

# Package ‘ggspectra’

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**Type** Package

**Title** Extensions to 'ggplot2' for Radiation Spectra

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**Description** Additional annotations, stats, geoms and scales for plotting ``light'' spectra with 'ggplot2', together with specializations of ggplot() and autoplot() methods for spectral data and waveband definitions stored in objects of classes defined in package 'photobiology'. Part of the 'r4photobiology' suite, Aphalo P. J. (2015) <doi:10.19232/uv4pb.2015.1.14>.

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**Imports** photobiologyWavebands (>= 0.5.2), scales (>= 1.2.0), ggrepel (>= 0.9.2), lubridate (>= 1.9.0), rlang (>= 1.0.2), tibble (>= 3.1.5)

**Suggests** knitr (>= 1.38), rmarkdown (>= 2.13), magrittr (>= 2.0.3)

**URL** <https://docs.r4photobiology.info/ggspectra/>,  
<https://github.com/aphalo/ggspectra/>

**BugReports** <https://github.com/aphalo/ggspectra/issues/>

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ggspectra-package      *ggspectra: Extensions to 'ggplot2' for Radiation Spectra*

---

**Description**

Additional annotations, stats, geoms and scales for plotting "light" spectra with 'ggplot2', together with specializations of ggplot() and autoplot() methods for spectral data and waveband definitions stored in objects of classes defined in package 'photobiology'. Part of the 'r4photobiology' suite, Aphalo P. J. (2015) [doi:10.19232/uv4pb.2015.1.14](https://doi.org/10.19232/uv4pb.2015.1.14).

**Details**

Package 'ggspectra' provides a set of layer functions and autoplot() methods extending packages 'ggplot2' and 'photobiology'. The autoplot() methods specialised for objects of classes defined in package 'photobiology' facilitate in many respects the plotting of spectral data. The ggplot() methods specialised for objects of classes defined in package 'photobiology' combined with the new layer functions and scales easy the task of flexibly plotting radiation-related spectra and of annotating the resulting plots.

**These methods, layer functions and scales are specialized and work only with certain types of data and ways of expressing physical quantities. Most importantly, all statistics expect the values mapped to the x aesthetic to be wavelengths expressed in nanometres (nm), which is ensured when the data are stored in data objects of classes defined in package 'photobiology'. The support for scale transforms is manual and only partial. Flipping is not supported.**

Although originally aimed at plots relevant to photobiology, many of the functions in the package are also useful for plotting other UV, VIS and NIR spectra of light emission, transmittance, reflectance, absorptance, and responses.

The available summary quantities are both simple statistical summaries and response-weighted summaries. Simple derived quantities represent summaries of a given range of wavelengths, and can be expressed either in energy or photon based units. Derived biologically effective quantities are used to quantify the effect of radiation on different organisms or processes within organisms. These effects can range from damage to perception of informational light signals. Additional features of spectra may be important and worthwhile annotating in plots. Of these, local maxima (peaks), minima (valleys) and spikes present in spectral data can also be annotated with statistics from 'ggspectra'.

Package 'ggspectra' is useful solely for plotting spectral data as most functions depend on the x aesthetic being mapped to a variable containing wavelength values expressed in nanometres. It works well together with many other extensions to package 'ggplot2' such as packages 'ggrepel', 'gganimate' and 'cowplot'.

This package is part of a suite of R packages for photobiological calculations described at the [r4photobiology](<https://www.r4photobiology.info>) web site.

### Note

This package makes use of the new features of 'ggplot2' >= 2.0.0 that make writing this kind of extensions easy and is consequently not compatible with earlier versions of 'ggplot2'.

### Author(s)

**Maintainer:** Pedro J. Aphalo <[pedro.aphalo@helsinki.fi](mailto:pedro.aphalo@helsinki.fi)> ([ORCID](#))

Other contributors:

- Titta K. Kotilainen ([ORCID](#)) [contributor]

### References

Aphalo, Pedro J. (2015) The r4photobiology suite. UV4Plants Bulletin, 2015:1, 21-29. [doi:10.19232/uv4pb.2015.1.14](https://doi.org/10.19232/uv4pb.2015.1.14).

ggplot2 web site at <https://ggplot2.tidyverse.org/>

ggplot2 source code at <https://github.com/tidyverse/ggplot2>

Function multiplot from <http://www.cookbook-r.com/>

### See Also

Useful links:

- <https://docs.r4photobiology.info/ggspectra/>
- <https://github.com/aphalo/ggspectra/>
- Report bugs at <https://github.com/aphalo/ggspectra/issues/>

### Examples

```
library(photobiologyWavebands)
```

```
ggplot(sun.spct) +  
  geom_line() +
```

```

stat_peaks(span = NULL)

ggplot(sun.spct, aes(w.length, s.e.irrad)) +
  geom_line() +
  stat_peaks(span = 21, geom = "point", colour = "red") +
  stat_peaks(span = 51, geom = "text", colour = "red", vjust = -0.3,
             label.fmt = "%3.0f nm")

ggplot(polyester.spct, range = UV()) + geom_line()

autoplot(sun.spct)

autoplot(polyester.spct,
         UV_bands(),
         range = UV(),
         annotations = c("=", "segments", "labels"))

```

---

Afr\_label

*Absorptance axis labels*


---

## Description

Generate cps axis labels in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

## Usage

```

Afr_label(
  unit.exponent = ifelse(pc.out, -2, 0),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.Afr"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE)
)

Rfr_total_label(
  unit.exponent = ifelse(pc.out, -2, 0),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE)
)

```

**Arguments**

<code>unit.exponent</code>	integer
<code>format</code>	character string, "R", "R.expression", "R.character", or "LaTeX".
<code>label.text</code>	character Textual portion of the labels.
<code>scaled</code>	logical If TRUE relative units are assumed.
<code>normalized</code>	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
<code>axis.symbols</code>	logical If TRUE symbols of the quantities are added to the name. Supported only by <code>format = "R.expression"</code> .
<code>pc.out</code>	logical, if TRUE use percent as default instead of fraction of one.

**Value**

a character string or an R expression.

**Examples**

```
Afr_label()
Afr_label(format = "R.expression", axis.symbols = FALSE)
Afr_label(-2)
Afr_label(-3)
Afr_label(format = "R.expression")
Afr_label(format = "LaTeX")
Afr_label(-2, format = "LaTeX")
```

```
Rfr_total_label()
Rfr_total_label(axis.symbols = FALSE)
Rfr_total_label(-2)
Rfr_total_label(-3)
Rfr_total_label(format = "R.expression")
Rfr_total_label(format = "LaTeX")
Rfr_total_label(-3, format = "LaTeX")
```

---

autoplot.calibration\_spct

*Plot one or more irradiance-calibration spectra.*

---

**Description**

These methods return a ggplot object with an annotated plot of the spectral data contained in a `calibration_spct` or a `calibration_mspct` object.

**Usage**

```

## S3 method for class 'calibration_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default = list(UVC(), UVB(), UVA(),
    PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  unit.out = "ignored",
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = "mean",
  span = NULL,
  wls.target = "HM",
  annotations = NULL,
  geom = "line",
  time.format = "",
  tz = "UTC",
  norm = NA,
  text.size = 2.5,
  idfactor = NULL,
  facets = FALSE,
  plot.data = "as.is",
  ylim = c(NA, NA),
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

## S3 method for class 'calibration_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  unit.out = "ignored",
  norm = NA,
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  plot.data = "as.is",
  idfactor = TRUE,
  facets = FALSE,
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

```

**Arguments**

object	a calibration_spct object or a calibration_mspct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.

range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
unit.out	character IGNORED.
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections Plot <b>Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
norm	numeric or character. Normalization to apply before plotting, If object is already normalized, the normalization is updated when a unit conversion applied.
text.size	numeric size of text in the plot decorations.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spct contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.
ylim	numeric y axis limits,
object.label	character The name of the object being plotted.
na.rm	logical.

## Details

The plot object returned is a ggplot (an object of class "gg") and it can be added to or modified as any other ggplot. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', plotmath expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless reserve.space is passed to parameter annotations.

The generic of the `autoplot` method is defined in package `'ggplot2'`. Package `'ggspectra'` defines specializations for the different classes for storage of spectral data defined in package `photobiology`.

For details about normalization and arguments to parameter `norm`, please, see `normalize`. If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with `'photobiology'` ( $\leq 0.10.9$ ) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with `fscale` methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named `"spct.idx"`, but if a different name is passed, it will be used instead.

## Value

A `ggplot` object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

## Plot Annotations

The recognized annotation names are: `"summaries"`, `"peaks"`, `"peak.labels"`, `"valleys"`, `"valley.labels"`, `"wls"`, `"wls.labels"`, `"colour.guide"`, `"color.guide"`, `"boxes"`, `"segments"`, `"labels"`. In addition, `"+"` is interpreted as a request to add to the already present default annotations, `"-"` as request to remove annotations and `"="` or missing `"+"` and `"-"` as a request to reset annotations to those requested. If used, `"+"`, `"-"` or `"="` must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass `" "` or `c("=", " ")` as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding `"peak.labels"` replaces `"peaks"` if present.

## Title Annotations

metadata retrieved from object `object` is passed to `ggplot2::ggtitle()` as arguments for `title`, `subtitle` and `caption`. The specification for the title is passed as argument to `annotations`, and consists in the keyword `title` with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to `title`, `subtitle` and `caption`. The recognized keywords are: `"objt"`, `"class"`, `"what"`, `"when"`, `"where"`, `"how"`, `"inst.name"`, `"inst.sn"`, `"comment"` and `"none"` are recognized as modifiers to `"title"`; `"none"` is a placeholder. Default is `"title:objt"` or no title depending on the context.

## See Also

`normalize`, `calibration_spct`, `waveband`, `photobiologyWavebands-package` and `autoplot`

Other `autoplot` methods: `autoplot.cps_spct()`, `autoplot.filter_spct()`, `autoplot.object_spct()`, `autoplot.raw_spct()`, `autoplot.reflector_spct()`, `autoplot.response_spct()`, `autoplot.source_spct()`, `autoplot.waveband()`, `set_annotations_default()`

**Examples**

```
# to be added
```

---

```
autoplot.cps_spct      Plot one or more detector-counts-per-second spectra.
```

---

**Description**

These methods return a ggplot object with an annotated plot of a cps\_spct or a cps\_mspct object.

**Usage**

```
## S3 method for class 'cps_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default = list(UVC(), UVB(), UVA(),
    PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  unit.out = NULL,
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = "mean",
  span = NULL,
  wls.target = "HM",
  annotations = NULL,
  geom = "line",
  time.format = "",
  tz = "UTC",
  text.size = 2.5,
  idfactor = NULL,
  facets = FALSE,
  plot.data = "as.is",
  ylim = c(NA, NA),
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)
```

```
## S3 method for class 'cps_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  unit.out = NULL,
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
```

```

    idfactor = TRUE,
    facets = FALSE,
    plot.data = "as.is",
    object.label = deparse(substitute(object)),
    na.rm = TRUE
  )

```

## Arguments

object	a cps_spct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting, If object is already normalized, the normalization is updated when a unit conversion applied.
unit.out	character IGNORED.
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections Plot <b>Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
text.size	numeric size of text in the plot decorations.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spct contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.

<code>ylim</code>	numeric y axis limits,
<code>object.label</code>	character The name of the object being plotted.
<code>na.rm</code>	logical.

