant Cell Environ 31:954-967
- Medlyn BE, Dreyer E, Ellsworth D, Forstreuter M, Harley PC, Kirschbaum MUF, Le Roux X, Montpied P, Strassemeyer J, Walcroft A, Wang K, Loutstau D. 2002. Temperature response of parameters of a biochemically based model of photosynthesis. II. A review of experimental data. Plant Cell Environ 25:1167-1179

Index

.get_gbc, 12
.get_gbc (CO2_conductance), 11
.get_gmc (CO2_conductance), 11
.get_gsc (CO2_conductance), 11
.get_gtc (CO2_conductance), 11
.get_guc (CO2_conductance), 11

A_demand (A_supply), 6
A_supply, 6
analyze_sensitivity, 3
aq_response, 6

bake, 8, 9, 52
bake_par, 9, 10, 50
baked-class, 9

calculate_j (calculate_jmax), 10
calculate_jmax, 10
calculated, 7, 12, 43, 46, 48, 49
check_dependencies, 11
CO2_conductance, 11
compile_data, 13
compute_sensitivity, 14
constants, 16, 50, 52
cross_df, 52

enviro_par, 17, 50, 52

fit_aci_response, 17
fit_aq_response, 21
fit_g_mc_variableJ, 26
fit_gs_model, 23
fit_hydra_vuln_curve, 28
fit_many, 31
fit_PV_curve, 32
fit_r_light_kok, 35
fit_r_light_WalkerOrt
 (fit_r_light_kok), 35
fit_r_light_yin(fit_r_light_kok), 35
fit_t_response, 38
future_map, 52

FvCB, 42, 52

gc2gw (gw2gc), 45
gs_mod_ballberry, 44
gs_mod_leuning (gs_mod_ballberry), 44
gs_mod_opti (gs_mod_ballberry), 44
gs_mod_optifull (gs_mod_ballberry), 44
gw2gc, 45

J, 46

leaf_par, 8, 9, 47, 50, 52

make_bakepar (make_parameters), 47
make_constants, 8
make_constants (make_parameters), 47
make_enviropar (make_parameters), 47
make_leafpar (make_parameters), 47
make_parameters, 16, 17, 47, 47, 48, 51, 52

parameter_names, 50
photo, 52
photo (photosynthesis), 51
photosynthesis, 51, 52
photosynthesis-package, 3
ppm2pa, 54
print_graphs, 55

read_li6800, 57

t_response_arrhenius, 57
t_response_arrhenius_kruse
 (t_response_arrhenius), 57
t_response_arrhenius_medlyn
 (t_response_arrhenius), 57
t_response_arrhenius_topt
 (t_response_arrhenius), 57
t_response_calc_dS
 (t_response_arrhenius), 57
t_response_calc_topt
 (t_response_arrhenius), 57

t_response_heskel
 (t_response_arrhenius), 57
t_response_mmrt (t_response_arrhenius),
 57
temp_resp1 (bake), 8
temp_resp2 (bake), 8
tleaf, 16, 17, 47, 48, 51, 52

W_carbox (FvCB), 42
W_regen (FvCB), 42
W_tpu (FvCB), 42