## Details

The plot object returned is a `ggplot` (an object of class "gg") and it can be added to or modified as any other `ggplot`. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', *plotmath* expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter annotations.

The generic of the `autoplot` method is defined in package 'ggplot2'. Package 'ggspectra' defines specializations for the different classes for storage of spectral data defined in package `photobiology`.

For details about normalization and arguments to parameter `norm`, please, see `normalize`. If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with 'photobiology' (<= 0.10.9) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with `fscale` methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named "spect.idx", but if a different name is passed, it will be used instead.

## Value

A `ggplot` object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

## Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or `c("=", "")` as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

## Title Annotations

metadata retrieved from object `object` is passed to `ggplot2::ggtitle()` as arguments for `title`, `subtitle` and `caption`. The specification for the title is passed as argument to annotations,

and consists in the keyword `title` with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title, subtitle and caption. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

### See Also

[normalize](#), [cps\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: [autoplot.calibration\\_spct\(\)](#), [autoplot.filter\\_spct\(\)](#), [autoplot.object\\_spct\(\)](#), [autoplot.raw\\_spct\(\)](#), [autoplot.reflector\\_spct\(\)](#), [autoplot.response\\_spct\(\)](#), [autoplot.source\\_spct\(\)](#), [autoplot.waveband\(\)](#), [set\\_annotatations\\_default\(\)](#)

### Examples

```
autoplot(white_led.cps_spct)
autoplot(white_led.cps_spct, geom = "spct")
autoplot(normalize(white_led.cps_spct, norm = "max"))

two_leds.mspct <-
  cps_mspct(list("LED 1" = white_led.cps_spct,
                "LED 2" = white_led.cps_spct / 2))
autoplot(two_leds.mspct)
autoplot(two_leds.mspct, idfactor = "Spectra")
autoplot(two_leds.mspct, plot.data = "mean")
```

---

`autoplot.filter_spct` *Plot one or more "filter" spectra.*

---

### Description

These methods return a ggplot object of an annotated plot from spectral data contained in a `filter_spct` or a `filter_mspct` object. Data can be expressed as absorbance, absorptance or transmittance.

### Usage

```
## S3 method for class 'filter_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default = list(UVC(), UVB(), UVA(),
    PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  plot.qty = getOption("photobiology.filter.qty", default = "transmittance"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = NULL,
```

```

span = NULL,
wls.target = "HM",
annotations = NULL,
geom = "line",
time.format = "",
tz = "UTC",
text.size = 2.5,
chroma.type = "CMF",
idfactor = NULL,
facets = FALSE,
plot.data = "as.is",
ylim = c(NA, NA),
object.label = deparse(substitute(object)),
na.rm = TRUE
)

## S3 method for class 'filter_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  plot.qty = getOption("photobiology.filter.qty", default = "transmittance"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  plot.data = "as.is",
  idfactor = TRUE,
  facets = FALSE,
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

```

### Arguments

object	a filter_spect object or a filter_mspct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting. If object is already normalized, the normalization is updated when a unit conversion applied.
plot.qty	character string one of "transmittance" or "absorbance".
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".

span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections <b>Plot Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
text.size	numeric size of text in the plot decorations.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spct contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.
ylim	numeric y axis limits,
object.label	character The name of the object being plotted.
na.rm	logical.

## Details

The plot object returned is a ggplot (an object of class "gg") and it can be added to or modified as any other ggplot. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', plotmath expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter `annotations`.

The generic of the [autoplot](#) method is defined in package 'ggplot2'. Package 'ggspectra' defines specializations for the different classes for storage of spectral data defined in package [photobiology](#).

For details about normalization and arguments to parameter `norm`, please, see [normalize](#). If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with 'photobiology' (<= 0.10.9) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with [fscale](#) methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does

not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named "spct.idx", but if a different name is passed, it will be used instead.

### Value

A ggplot object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

### Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or c("=", "") as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

### Title Annotations

metadata retrieved from object object is passed to ggplot2::ggtitle() as arguments for title, subtitle and caption. The specification for the title is passed as argument to annotations, and consists in the keyword title with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title, subtitle and caption. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

### Note

The plotting of absorbance is an exception to scale limits as the y-axis is not extended past 6 a.u. In the case of absorbance, values larger than 6 a.u. are rarely meaningful due to stray light during measurement. However, when transmittance values below the detection limit are rounded to zero, and later converted into absorbance, values Inf a.u. result, disrupting the plot. Scales are further expanded so as to make space for the annotations.

If idfactor = NULL, the default for single spectra, the name of the factor is retrieved from metadata or if no metadata found, the default "spct.idx" is tried. The default for multiple spectra is to create a factor named "spct.idx", but if a different name is passed, it will be used instead, possibly renaming a pre-existing one.

### See Also

[normalize](#), [filter\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: `autoplot.calibration_spct()`, `autoplot.cps_spct()`, `autoplot.object_spct()`, `autoplot.raw_spct()`, `autoplot.reflector_spct()`, `autoplot.response_spct()`, `autoplot.source_spct()`, `autoplot.waveband()`, `set_annotations_default()`

## Examples

```
# one spectrum
autoplot(yellow_gel.spct)
autoplot(yellow_gel.spct, geom = "spct")
autoplot(yellow_gel.spct, plot.qty = "transmittance")
autoplot(yellow_gel.spct, plot.qty = "absorbance")
autoplot(yellow_gel.spct, pc.out = TRUE)
autoplot(yellow_gel.spct, annotations = c("+", "wls"))

# spectra for two filters in long form
autoplot(two_filters.spct)
autoplot(two_filters.spct, idfactor = TRUE)
autoplot(two_filters.spct, idfactor = "Spectra")
autoplot(two_filters.spct, facets = TRUE)

# spectra for two filters as a collection
autoplot(two_filters.mspect)
autoplot(two_filters.mspect, idfactor = "Spectra")
autoplot(two_filters.mspect, facets = TRUE)
```

---

`autoplot.object_spct` *Plot one or more "object" spectra.*

---

## Description

These methods return a ggplot object with an annotated plot of an `object_spct` or an `object_spct` object. These objects contain spectral transmittance, reflectance and possibly absorbance data. As these quantities add up to one, only two are needed.

## Usage

```
## S3 method for class 'object_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default = list(UVC(), UVB(), UVA(),
    PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  plot.qty = "all",
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = NULL,
  span = NULL,
```

```

wls.target = "HM",
annotations = NULL,
geom = NULL,
time.format = "",
tz = "UTC",
stacked = plot.qty == "all",
text.size = 2.5,
chroma.type = "CMF",
idfactor = NULL,
facets = NULL,
plot.data = "as.is",
ylim = c(NA, NA),
object.label = deparse(substitute(object)),
na.rm = TRUE
)

## S3 method for class 'object_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  plot.qty = getOption("photobiology.filter.qty", default = "all"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  plot.data = "as.is",
  idfactor = TRUE,
  facets = plot.qty == "all",
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

```

### Arguments

object	an object_spct object
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting. If object is already normalized, the normalization is updated when a unit conversion applied.
plot.qty	character string, one of "all", "transmittance", "absorbance", "absorptance", or "reflectance".
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".

span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections <b>Plot Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
stacked	logical Whether to use <code>position_stack()</code> or <code>position_identity()</code> .
text.size	numeric size of text in the plot decorations.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spct contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.
ylim	numeric y axis limits,
object.label	character The name of the object being plotted.
na.rm	logical.

## Details

The plot object returned is a ggplot (an object of class "gg") and it can be added to or modified as any other ggplot. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', plotmath expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter annotations.

The generic of the [autoplot](#) method is defined in package 'ggplot2'. Package 'ggspectra' defines specializations for the different classes for storage of spectral data defined in package [photobiology](#).

For details about normalization and arguments to parameter `norm`, please, see [normalize](#). If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with 'photobiology' (<= 0.10.9) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with [fscale](#) methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named `"spct.idx"`, but if a different name is passed, it will be used instead.

### Value

A ggplot object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

### Plot Annotations

The recognized annotation names are: `"summaries"`, `"peaks"`, `"peak.labels"`, `"valleys"`, `"valley.labels"`, `"wls"`, `"wls.labels"`, `"colour.guide"`, `"color.guide"`, `"boxes"`, `"segments"`, `"labels"`. In addition, `"+"` is interpreted as a request to add to the already present default annotations, `"-"` as request to remove annotations and `"="` or missing `"+"` and `"-"` as a request to reset annotations to those requested. If used, `"+"`, `"-"` or `"="` must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass `""` or `c("=", "")` as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding `"peak.labels"` replaces `"peaks"` if present.

### Title Annotations

metadata retrieved from object `object` is passed to `ggplot2::ggtitle()` as arguments for `title`, `subtitle` and `caption`. The specification for the title is passed as argument to `annotations`, and consists in the keyword `title` with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to `title`, `subtitle` and `caption`. The recognized keywords are: `"objt"`, `"class"`, `"what"`, `"when"`, `"where"`, `"how"`, `"inst.name"`, `"inst.sn"`, `"comment"` and `"none"` are recognized as modifiers to `"title"`; `"none"` is a placeholder. Default is `"title:objt"` or no title depending on the context.

### Note

In the case of multiple spectra contained in the argument to `object` plotting is for `plot.qty = "all"` is always done using facets. Other plot quantities are handled by the methods for `filter_spct` and `reflector_spct` objects after on-the-fly conversion and the use of facets is possible but not the default.

If `idfactor = NULL`, the default for single spectra, the name of the factor is retrieved from metadata or if no metadata found, the default `"spct.idx"` is tried. The default for multiple spectra is to create a factor named `"spct.idx"`, but if a different name is passed, it will be used instead, possibly renaming a pre-existing one.

### See Also

[normalize](#), [object\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: `autoplot.calibration_spct()`, `autoplot.cps_spct()`, `autoplot.filter_spct()`, `autoplot.raw_spct()`, `autoplot.reflector_spct()`, `autoplot.response_spct()`, `autoplot.source_spct()`, `autoplot.waveband()`, `set_annotations_default()`

## Examples

```
low_res.spct <- thin_wl(Ler_leaf.spct,
                      max.wl.step = 20,
                      max.slope.delta = 0.01,
                      col.names = "Tfr")
autoplot(low_res.spct)
autoplot(low_res.spct, geom = "line")

two_leaves.mspct <-
  object_mspct(list("Arabidopsis leaf 1" = low_res.spct,
                  "Arabidopsis leaf 2" = low_res.spct))
autoplot(two_leaves.mspct, idfactor = "Spectra")
```

---

`autoplot.raw_spct`      *Plot one or more raw-detector-counts spectra.*

---

## Description

These methods construct a ggplot object with an annotated plot of a `raw_spct` or a `raw_mspct` object.

## Usage

```
## S3 method for class 'raw_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default = list(UVC(), UVB(), UVA(),
    PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  unit.out = "counts",
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = "mean",
  span = NULL,
  wls.target = "HM",
  annotations = NULL,
  geom = "line",
  time.format = "",
  tz = "UTC",
  text.size = 2.5,
  idfactor = NULL,
```

```

    facets = FALSE,
    plot.data = "as.is",
    ylim = c(NA, NA),
    object.label = deparse(substitute(object)),
    na.rm = TRUE
  )

## S3 method for class 'raw_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  unit.out = "counts",
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  idfactor = TRUE,
  facets = FALSE,
  plot.data = "as.is",
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

```

## Arguments

object	a raw_spct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting. If object is already normalized, the normalization is updated when a unit conversion applied.
unit.out	character IGNORED.
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections Plot <b>Annotations</b> and <b>Title Annotations</b> .

geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
text.size	numeric size of text in the plot decorations.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spct contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.
ylim	numeric y axis limits,
object.label	character The name of the object being plotted.
na.rm	logical.

## Details

The plot object returned is a ggplot (an object of class "gg") and it can be added to or modified as any other ggplot. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', plotmath expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter annotations.

The generic of the `autoplot` method is defined in package 'ggplot2'. Package 'ggspectra' defines specializations for the different classes for storage of spectral data defined in package [photobiology](#).

For details about normalization and arguments to parameter `norm`, please, see [normalize](#). If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with 'photobiology' (<= 0.10.9) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with `fscale` methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named "spct.idx", but if a different name is passed, it will be used instead.

## Value

A ggplot object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

## Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or c("=", "") as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

## Title Annotations

metadata retrieved from object object is passed to ggplot2::ggtitle() as arguments for title, subtitle and caption. The specification for the title is passed as argument to annotations, and consists in the keyword title with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title, subtitle and caption. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

## See Also

[normalize](#), [raw\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: [autoplot.calibration\\_spct\(\)](#), [autoplot.cps\\_spct\(\)](#), [autoplot.filter\\_spct\(\)](#), [autoplot.object\\_spct\(\)](#), [autoplot.reflector\\_spct\(\)](#), [autoplot.response\\_spct\(\)](#), [autoplot.source\\_spct\(\)](#), [autoplot.waveband\(\)](#), [set\\_annotations\\_default\(\)](#)

## Examples

```
low_res.raw_spct <- thin_wl(white_led.raw_spct,
                          max.wl.step = 20,
                          max.slope.delta = 0.05,
                          col.names = "counts_3")

autoplot(low_res.raw_spct)
autoplot(low_res.raw_spct, annotations = "")

two_leds.mspct <-
  raw_mspct(list("LED 1" = low_res.raw_spct,
                "LED 2" = low_res.raw_spct))

autoplot(two_leds.mspct)
autoplot(two_leds.mspct, facets = 1) # one column
```

---

`autoplot.reflector_spct`*Plot one or more reflector spectra.*

---

### Description

These methods return a ggplot object for an annotated plot from spectral data stored in a `reflector_spct` or a `reflector_mspct` object.

### Usage

```
## S3 method for class 'reflector_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default = list(UVC(), UVB(), UVA(),
    PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  plot.qty = getOption("photobiology.reflector.qty", default = "reflectance"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = NULL,
  span = NULL,
  wls.target = "HM",
  annotations = NULL,
  geom = "line",
  time.format = "",
  tz = "UTC",
  text.size = 2.5,
  chroma.type = "CMF",
  idfactor = NULL,
  facets = FALSE,
  plot.data = "as.is",
  ylim = c(NA, NA),
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

## S3 method for class 'reflector_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  plot.qty = getOption("photobiology.reflector.qty", default = "reflectance"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  plot.data = "as.is",
```

```

    idfactor = TRUE,
    facets = FALSE,
    object.label = deparse(substitute(object)),
    na.rm = TRUE
  )

```

## Arguments

object	a reflector_spct object or a reflector_mspct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting, If object is already normalized, the normalization is updated when a unit conversion applied.
plot.qty	character string (currently ignored).
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections <b>Plot Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <code>strptime</code> .
tz	character Time zone to use for title and/or subtitle.
text.size	numeric size of text in the plot decorations.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spct contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.

<code>ylim</code>	numeric y axis limits,
<code>object.label</code>	character The name of the object being plotted.
<code>na.rm</code>	logical.

## Details

The plot object returned is a `ggplot` (an object of class "gg") and it can be added to or modified as any other `ggplot`. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', *plotmath* expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter annotations.

The generic of the `autoplot` method is defined in package 'ggplot2'. Package 'ggspectra' defines specializations for the different classes for storage of spectral data defined in package `photobiology`.

For details about normalization and arguments to parameter `norm`, please, see `normalize`. If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with 'photobiology' (<= 0.10.9) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with `fscale` methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named "spt.idx", but if a different name is passed, it will be used instead.

## Value

A `ggplot` object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

## Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or `c("=", "")` as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

## Title Annotations

metadata retrieved from object `object` is passed to `ggplot2::ggtitle()` as arguments for `title`, `subtitle` and `caption`. The specification for the title is passed as argument to annotations,

and consists in the keyword `title` with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title, subtitle and caption. The recognized keywords are: `"objt"`, `"class"`, `"what"`, `"when"`, `"where"`, `"how"`, `"inst.name"`, `"inst.sn"`, `"comment"` and `"none"` are recognized as modifiers to `"title"`; `"none"` is a placeholder. Default is `"title:objt"` or no title depending on the context.

### See Also

[normalize](#), [reflector\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: [autoplot.calibration\\_spct\(\)](#), [autoplot.cps\\_spct\(\)](#), [autoplot.filter\\_spct\(\)](#), [autoplot.object\\_spct\(\)](#), [autoplot.raw\\_spct\(\)](#), [autoplot.response\\_spct\(\)](#), [autoplot.source\\_spct\(\)](#), [autoplot.waveband\(\)](#), [set\\_annotatations\\_default\(\)](#)

### Examples

```
autoplot(Ler_leaf_rflt.spct)
autoplot(Ler_leaf_rflt.spct, geom = "spct")
autoplot(Ler_leaf_rflt.spct, annotations = c("+", "valleys"))

two_leaves.mspct <-
  reflector_mspct(list("Arabidopsis leaf 1" = Ler_leaf_rflt.spct,
                     "Arabidopsis leaf 2" = Ler_leaf_rflt.spct / 2))
autoplot(two_leaves.mspct)
autoplot(two_leaves.mspct, idfactor = "Spectra")
autoplot(two_leaves.mspct, facets = 2)
```

---

```
autoplot.response_spct
```

*Plot one or more response spectra.*

---

### Description

These methods return a ggplot object with an annotated plot of the spectral data contained in a `response_spct` or a `response_mspct` object. Spectral responsitivity can be expressed either on an energy basis or a photon or quantum basis.

### Usage

```
## S3 method for class 'response_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default =
    list(photobiologyWavebands::UVC(), photobiologyWavebands::UVB(),
         photobiologyWavebands::UVA(), photobiologyWavebands::PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
```

```

    unit.out = getOption("photobiology.radiation.unit", default = "energy"),
    pc.out = getOption("ggspectra.pc.out", default = FALSE),
    label.qty = NULL,
    span = NULL,
    wls.target = "HM",
    annotations = NULL,
    geom = "line",
    time.format = "",
    tz = "UTC",
    text.size = 2.5,
    idfactor = NULL,
    facets = FALSE,
    plot.data = "as.is",
    ylim = c(NA, NA),
    object.label = deparse(substitute(object)),
    na.rm = TRUE
  )

## S3 method for class 'response_mspct'
autoplot(
  object,
  ...,
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  unit.out = getOption("photobiology.radiation.unit", default = "energy"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  plot.data = "as.is",
  facets = FALSE,
  idfactor = TRUE,
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

```

### Arguments

object	a response_spct object or a response_mspct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting. If object is already normalized, the normalization is updated when a unit conversion applied.
unit.out	character string indicating type of radiation units to use for plotting: "photon" or its synonym "quantum", or "energy".
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.

label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections Plot <b>Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spect" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
text.size	numeric size of text in the plot decorations.
idfactor	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
facets	logical or integer Indicating if facets are to be created for the levels of idfactor when spect contain multiple spectra in long form.
plot.data	character Data to plot. Default is "as.is" plotting one line per spectrum. When passing "mean", "median", "sum", "prod", "var", "sd", "se" as argument all the spectra must contain data at the same wavelength values.
ylim	numeric y axis limits,
object.label	character The name of the object being plotted.
na.rm	logical.

## Details

The plot object returned is a ggplot (an object of class "gg") and it can be added to or modified as any other ggplot. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In 'ggplot2', plotmath expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter annotations.

The generic of the [autoplot](#) method is defined in package 'ggplot2'. Package 'ggspectra' defines specializations for the different classes for storage of spectral data defined in package [photobiology](#).

For details about normalization and arguments to parameter `norm`, please, see [normalize](#). If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with 'photobiology' ( $\leq 0.10.9$ ) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with [fscale](#) methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does

not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named "spct.idx", but if a different name is passed, it will be used instead.

### Value

A ggplot object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

### Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or c("=", "") as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

### Title Annotations

metadata retrieved from object object is passed to ggplot2::ggtitle() as arguments for title, subtitle and caption. The specification for the title is passed as argument to annotations, and consists in the keyword title with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title, subtitle and caption. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

### See Also

[normalize](#), [response\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: [autoplot.calibration\\_spct\(\)](#), [autoplot.cps\\_spct\(\)](#), [autoplot.filter\\_spct\(\)](#), [autoplot.object\\_spct\(\)](#), [autoplot.raw\\_spct\(\)](#), [autoplot.reflector\\_spct\(\)](#), [autoplot.source\\_spct\(\)](#), [autoplot.waveband\(\)](#), [set\\_annotations\\_default\(\)](#)

### Examples

```
autoplot(photodiode.spct)
autoplot(photodiode.spct, geom = "spct")
autoplot(photodiode.spct, unit.out = "photon")
autoplot(photodiode.spct, annotations = "")

two_sensors.mspct <-
  response_mspct(list("Photodiode" = photodiode.spct,
                    "Coupled charge device" = ccd.spct)) |>
  normalize()
```

```

autoplot(two_sensors.mspct, unit.out = "photon")
autoplot(two_sensors.mspct, idfactor = "Spectra")
autoplot(two_sensors.mspct, facets = 2)
autoplot(two_sensors.mspct, geom = "spct")

```

---

autoplot.source\_spct *Plot one or more light-source spectra.*

---

### Description

These methods return a ggplot object with an annotated plot of the spectral data contained in a source\_spct or a source\_mspct object.

### Usage

```

## S3 method for class 'source_spct'
autoplot(
  object,
  ...,
  w.band = getOption("photobiology.plot.bands", default =
    list(photobiologyWavebands::UVC(), photobiologyWavebands::UVB(),
    photobiologyWavebands::UVA(), photobiologyWavebands::PhR())),
  range = getOption("ggspectra.wlrange", default = NULL),
  norm = NA,
  unit.out = getOption("photobiology.radiation.unit", default = "energy"),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  label.qty = NULL,
  span = NULL,
  wls.target = "HM",
  annotations = NULL,
  geom = "line",
  time.format = "",
  tz = "UTC",
  text.size = 2.5,
  chroma.type = "CMF",
  idfactor = NULL,
  facets = FALSE,
  plot.data = "as.is",
  ylim = c(NA, NA),
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)

## S3 method for class 'source_mspct'
autoplot(

```

```

    object,
    ...,
    range = getOption("ggspectra.wlrange", default = NULL),
    norm = NA,
    unit.out = getOption("photobiology.radiation.unit", default = "energy"),
    pc.out = getOption("ggspectra.pc.out", default = FALSE),
    idfactor = TRUE,
    facets = FALSE,
    plot.data = "as.is",
    object.label = deparse(substitute(object)),
    na.rm = TRUE
  )

```

### Arguments

object	a source_spct or a source_mspct object.
...	in the case of collections of spectra, additional arguments passed to the plot methods for individual spectra, otherwise currently ignored.
w.band	a single waveband object or a list of waveband objects.
range	an R object on which range() returns a vector of length 2, with minimum and maximum wavelengths (nm).
norm	numeric or character. Normalization to apply before plotting. If object is already normalized, the normalization is updated when a unit conversion applied.
unit.out	character string indicating type of radiation units to use for plotting: "photon" or its synonym "quantum", or "energy".
pc.out	logical, if TRUE use percent instead of fraction of one for normalized spectral data.
label.qty	character string giving the type of summary quantity to use for labels, one of "mean", "total", "contribution", and "relative".
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centred at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
annotations	a character vector. For details please see sections <b>Plot Annotations</b> and <b>Title Annotations</b> .
geom	character The name of a ggplot geometry, currently only "area", "spct" and "line". The default NULL selects between them based on stacked.
time.format	character Format as accepted by <a href="#">strptime</a> .
tz	character Time zone to use for title and/or subtitle.
text.size	numeric size of text in the plot decorations.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.

<code>idfactor</code>	character Name of an index factor used to identify each spectrum when multiple spectra are included in a plot. It is used as title to the guide in the plot and can include embedded spaces and new lines.
<code>facets</code>	logical or integer Indicating if facets are to be created for the levels of <code>idfactor</code> when <code>spect</code> contain multiple spectra in long form.
<code>plot.data</code>	character Data to plot. Default is <code>"as.is"</code> plotting one line per spectrum. When passing <code>"mean"</code> , <code>"median"</code> , <code>"sum"</code> , <code>"prod"</code> , <code>"var"</code> , <code>"sd"</code> , <code>"se"</code> as argument all the spectra must contain data at the same wavelength values.
<code>ylim</code>	numeric y axis limits,
<code>object.label</code>	character The name of the object being plotted.
<code>na.rm</code>	logical.

### Details

The plot object returned is a `ggplot` (an object of class `"gg"`) and it can be added to or modified as any other `ggplot`. The axis labels are encoded as *plotmath* expressions as they contain superscripts and special characters. In `'ggplot2'`, *plotmath* expressions do not obey theme settings related to text fonts, except for size.

Scale limits are expanded so as to make space for the annotations. If annotations are disabled, limits are not expanded unless `reserve.space` is passed to parameter annotations.

The generic of the `autoplot` method is defined in package `'ggplot2'`. Package `'ggspectra'` defines specializations for the different classes for storage of spectral data defined in package `photobiology`.

For details about normalization and arguments to parameter `norm`, please, see `normalize`. If `norm = NA`, the default, `normalize()` is not called. All other values passed as argument to `norm` result in a call to `normalize()` with this value as its argument. In the case of objects created with `'photobiology'` ( $\leq 0.10.9$ ) `norm = "undo"` is not supported. Be aware that calls to `normalize()` remove any scaling previously applied with `fscale` methods.

For multiple spectra in long form spectral objects, with `idfactor = NULL`, the default, the name of the factor is retrieved from metadata. If the character string passed as argument to `idfactor` does not match the one retrieved from the object, results in renaming of the pre-existing factor. The default for collections of spectra is to create a factor named `"spect.idx"`, but if a different name is passed, it will be used instead.

### Value

A `ggplot` object with a number of layers that depends on the data and annotations. The data member retains its original class and metadata attributes.

### Plot Annotations

The recognized annotation names are: `"summaries"`, `"peaks"`, `"peak.labels"`, `"valleys"`, `"valley.labels"`, `"wls"`, `"wls.labels"`, `"colour.guide"`, `"color.guide"`, `"boxes"`, `"segments"`, `"labels"`. In addition, `"+"` is interpreted as a request to add to the already present default annotations, `"-"` as request to remove annotations and `"="` or missing `"+"` and `"-"` as a request to reset annotations to those requested. If used, `"+"`, `"-"` or `"="` must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations

one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or c("", "") as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

### Title Annotations

metadata retrieved from object object is passed to `ggplot2::ggtitle()` as arguments for title, subtitle and caption. The specification for the title is passed as argument to annotations, and consists in the keyword title with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title, subtitle and caption. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

### See Also

[normalize](#), [source\\_spct](#), [waveband](#), [photobiologyWavebands-package](#) and [autoplot](#)

Other autoplot methods: [autoplot.calibration\\_spct\(\)](#), [autoplot.cps\\_spct\(\)](#), [autoplot.filter\\_spct\(\)](#), [autoplot.object\\_spct\(\)](#), [autoplot.raw\\_spct\(\)](#), [autoplot.reflector\\_spct\(\)](#), [autoplot.response\\_spct\(\)](#), [autoplot.waveband\(\)](#), [set\\_annotations\\_default\(\)](#)

### Examples

```
autoplot(sun.spct)
autoplot(sun.spct, geom = "spct")
autoplot(sun.spct, unit.out = "photon")
autoplot(normalize(sun.spct))
autoplot(normalize(sun.spct), pc.out = TRUE)

# multiple spectra in long form
autoplot(sun_evening.spct)
autoplot(sun_evening.spct, facets = 1) # one column
autoplot(sun_evening.spct, facets = 2) # two columns
autoplot(sun_evening.spct, plot.data = "mean")
autoplot(sun_evening.spct, idfactor = "Sequence")

# multiple spectra as a collection
autoplot(sun_evening.mspct)
autoplot(sun_evening.mspct, facets = 1) # one column
autoplot(sun_evening.mspct, facets = 2) # two columns
autoplot(sun_evening.mspct, plot.data = "mean")
autoplot(sun_evening.mspct, idfactor = "Time")
```

---

autoplot.waveband      *Create a complete ggplot for a waveband descriptor.*

---

## Description

Construct a ggplot object with an annotated plot of a waveband object.

## Usage

```
## S3 method for class 'waveband'
autoplot(
  object,
  ...,
  w.length = NULL,
  range = c(280, 800),
  fill = 0,
  span = NULL,
  wls.target = "HM",
  unit.in = getOption("photobiology.radiation.unit", default = "energy"),
  unit.out = unit.in,
  annotations = NULL,
  geom = "line",
  wb.trim = TRUE,
  norm = NA,
  text.size = 2.5,
  ylim = c(NA, NA),
  object.label = deparse(substitute(object)),
  na.rm = TRUE
)
```

## Arguments

object	a waveband object.
...	arguments passed along by name to autoplot.response_spct().
w.length	numeric vector of wavelengths (nm).
range	an R object on which range() returns a vector of length 2, with min and max wavelengths (nm).
fill	value to use as response for wavelengths outside the waveband range.
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centered at that element.
wls.target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.

<code>unit.in, unit.out</code>	the type of unit we assume as reference: "energy" or "photon" based for the waveband definition and the implicit matching response plotted.
<code>annotations</code>	a character vector. For details please see section Plot Annotations.
<code>geom</code>	character The name of a ggplot geometry, currently only "area", "spct" and "line".
<code>wb.trim</code>	logical. Passed to <code>trim_wl</code> . Relevant only when the waveband extends partly outside range.
<code>norm</code>	numeric or character Normalization wavelength (nm) or character string "max" or other criterion for normalization.
<code>text.size</code>	numeric size of text in the plot decorations.
<code>ylim</code>	numeric y axis limits,
<code>object.label</code>	character The name of the object being plotted.
<code>na.rm</code>	logical.

### Details

A `response_spct` object is created based on the waveband object. A waveband object can describe either a simple wavelength range or a (biological) spectral weighting function (BSWF). See [autoplot.response\\_spct](#) for additional details.

Effectiveness spectra are plotted expressing the spectral effectiveness either as  $1\text{mol}^{-1}\text{nm}$  photons of  $1\text{J}^{-1}\text{nm}$  which can be selected through formal argument `unit.out`. The value of `unit.in` has no effect on the result when using BSWFs, as BSWFs are defined based on a certain base of expression, which is enforced. In contrast, for wavebands which only define a wavelength range, changing the assumed reference irradiance units, changes the responsivity according to Planck's law.

Unused arguments are passed along, which means that other plot aspects can be controlled by providing arguments for the plot method of the `response_spct` class.

### Value

a ggplot object.

### Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or `c("=", "")` as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

**See Also**

[autoplot.response\\_spct](#), [waveband](#).

Other autoplot methods: [autoplot.calibration\\_spct\(\)](#), [autoplot.cps\\_spct\(\)](#), [autoplot.filter\\_spct\(\)](#), [autoplot.object\\_spct\(\)](#), [autoplot.raw\\_spct\(\)](#), [autoplot.reflector\\_spct\(\)](#), [autoplot.response\\_spct\(\)](#), [autoplot.source\\_spct\(\)](#), [set\\_annotatations\\_default\(\)](#)

**Examples**

```
autoplot(waveband(c(400, 500)))
autoplot(waveband(c(400, 500)), geom = "spct")
```

---

autotitle

*Add title, subtitle and caption to a spectral plot*

---

**Description**

Add a title, subtitle and caption to a spectral plot based on automatically extracted metadata from an spectral object.

**Usage**

```
autotitle(
  object,
  object.label = deparse(substitute(object)),
  annotations = "title",
  time.format = NULL,
  tz = "",
  default.title = "title:objt"
)
```

```
ggtitle_spct(
  object,
  object.label = deparse(substitute(object)),
  annotations = "title",
  time.format = NULL,
  tz = "",
  default.title = "title:objt"
)
```

**Arguments**

<code>object</code>	generic_spct or generic_mspct The spectral object plotted.
<code>object.label</code>	character The name of the object being plotted.
<code>annotations</code>	character vector Annotations as described for <code>plot()</code> methods, values unrelated to title are ignored.

`time.format` character Format as accepted by `strptime`.  
`tz` character time zone used in labels.  
`default.title` character vector The default used for annotations = "title".

**Value**

The return value of `ggplot2::labs()`.

**Title Annotations**

metadata retrieved from object `object` is passed to `ggplot2::ggtitle()` as arguments for `title`, `subtitle` and `caption`. The specification for the title is passed as argument to `annotations`, and consists in the keyword `title` with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to `title`, `subtitle` and `caption`. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

**Note**

Method renamed as `autotitle()` to better reflect its function; `ggtitle_spct()` is deprecated but will remain available for backwards compatibility.

**Examples**

```

p <- ggplot(sun.spct) +
  geom_line()

p + autotitle(sun.spct)
p + autotitle(sun.spct, object.label = "The terrestrial solar spectrum")
p + autotitle(sun.spct, annotations = "title:objt:class")
p + autotitle(sun.spct, annotations = "title:where:when:how")

p <- ggplot(sun_evening.spct) +
  aes(linetype = spct.idx) +
  geom_line()

p + autotitle(sun_evening.spct, annotations = "title:objt:class")
p + autotitle(sun_evening.spct, annotations = "title:where:when:how")
p + autotitle(sun_evening.spct, annotations = "title:none:none:how")

p <- ggplot(sun_evening.mspct) +
  aes(linetype = spct.idx) +
  geom_line()

p + autotitle(sun_evening.mspct, annotations = "title:objt:class")
  
```

---

axis_labels_uk	<i>Default text for axis labels</i>
----------------	-------------------------------------

---

### Description

Texts used by default for axis labels in plots are recalled from character vectors returned by these functions. The aim is that their default values can be easily changed or translated to other languages. They contain only the text part, but not symbols or units of expression.

### Usage

```
axis_labels_uk(append = "", sep = "")  
  
axis_labels_uk_comma()  
  
axis_labels_none()  
  
axis_labels()
```

### Arguments

append	character	The string to be appended to each label,
sep	character	Passed to function paste as argument for parameter sep.

### Details

By default `axis_labels()` contains a copy of `axis_labels_uk_comma()`. By assigning to this name a user function that returns a named character vector using the same names as those returned by these functions, it is possible to temporarily change the default texts.

Currently only UK English label texts are predefined and `axis_labels()` is a synonym of `axis_labels_uk()`.

### Value

A character vector

### Examples

```
axis_labels()[["w.length"]] # ending in a comma  
axis_labels_uk()[["w.length"]] # no comma  
axis_labels_none()[["w.length"]] # empty label
```

---

A_label	<i>Absorbance axis labels</i>
---------	-------------------------------

---

### Description

Generate cps axis labels in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

### Usage

```
A_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  Tfr.type
)
```

```
A_internal_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
A_total_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

### Arguments

unit.exponent	integer
format	character string, "R", "R.expresion", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).

`axis.symbols` logical If TRUE symbols of the quantities are added to the name. Supported only by `format = "R.expression"`.

`Tfr.type` character, either "total" or "internal".

**Value**

a character string or an R expression.

**Note**

Default for `label.text` depends on the value passed as argument to `Tfr.type`.

**Examples**

```
A_label(Tfr.type = "internal")
A_label(Tfr.type = "total")
A_label(Tfr.type = "total", axis.symbols = FALSE)

A_internal_label()
A_internal_label(format = "R.expression", axis.symbols = FALSE)
A_internal_label(-3)
A_internal_label(format = "R.expression")
A_internal_label(format = "LaTeX")
A_internal_label(-3, format = "LaTeX")

A_total_label()
A_total_label(format = "R.expression", axis.symbols = FALSE)
A_total_label(-3)
A_total_label(format = "R.expression")
A_total_label(format = "LaTeX")
A_total_label(-3, format = "LaTeX")
```

---

`black_or_white`      *Chose black vs. white color based on weighted mean of RGB channels*

---

**Description**

Chose black or white color based on a color to be used as background. Usefull when using `geom_text` on top of tiles or bars, or `geom_label` with a variable fill.

**Usage**

```
black_or_white(colors, threshold = 0.45)
```

**Arguments**

colors            character A vector of color definitions.  
 threshold        numeric in range 0 to 1.

**Examples**

```
black_or_white("red")
black_or_white(colors()[1:10])
```

---

color_chart	<i>Create a color checker chart</i>
-------------	-------------------------------------

---

**Description**

Color-checker-chart ggplot labelled with color names or with indexes of the colors in the vector passed as first argument.

**Usage**

```
color_chart(
  colors = grDevices::colors(),
  ncol = NULL,
  use.names = NULL,
  text.size = 2,
  text.color = NULL,
  grid.color = "white"
)
```

**Arguments**

colors            character A vector of color definitions.  
 ncol             integer Number of column in the checker grid.  
 use.names        logical Force use of names or indexes.  
 text.size        numeric Size of the text labels drawn on each color tile.  
 text.color       character Color definition, used for text on tiles.  
 grid.color       character Color definition, used for grid lines between tiles.

**Note**

Default text.color uses black\_or\_white() to ensure enough contrast. Default for use.names depends on number of columns in the grid, indexes are used when columns are seven or more.

**Examples**

```
color_chart()
color_chart(grep("dark", colors()), value = TRUE, text.size = 3.5)
```

---

counts_label	<i>Raw-counts axis labels</i>
--------------	-------------------------------

---

### Description

Generate axis labels in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

### Usage

```
counts_label(  
  unit.exponent = 3,  
  format = getOption("photobiology.math", default = "R.expression"),  
  label.text = axis_labels()[["counts"]],  
  scaled = FALSE,  
  normalized = FALSE,  
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)  
)
```

### Arguments

unit.exponent	integer
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".

### Value

a character string or an R expression.

### Examples

```
counts_label()  
counts_label("R.expression")  
counts_label("LaTeX")
```

---

cps_label	<i>Counts-per-second axis labels</i>
-----------	--------------------------------------

---

### Description

Generate pixel response rate axis labels in cps units. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

### Usage

```
cps_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["cps"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

### Arguments

unit.exponent	integer
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".

### Value

a character string or an R expression.

### Examples

```
cps_label()
cps_label(3)
cps_label(format = "R.expression")
cps_label(format = "R.character")
cps_label(format = "LaTeX")
cps_label(3, format = "LaTeX")
```

---

exponent2prefix	<i>SI unit prefixes</i>
-----------------	-------------------------

---

### Description

Convert SI unit prefixes into exponents of ten of multipliers and vice-versa.

### Usage

```
exponent2prefix(
  exponent,
  char.set = getOption("photobiology.fancy.chars", default = "utf8")
)

exponent2factor(exponent = 0, if.zero.exponent = "1")

exponent2prefix_name(exponent)

prefix_name2exponent(name)

prefix2exponent(
  prefix,
  char.set = getOption("photobiology.fancy.chars", default = "utf8")
)

has_SI_prefix(exponent)

nearest_SI_exponent(exponent)
```

### Arguments

exponent	numeric	The power of 10 of the unit multiplier.
char.set	character	How to encode Greek letters and other fancy characters in prefixes: "utf8", "ascii", "LaTeX". The difference between "utf8" and "ascii" is that the first uses UTF8 character "micro" (similar to Greek mu) and the second uses "u".
if.zero.exponent	character	string to return when exponent is equal to zero.
name	character	Long SI name of multiplier.
prefix	character	Unit prefix used for multiplier.

### Note

To change the default `char.set`, set R option `"photobiology.fancy.chars"`. Implementation is based on a table of data and extensible to any alphabet supported by R character objects by expanding the table.

**Examples**

```
exponent2prefix(3)
exponent2prefix(0)
exponent2prefix(-6)
```

```
exponent2factor(3)
exponent2factor(0)
exponent2factor(0, NULL)
exponent2factor(0, "")
exponent2factor(-6)
```

---

geom\_spct

*Spectral data plots.*


---

**Description**

For each continuous x value, geom\_spct displays a y interval. geom\_spct is a special case of geom\_area, where the minimum of the range is fixed to 0, but stacking is not enabled.

**Usage**

```
geom_spct(
  mapping = NULL,
  data = NULL,
  stat = "identity",
  position = "identity",
  ...,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)
```

**Arguments**

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A data frame. If specified, overrides the default data frame defined at the top level of the plot.
stat	The statistical transformation to use on the data for this layer, as a string.
position	Position adjustment, either as a string, or the result of a call to a position adjustment function.
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.

<code>na.rm</code>	If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Details

An spectrum plot is the analog of a line plot (see [geom\\_path](#)), and can be used to show how y varies over the range of x. The difference is that the area under the line is filled.

### Aesthetics

See [geom\\_ribbon](#)

### See Also

[geom\\_ribbon](#) for stacked areas, [geom\\_path](#) for lines (lines), [geom\\_point](#) for scatter plots.

### Examples

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) + geom_spct()
```

---

ggplot

*Create a new ggplot plot from spectral data.*

---

### Description

`ggplot()` initializes a `ggplot` object. It can be used to declare the input spectral object for a graphic and to optionally specify the set of plot aesthetics intended to be common throughout all subsequent layers unless specifically overridden.

### Usage

```
## S3 method for class 'source_spct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  unit.out = getOption("photobiology.radiation.unit", default = "energy"),
  environment = parent.frame()
)
```

```
## S3 method for class 'response_spct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  unit.out = getOption("photobiology.radiation.unit", default = "energy"),
  environment = parent.frame()
)

## S3 method for class 'filter_spct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  plot.qty = getOption("photobiology.filter.qty", default = "transmittance"),
  environment = parent.frame()
)

## S3 method for class 'reflector_spct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  plot.qty = NULL,
  environment = parent.frame()
)

## S3 method for class 'cps_spct'
ggplot(data, mapping = NULL, ..., range = NULL, environment = parent.frame())

## S3 method for class 'calibration_spct'
ggplot(data, mapping = NULL, ..., range = NULL, environment = parent.frame())

## S3 method for class 'raw_spct'
ggplot(data, mapping = NULL, ..., range = NULL, environment = parent.frame())

## S3 method for class 'object_spct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  plot.qty = getOption("photobiology.object.qty", default = "all"),
  environment = parent.frame()
)
```

```
)

## S3 method for class 'generic_spct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  spct_class,
  environment = parent.frame()
)

## S3 method for class 'generic_mspct'
ggplot(data, mapping = NULL, ..., range = NULL, environment = parent.frame())

## S3 method for class 'filter_mspct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  plot.qty = getOption("photobiology.filter.qty", default = "transmittance"),
  environment = parent.frame()
)

## S3 method for class 'source_mspct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  unit.out = getOption("photobiology.radiation.unit", default = "energy"),
  environment = parent.frame()
)

## S3 method for class 'object_mspct'
ggplot(
  data,
  mapping = NULL,
  ...,
  range = NULL,
  plot.qty = getOption("photobiology.object.qty", default = ifelse(length(data) > 1L,
    "as.is", "all")),
  environment = parent.frame()
)
```

**Arguments**

<code>data</code>	Default spectrum dataset to use for plot. If not a spectrum, the methods used will be those defined in package <code>ggplot2</code> . See <a href="#">ggplot</a> . If not specified, must be supplied in each layer added to the plot.
<code>mapping</code>	Default list of aesthetic mappings to use for plot. If not specified, in the case of spectral objects, a default mapping will be used.
<code>...</code>	Other arguments passed on to methods.
<code>range</code>	an R object on which <code>range()</code> returns a vector of length 2, with min and max wavelengths (nm).
<code>unit.out</code>	character string indicating type of units to use for plotting spectral irradiance or spectral response, "photon" or "energy".
<code>environment</code>	If a variable defined in the aesthetic mapping is not found in the data, <code>ggplot</code> will look for it in this environment. It defaults to using the environment in which <code>ggplot()</code> is called.
<code>plot.qty</code>	character string One of "transmittance", "absorptance" or "absorbance" for <code>filter_spct</code> objects, and in addition to these "reflectance", "all" or "as.is" for <code>object_spct</code> objects.
<code>spct_class</code>	character Class into which a <code>generic_spct</code> object will be converted before plotting. The column names in data should match those expected by the class constructor (see <a href="#">setGenericSpct</a> ); other arguments should be passed by name).

**Details**

`ggplot()` is typically used to construct a plot incrementally, using the `+` operator to add layers to the existing `ggplot` object. This is advantageous in that the code is explicit about which layers are added and the order in which they are added. For complex graphics with multiple layers, initialization with `ggplot` is recommended.

We show seven common ways to invoke `ggplot` for spectra and collections of spectra:

- `ggplot(spct)`
- `ggplot(spct, unit.out = <unit.to.use>)`
- `ggplot(spct, plot.qty = <quantity.to.plot>)`
- `ggplot(spct, range = <wavelength.range>)`
- `ggplot(spct) + aes(<other aesthetics>)`
- `ggplot(spct, aes(x, y, <other aesthetics>))`
- `ggplot(spct, aes())`

The first method is recommended if all layers use the same data and the same set of automatic default `x` and `y` aesthetics. The second, third and fourth use automatic default `x` and `y` aesthetics but first transform or trim the spectral data to be plotted. The fifth uses automatic default `x` and `y` aesthetics and adds mappings for other aesthetics. These patterns can be combined as needed. The sixth disables the use of a default automatic mapping, while the seventh delays the mapping of aesthetics and can be convenient when using different mappings for different geoms.

## Object spectra

In the case of class `object_spct`, the arguments `"all"` and `"as.is"` if passed to `plot.qty`, indicate in the first case that the data are to be converted into long form, to allow stacking, while in the second case data is copied unchanged to the plot object. `"reflectance"` passed to `plot.qty` converts data into a `reflector_spct` object and `"absorbance"`, `"absorptance"` and `"reflectance"`, convert data into a `filter_spct`.

## Collections of spectra

The method for collections of spectra accepts arguments for the same parameters as the corresponding methods for single spectra. Heterogeneous generic collections of spectra are not supported. When plotting collections of spectra the factor `spect.idx` contains as levels the names of the individual members of the collection, and can be mapped to aesthetics or used for faceting.

## Note

Current implementation does not merge the default mapping with user supplied mapping. If user supplies a mapping, it is used as is, and variables should be present in the spectral object. In contrast, when using the default mapping, unit or quantity conversions are done on the fly when needed. To add to the default mapping, `aes()` can be used by itself to compose the `ggplot`. In all cases, except when an `object_spct` is converted into long form, the data member of the returned plot object retains its class and attributes.

`plot.qty` is ignored for reflectors.

## Examples

```
ggplot(sun.spct) + geom_line()
ggplot(sun.spct, unit.out = "photon") + geom_line()

ggplot(yellow_gel.spct) + geom_line()
ggplot(yellow_gel.spct, plot.qty = "absorbance") + geom_line()

ggplot(Ler_leaf.spct) + facet_grid(~variable) + geom_line()
ggplot(Ler_leaf.spct) + aes(linetype = variable) + geom_line()
```

---

multipliers\_label

*Calibration multipliers axis labels*

---

## Description

Calibration multipliers axis labels. Output can be selected as character, expression (R default devices) or LaTeX (for `tikz` device).

**Usage**

```

multipliers_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["e.mult"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

```

**Arguments**

unit.exponent	integer
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".

**Value**

a character string or an R expression.

**Examples**

```

multipliers_label()
multipliers_label(3)
multipliers_label(format = "R.expression")
multipliers_label(format = "R.character")
multipliers_label(format = "LaTeX")
multipliers_label(3, format = "LaTeX")

```

---

multiplot

*Multiple plot function*


---

**Description**

Grid based; allows multiple plots arranged in a matrix and printed to any R device. ggplot objects can be passed in ..., or to plotlist (as a list of ggplot objects)

**Usage**

```
multiplot(
  ...,
  plotlist = NULL,
  ncol = 1,
  cols = ncol,
  layout = NULL,
  title = "",
  title.position = "left",
  title.fontsize = 12,
  title.fontfamily = "sans",
  title.fontface = "bold",
  title.colour = "black"
)
```

**Arguments**

...	one or more ggplot objects.
plotlist	list of ggplot objects.
ncol, cols	numerical Number of columns in layout.
layout	A numeric matrix specifying the layout. If present, 'cols' is ignored.
title	character vector Title of the composite plot.
title.position	numeric or character, the horizontal position of the title.
title.fontsize	numeric
title.fontfamily	character e.g. "sans", "serif", "mono".
title.fontface	character e.g. "plain", "bold", "italic", "bold.italic".
title.colour	character e.g. "black", "red".

**Details**

ggplot objects can be passed in ..., or to plotlist (as a list of ggplot objects) If the layout is something like `matrix(c(1,2,3,3), nrow=2, byrow=TRUE)`, then plot 1 will go in the upper left, 2 will go in the upper right, and 3 will go all the way across the bottom.

**Note**

Modified from example by Winston Chang found in the Cookbook for R Licenced under CC BY-SA

**References**

<http://www.cookbook-r.com/>

## Examples

```
multiplot(plot(sun.spct), plot(yellow_gel.spct), ncol = 1)
multiplot(plot(sun.spct), plot(yellow_gel.spct), ncol = 1,
          title = "The sun and a yellow filter")
```

---

plot.generic\_spct      *Deprecated plot methods*

---

## Description

These plot() methods return a ggplot object with an annotated plot of an object of a class derived from generic\_spct, of a class derived from generic\_mspct or of an object of class waveband for which an autoplot() method exists. They are implemented as wrappers of autoplot(). The generic for plot() is defined by base R and specializations for objects of diverse classes are provided various packages and R itself. The generic for autoplot() is defined by package 'ggplot2'.

## Usage

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

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

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

## Arguments

x                    An R object derived from class generic\_spct or derived from class generic\_mspct.  
...                    Named arguments passed to autoplot() methods.

## Value

a ggplot object.

## Deprecation warning!

These plot() specializations are provided for backwards compatibility, but all new or updated code should call autoplot() instead of plot() on objects of spectral and waveband classes defined in package 'photobiology'.

These methods add support for plot() specializations as these specializations were provided by package 'ggspectra' years ago, before 'ggplot2' had an autoplot() generic. As these methods return ggplots autoplot is a more suitable name for them.

**See Also**

[autoplot.calibration\\_spct](#), [autoplot.cps\\_spct](#), [autoplot.filter\\_spct](#), [autoplot.raw\\_spct](#), [autoplot.response\\_spct](#), [autoplot.source\\_spct](#) and [autoplot.waveband](#).

**Examples**

```
plot(sun.spct) # deprecated syntax, to be avoided
autoplot(sun.spct) # current syntax, to be used
```

---

Rfr\_label

*Reflectance axis labels*


---

**Description**

Generate spectral reflectance labels in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

**Usage**

```
Rfr_label(
  unit.exponent = ifelse(pc.out, -2, 0),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  Rfr.type
)

Rfr_specular_label(
  unit.exponent = ifelse(pc.out, -2, 0),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE)
)
```

**Arguments**

```
unit.exponent  integer
format         character string, "R", "R.expression", "R.character", or "LaTeX".
label.text     character Textual portion of the labels.
```

scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in nanometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
pc.out	logical, if TRUE use percent as default instead of fraction of one.
Rfr.type	character, either "total" or "specular".

**Value**

a character string or an R expression.

**Note**

Default for label.text depends on the value passed as argument to Rfr.type.

**Examples**

```
Rfr_label(Rfr.type = "specular")
Rfr_label(Rfr.type = "total")

Rfr_specular_label()
Rfr_specular_label(axis.symbols = FALSE)
Rfr_specular_label(-2)
Rfr_specular_label(-3)
Rfr_specular_label(format = "R.expression")
Rfr_specular_label(format = "LaTeX")
Rfr_specular_label(-3, format = "LaTeX")
```

---

s.e.irrad_label	<i>Spectral irradiance axis labels</i>
-----------------	--

---

**Description**

Generate axis labels for spectral irradiance in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

**Usage**

```
s.e.irrad_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.e.irrad"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
```

```

)

s.q.irrad_label(
  unit.exponent = ifelse(normalized, 0, -6),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.q.irrad"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

```

### Arguments

unit.exponent	integer.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".

### Value

a character string or an R expression.

### Examples

```

str(s.e.irrad_label())
str(s.e.irrad_label(axis.symbols = FALSE))
str(s.e.irrad_label(format = "R.expression"))
str(s.e.irrad_label(format = "LaTeX"))

```

```

str(s.q.irrad_label())
str(s.q.irrad_label(axis.symbols = FALSE))
str(s.q.irrad_label(format = "R.expression"))
str(s.q.irrad_label(format = "LaTeX"))

```

---

s.e.response_label	<i>spectral response and action axis labels</i>
--------------------	---

---

### Description

Generate axis labels for response or action spectra in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

**Usage**

```
s.e.response_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.e.response"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
s.q.response_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.q.response"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
s.e.action_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.e.action"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
s.q.action_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.q.action"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

**Arguments**

unit.exponent	integer
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".

**Value**

a character string or an R expression.

**Examples**

```
s.e.response_label()
s.e.response_label(format = "R.expression")
s.e.response_label(format = "R.character")
s.e.response_label(format = "LaTeX")
s.e.response_label(unit.exponent = 3, format = "R.character")
s.q.response_label(format = "R.character")
s.e.action_label(format = "R.character")
s.q.action_label(format = "R.character")
s.e.response_label(scaled = TRUE)
s.e.response_label(scaled = TRUE, format = "R.character")
s.e.response_label(scaled = TRUE, format = "LaTeX")
s.e.response_label(normalized = 300)
s.e.response_label(normalized = 300, format = "R.character")
s.e.response_label(normalized = 300, format = "LaTeX")
s.q.response_label(scaled = TRUE)
s.q.response_label(scaled = TRUE, format = "R.character")
s.q.response_label(scaled = TRUE, format = "LaTeX")
s.q.response_label(normalized = 300)
s.q.response_label(normalized = 300, format = "R.character")
s.q.response_label(normalized = 300, format = "LaTeX")
```

---

scale\_x\_energy\_eV\_continuous

*Energy per photon x-scale*

---

**Description**

Scale x continuous with defaults suitable for wavelengths expressed as energy per photon [eV] or [J].

**Usage**

```
scale_x_energy_eV_continuous(
  unit.exponent = 0,
  name = w_energy_eV_label(unit.exponent = unit.exponent, label.text = label.text,
    axis.symbols = axis.symbols),
  breaks = scales::pretty_breaks(n = 7),
  labels = SI_pl_format(exponent = unit.exponent),
  label.text = axis_labels()[["energy"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)
```

```

scale_x_energy_J_continuous(
  unit.exponent = -18,
  name = w_energy_J_label(unit.exponent = unit.exponent, label.text = label.text,
    axis.symbols = axis.symbols),
  breaks = scales::pretty_breaks(n = 7),
  labels = SI_pl_format(exponent = unit.exponent),
  label.text = axis_labels()[["energy"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

```

### Arguments

<code>unit.exponent</code>	integer
<code>name</code>	The name of the scale, used for the axis-label.
<code>breaks</code>	The positions of ticks or a function to generate them.
<code>labels</code>	The tick labels or a function to generate them from the tick positions.
<code>label.text</code>	character Textual portion of the labels.
<code>axis.symbols</code>	logical If TRUE symbols of the quantities are added to the name. Supported only by <code>format = "R.expression"</code> .
<code>...</code>	other named arguments passed to <code>scale_y_continuous</code>

### Details

This scale automates the generation of axis labels when the variable mapped to the  $x$  aesthetic contains numeric values for wavelengths expressed as energy per photon. This is **not** how spectral data are stored in all the packages of the R for Photobiology suite and can be used in plots built with `ggplot2()` with explicit mapping using a conversion function. If desired, a secondary axis can be added manually as described in [sec\\_axis](#).

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

### Examples

```

ggplot(sun.spct, aes(x = wl2energy(w.length, unit = "joule"), y = s.e.irrad)) +
  geom_line() +
  scale_x_energy_J_continuous()

ggplot(sun.spct, aes(x = wl2energy(w.length, unit = "joule"), y = s.e.irrad)) +
  geom_line() +
  scale_x_energy_J_continuous(unit.exponent = -19)

ggplot(sun.spct, aes(x = wl2energy(w.length, unit = "eV"), y = s.e.irrad)) +
  geom_line() +
  scale_x_energy_eV_continuous()

```

```
ggplot(sun.spct, aes(x = wl2energy(w.length, unit = "eV"), y = s.e.irrad)) +
  geom_line() +
  scale_x_energy_eV_continuous(unit.exponent = -3)
```

---

scale\_x\_frequency\_continuous

*Frequency x-scale*

---

### Description

Scale x continuous with defaults suitable for wavelengths expressed as frequencies [Hz].

### Usage

```
scale_x_frequency_continuous(
  unit.exponent = 12,
  name = w_frequency_label(unit.exponent = unit.exponent, label.text = label.text,
    axis.symbols = axis.symbols),
  breaks = scales::pretty_breaks(n = 7),
  labels = SI_pl_format(exponent = unit.exponent),
  label.text = axis_labels()[["freq"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)
```

### Arguments

unit.exponent	integer
name	The name of the scale, used for the axis-label.
breaks	The positions of ticks or a function to generate them.
labels	The tick labels or a function to generate them from the tick positions.
label.text	character Textual portion of the labels.
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
...	other named arguments passed to scale_y_continuous

### Details

This scale automates the generation of axis labels when the variable mapped to the  $x$  aesthetic contains numeric values for wavelengths expressed as frequency. This is **not** how spectral data are stored in the packages of the R for Photobiology suite and can be only used in plots built with `ggplot2()` with explicit mapping using a conversion function. If desired, a secondary axis can be added manually as described in [sec\\_axis](#).

**Note**

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

**Examples**

```
ggplot(sun.spct, aes(x = wl2frequency(w.length), y = s.e.irrad)) +
  geom_line() +
  scale_x_frequency_continuous()
```

```
ggplot(sun.spct, aes(x = wl2frequency(w.length), y = s.e.irrad)) +
  geom_line() +
  scale_x_frequency_continuous(14)
```

---

scale\_x\_wavenumber\_continuous  
*Wavenumber x-scale*

---

**Description**

Scale x continuous with defaults suitable for wavelengths expressed as wavenumbers [ $m^{-2}$ ].

**Usage**

```
scale_x_wavenumber_continuous(
  unit.exponent = -6,
  name = w_number_label(unit.exponent = unit.exponent, label.text = label.text,
    axis.symbols = axis.symbols),
  breaks = scales::pretty_breaks(n = 7),
  labels = SI_pl_format(exponent = -unit.exponent),
  label.text = axis_labels()[["w.number"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)
```

**Arguments**

unit.exponent	integer
name	The name of the scale, used for the axis-label.
breaks	The positions of ticks or a function to generate them.
labels	The tick labels or a function to generate them from the tick positions.
label.text	character Textual portion of the labels.
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
...	other named arguments passed to scale_y_continuous

**Details**

This scale automates the generation of axis labels when the variable mapped to the  $x$  aesthetic contains numeric values for wavelengths expressed wavenumbers. This is **not** how spectral data are stored in all the packages of the R for Photobiology suite and can be used in plots built with `ggplot2()` with explicit mapping using a conversion function. If desired, a secondary axis can be added manually as described in [sec\\_axis](#).

**Note**

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

**Examples**

```
ggplot(sun.spct, aes(x = wl2wavenumber(w.length), y = s.e.irrad)) +
  geom_line() +
  scale_x_wavenumber_continuous()

ggplot(sun.spct, aes(x = wl2wavenumber(w.length), y = s.e.irrad)) +
  geom_line() +
  scale_x_wavenumber_continuous(unit.exponent = -5)
```

---

scale\_x\_wl\_continuous *Wavelength x-scale*

---

**Description**

Scale  $x$  continuous with defaults suitable for wavelengths in nanometres.

**Usage**

```
scale_x_wl_continuous(
  unit.exponent = -9,
  name = w_length_label(unit.exponent = unit.exponent, label.text = label.text,
    axis.symbols = axis.symbols),
  breaks = scales::pretty_breaks(n = 7),
  labels = SI_pl_format(exponent = unit.exponent + 9),
  label.text = axis_labels()[["w.length"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)
```

**Arguments**

<code>unit.exponent</code>	integer
<code>name</code>	The name of the scale, used for the axis-label.
<code>breaks</code>	The positions of ticks or a function to generate them.

labels	The tick labels or a function to generate them from the tick positions.
label.text	character Textual portion of the labels.
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
...	other named arguments passed to scale_y_continuous

### Details

This scale automates the generation of axis labels when the variable mapped to the  $x$  aesthetic contains numeric values for wavelengths expressed in nanometres. This is how spectral data are stored in all the packages of the R for Photobiology suite, including the the expected data by the autoplot() methods defined in 'ggspectra'.

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

### Examples

```
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous()

ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(unit.exponent = -6)

ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(label.text = "Longitud de onda,")

autoplot(sun.spct) +
  scale_x_wl_continuous(label.text = "Longitud de onda,",
                        unit.exponent = -6)
```

---

scale\_y\_Afr\_continuous

*Absorptance y-scale*

---

### Description

Scale y continuous with defaults suitable for spectral absorbance.

**Usage**

```

scale_y_Afr_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Afr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = round(normalized, 1), axis.symbols =
    axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.Afr"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  ...
)

```

**Arguments**

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
limits	One of NULL for default based on data range, a numeric vector of length two (NA allowed) or a function that accepts the data-based limits as argument and returns new limits.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
pc.out	logical, if TRUE use percent as default instead of fraction of one.
...	other named arguments passed to scale_y_continuous

**Note**

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

**Examples**

```

Afr_as_default()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Afr_continuous() +
  scale_x_wl_continuous()

```

```

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Afr_continuous(unit.exponent = -2) +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Afr_continuous(unit.exponent = -3) +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Afr_continuous(axis.symbols = FALSE) +
  scale_x_wl_continuous(axis.symbols = FALSE)

unset_filter_qty_default()

```

---

scale\_y\_A\_continuous *Absorbance y-scale*

---

## Description

Scale y continuous with defaults suitable for spectral absorbance.

## Usage

```

scale_y_A_continuous(
  unit.exponent = 0,
  name = A_label(unit.exponent = unit.exponent, format = format, label.text = label.text,
    scaled = scaled, normalized = ifelse(is.numeric(normalized), round(normalized, 1),
    unique(normalized))), axis.symbols = axis.symbols, Tfr.type = Tfr.type),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  Tfr.type,
  ...
)

scale_y_A_internal_continuous(
  unit.exponent = 0,
  name = A_label(unit.exponent = unit.exponent, format = format, label.text = label.text,
    scaled = scaled, normalized = ifelse(is.numeric(normalized), round(normalized, 1),
    unique(normalized))), axis.symbols = axis.symbols, Tfr.type = "internal"),

```

```

    labels = SI_pl_format(exponent = unit.exponent),
    format = getOption("photobiology.math", default = "R.expression"),
    label.text = NULL,
    scaled = FALSE,
    normalized = FALSE,
    axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
    ...
  )

scale_y_A_total_continuous(
  unit.exponent = 0,
  name = A_label(unit.exponent = unit.exponent, format = format, label.text = label.text,
    scaled = scaled, normalized = ifelse(is.numeric(normalized), round(normalized, 1),
    unique(normalized))), axis.symbols = axis.symbols, Tfr.type = "total"),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

```

### Arguments

<code>unit.exponent</code>	integer
<code>name</code>	The name of the scale, used for the axis-label.
<code>labels</code>	The tick labels or a function to generate them.
<code>format</code>	character string, "R", "R.expression", "R.character", or "LaTeX".
<code>label.text</code>	character Textual portion of the labels.
<code>scaled</code>	logical If TRUE relative units are assumed.
<code>normalized</code>	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
<code>axis.symbols</code>	logical If TRUE symbols of the quantities are added to the name. Supported only by <code>format = "R.expression"</code> .
<code>Tfr.type</code>	character, either "total" or "internal".
<code>...</code>	other named arguments passed to <code>scale_y_continuous</code>

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

### Examples

```

ggplot(yellow_gel.spct, plot.qty = "absorbance") +
  geom_line() +
  scale_y_A_continuous(Tfr.type = getTfrType(yellow_gel.spct)) +
  scale_x_wl_continuous()

```

```

ggplot(yellow_gel.spct, plot.qty = "absorbance") +
  geom_line() +
  scale_y_A_internal_continuous() +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct, plot.qty = "absorbance") +
  geom_line() +
  scale_y_A_total_continuous() +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct, plot.qty = "absorbance") +
  geom_line() +
  scale_y_A_total_continuous(axis.symbols = FALSE) +
  scale_x_wl_continuous(axis.symbols = FALSE)

ggplot(yellow_gel.spct, plot.qty = "absorbance") +
  geom_line() +
  scale_y_A_internal_continuous(normalized = "none") +
  scale_x_wl_continuous()

```

---

scale\_y\_counts\_continuous

*Raw-counts y-scale*

---

## Description

Scale y continuous with defaults suitable for raw detector counts.

## Usage

```

scale_y_counts_continuous(
  unit.exponent = ifelse(normalized, 0, 3),
  name = counts_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["counts"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

scale_y_counts_tg_continuous(
  unit.exponent = ifelse(normalized, 0, 3),
  name = counts_label(unit.exponent = 0, format = format, label.text = label.text, scaled

```

```

    = scaled, normalized = ifelse(is.numeric(normalized), round(normalized, 1),
      unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_tg_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["counts"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

```

### Arguments

<code>unit.exponent</code>	integer
<code>name</code>	The name of the scale, used for the axis-label.
<code>labels</code>	The tick labels or a function to generate them.
<code>format</code>	character string, "R", "R.expression", "R.character", or "LaTeX".
<code>label.text</code>	character Textual portion of the labels.
<code>scaled</code>	logical If TRUE relative units are assumed.
<code>normalized</code>	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
<code>axis.symbols</code>	logical If TRUE symbols of the quantities are added to the name. Supported only by <code>format = "R.expression"</code> .
<code>...</code>	other named arguments passed to <code>scale_y_continuous</code>

### Note

This function only alters default arguments values for `name` and `labels`, please, see documentation for [scale\\_continuous](#) for other parameters.

### Examples

```

ggplot(white_led.raw_spct) +
  geom_line() +
  scale_y_counts_continuous() +
  scale_x_wl_continuous()

ggplot(white_led.raw_spct) +
  geom_line() +
  scale_y_counts_continuous(unit.exponent = 0) +
  scale_x_wl_continuous()

ggplot(white_led.raw_spct) +
  geom_line() +
  scale_y_counts_tg_continuous() +
  scale_x_wl_continuous()

ggplot(white_led.raw_spct) +
  geom_line() +

```

```

scale_y_counts_tg_continuous(unit.exponent = 0) +
scale_x_wl_continuous()

if (packageVersion("photobiology") > "0.11.4") {
  norm_led.raw_spct <- normalize(white_led.raw_spct, norm = "max")

  ggplot(norm_led.raw_spct) +
    geom_line() +
    scale_y_counts_continuous(unit.exponent = 0, normalized = "max") +
    scale_x_wl_continuous()
}

```

---

scale\_y\_cps\_continuous

*Counts-per-second y-scale*

---

### Description

Scale y continuous with defaults suitable for raw detector counts.

### Usage

```

scale_y_cps_continuous(
  unit.exponent = 0,
  name = cps_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["cps"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

```

### Arguments

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).

axis.symbols    logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".

...            other named arguments passed to scale\_y\_continuous

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

### Examples

```
ggplot(white_led.cps_spct) +
  geom_line() +
  scale_y_cps_continuous() +
  scale_x_wl_continuous()

ggplot(white_led.cps_spct) +
  geom_line() +
  scale_y_cps_continuous(3) +
  scale_x_wl_continuous()

ggplot(white_led.cps_spct * 1e-4) +
  geom_line() +
  scale_y_cps_continuous(scaled = TRUE) +
  scale_x_wl_continuous()

if (packageVersion("photobiology") > "0.11.4") {
  norm_led.cps_spct <- normalize(white_led.cps_spct, norm = "max")

  ggplot(norm_led.cps_spct) +
    geom_line() +
    scale_y_cps_continuous(normalized = is_normalized(norm_led.cps_spct)) +
    scale_x_wl_continuous()

  ggplot(norm_led.cps_spct) +
    geom_line() +
    scale_y_cps_continuous(normalized = getNormalized(norm_led.cps_spct)) +
    scale_x_wl_continuous()

  ggplot(norm_led.cps_spct) +
    geom_line() +
    scale_y_cps_continuous(normalized =
      normalization(norm_led.cps_spct)$norm.type) +
    scale_x_wl_continuous()
}
```

---

scale\_y\_multipliers\_continuous

*Calibration multipliers y-scale*

---

**Description**

Scale y continuous with defaults suitable for raw the calibration multipliers used to convert pixel response rate (counts per second) into energy irradiance units.

**Usage**

```
scale_y_multipliers_continuous(
  unit.exponent = 0,
  name = multipliers_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["e.mult"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)
```

**Arguments**

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
...	other named arguments passed to scale_y_continuous

**Note**

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

---

scale\_y\_Rfr\_continuous

*Reflectance y-scale*

---

**Description**

Scale y continuous with defaults suitable for spectral reflectance.

**Usage**

```

scale_y_Rfr_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Rfr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized))), axis.symbols = axis.symbols, Rfr.type =
    Rfr.type),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  Rfr.type,
  ...
)

scale_y_Rfr_specular_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Rfr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized))), axis.symbols = axis.symbols, Rfr.type =
    "specular"),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  ...
)

scale_y_Rfr_total_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Rfr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized))), axis.symbols = axis.symbols, Rfr.type =
    "total"),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,

```

```

axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
pc.out = getOption("ggspectra.pc.out", default = FALSE),
...
)

```

### Arguments

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
limits	One of NULL for default based on data range, a numeric vector of length two (NA allowed) or a function that accepts the data-based limits as argument and returns new limits.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
pc.out	logical, if TRUE use percent as default instead of fraction of one.
Rfr.type	character, either "total" or "spcular".
...	other named arguments passed to scale_y_continuous

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

### Examples

```

ggplot(Ler_leaf_rflt.spct) +
  geom_line() +
  scale_y_Rfr_continuous(Rfr.type = getRfrType(Ler_leaf_rflt.spct)) +
  scale_x_wl_continuous()

ggplot(Ler_leaf_rflt.spct) +
  geom_line() +
  scale_y_Rfr_continuous(unit.exponent = -2,
                        Rfr.type = getRfrType(Ler_leaf_rflt.spct)) +
  scale_x_wl_continuous()

ggplot(Ler_leaf_rflt.spct) +
  geom_line() +
  scale_y_Rfr_continuous(unit.exponent = -3,
                        Rfr.type = getRfrType(Ler_leaf_rflt.spct)) +
  scale_x_wl_continuous()

ggplot(Ler_leaf_rflt.spct) +

```

```

geom_line() +
scale_y_Rfr_specular_continuous() +
scale_x_wl_continuous()

ggplot(Ler_leaf_rflt.spct) +
  geom_line() +
  scale_y_Rfr_specular_continuous(axis.symbols = FALSE) +
  scale_x_wl_continuous(axis.symbols = FALSE)

ggplot(normalize(Ler_leaf_rflt.spct)) +
  geom_line() +
  scale_y_Rfr_continuous(Rfr.type = getRfrType(Ler_leaf_rflt.spct),
    normalized = "max") +
  scale_x_wl_continuous()

```

---

scale\_y\_s.e.irrad\_continuous

*Spectral irradiance y-scale*

---

## Description

Scale y continuous with defaults suitable for raw detector counts.

## Usage

```

scale_y_s.e.irrad_continuous(
  unit.exponent = 0,
  name = s.e.irrad_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.e.irrad"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

scale_y_s.q.irrad_continuous(
  unit.exponent = ifelse(normalized, 0, -6),
  name = s.q.irrad_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.q.irrad"]],

```

```

    scaled = FALSE,
    normalized = FALSE,
    axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
    ...
)

scale_y_s.e.irrad_log10(
  unit.exponent = 0,
  name = s.e.irrad_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.e.irrad"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

scale_y_s.q.irrad_log10(
  unit.exponent = ifelse(normalized, 0, -6),
  name = s.q.irrad_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.q.irrad"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

```

### Arguments

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the default name.
...	other named arguments passed to scale_y_continuous

**Note**

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

**Examples**

```

ggplot(sun.spct) +
  geom_line() +
  scale_y_s.e.irrad_continuous() +
  scale_x_wl_continuous()

ggplot(sun.spct) +
  geom_line() +
  scale_y_s.e.irrad_continuous(label.text = "") +
  scale_x_wl_continuous()

ggplot(sun.spct) +
  geom_line() +
  scale_y_s.e.irrad_continuous(label.text = "Irradiancia spectral,") +
  scale_x_wl_continuous(label.text = "Longitud de onda,")

ggplot(sun.spct) +
  geom_line() +
  scale_y_s.e.irrad_continuous(unit.exponent = -1) +
  scale_x_wl_continuous()

ggplot(sun.spct, unit.out = "photon") +
  geom_line() +
  scale_y_s.q.irrad_continuous() +
  scale_x_wl_continuous()

ggplot(clip_wl(sun.spct, c(295, NA))) +
  geom_line() +
  scale_y_s.e.irrad_log10() +
  scale_x_wl_continuous()

ggplot(clip_wl(sun.spct, c(295, NA)),
  unit.out = "photon") +
  geom_line(na.rm = TRUE) +
  scale_y_s.q.irrad_log10() +
  scale_x_wl_continuous()

if (packageVersion("photobiology") < "0.11.4") {
  photon_as_default()
  normalized_sun.spct <- normalize(e2q(sun.spct, action = "replace"))
  ggplot(normalized_sun.spct) +
    geom_line(na.rm = TRUE) +
    scale_y_s.q.irrad_continuous(normalized =
      normalization(normalized_sun.spct)$norm.wl) +
    scale_x_wl_continuous()

  ggplot(normalized_sun.spct) +
    geom_line(na.rm = TRUE) +

```

```

    scale_y_s.q.irrad_continuous(normalized =
                                normalization(normalized_sun.spct)$norm.type) +
    scale_x_wl_continuous()

unset_radiation_unit_default()
}

```

---

scale\_y\_s.e.response\_continuous

*Spectral response and action y-scales*

---

## Description

Scale y continuous with defaults suitable for response and action spectra.

## Usage

```

scale_y_s.e.response_continuous(
  unit.exponent = 0,
  name = s.e.response_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = -unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.e.response"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

scale_y_s.q.response_continuous(
  unit.exponent = 0,
  name = s.q.response_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
  labels = SI_pl_format(exponent = -unit.exponent),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["s.q.response"]],
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  ...
)

scale_y_s.e.action_continuous(

```

```

unit.exponent = 0,
name = s.e.action_label(unit.exponent = unit.exponent, format = format, label.text =
  label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
labels = SI_pl_format(exponent = -unit.exponent),
format = getOption("photobiology.math", default = "R.expression"),
label.text = axis_labels()[["s.e.action"]],
scaled = FALSE,
normalized = FALSE,
axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
...
)

scale_y_s.q.action_continuous(
  unit.exponent = 0,
  name = s.q.action_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
      round(normalized, 1), unique(normalized)), axis.symbols = axis.symbols),
labels = SI_pl_format(exponent = -unit.exponent),
format = getOption("photobiology.math", default = "R.expression"),
label.text = axis_labels()[["s.q.action"]],
scaled = FALSE,
normalized = FALSE,
axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
...
)

```

### Arguments

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
...	other named arguments passed to scale_y_continuous

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#).

**Examples**

```

ggplot(ccd.spct) +
  geom_line() +
  scale_y_s.e.action_continuous() + # per joule
  scale_x_wl_continuous()

ggplot(ccd.spct) +
  geom_line() +
  scale_y_s.e.response_continuous() + # per joule
  scale_x_wl_continuous()

ggplot(ccd.spct) +
  geom_line() +
  scale_y_s.e.response_continuous(unit.exponent = 6) + # per mega joule
  scale_x_wl_continuous()

ggplot(ccd.spct, unit.out = "photon") +
  geom_line() +
  scale_y_s.q.response_continuous() + # per mol
  scale_x_wl_continuous()

ggplot(ccd.spct, unit.out = "photon") +
  geom_line() +
  scale_y_s.q.response_continuous(unit.exponent = 3) + # per 1000 moles
  scale_x_wl_continuous()

norm_ccd.spct <- normalize(ccd.spct, norm = "max")
ggplot(norm_ccd.spct) +
  geom_line() +
  scale_y_s.e.response_continuous(normalized = getNormalized(norm_ccd.spct)) +
  scale_x_wl_continuous()

if (packageVersion("photobiology") > "0.11.4") {
  ggplot(norm_ccd.spct) +
    geom_line() +
    scale_y_s.e.response_continuous(normalized =
      normalization(norm_ccd.spct)$norm.type) +
    scale_x_wl_continuous()
}
photon_as_default()

norm_ccd.spct <- normalize(ccd.spct, norm = "max")
ggplot(norm_ccd.spct) +
  geom_line() +
  scale_y_s.q.response_continuous(normalized = getNormalized(norm_ccd.spct)) +
  scale_x_wl_continuous()

ggplot(norm_ccd.spct) +
  geom_line() +
  scale_y_s.q.response_continuous(unit.exponent = 2,
                                normalized = getNormalized(norm_ccd.spct)) +
  scale_x_wl_continuous()

```

```
unset_radiation_unit_default()
```

---

```
scale_y_Tfr_continuous
```

*Transmittance y-scale*

---

### Description

Scale y continuous with defaults suitable for spectral transmittance.

### Usage

```
scale_y_Tfr_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Tfr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized))), axis.symbols = axis.symbols, Tfr.type =
    Tfr.type),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  Tfr.type,
  ...
)

scale_y_Tfr_internal_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Tfr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized))), axis.symbols = axis.symbols, Tfr.type =
    "internal"),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  ...
)
```

```

)

scale_y_Tfr_total_continuous(
  unit.exponent = ifelse(pc.out, -2, 0),
  name = Tfr_label(unit.exponent = unit.exponent, format = format, label.text =
    label.text, scaled = scaled, normalized = ifelse(is.numeric(normalized),
    round(normalized, 1), unique(normalized))), axis.symbols = axis.symbols, Tfr.type =
    "total"),
  labels = SI_pl_format(exponent = unit.exponent),
  limits = c(0, 1),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  ...
)

```

### Arguments

unit.exponent	integer
name	The name of the scale, used for the axis-label.
labels	The tick labels or a function to generate them.
limits	One of NULL for default based on data range, a numeric vector of length two (NA allowed) or a function that accepts the data-based limits as argument and returns new limits.
format	character string, "R", "R.expression", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in nanometers (nm).
axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
pc.out	logical, if TRUE use percent as default instead of fraction of one.
Tfr.type	character, either "total" or "internal".
...	other named arguments passed to scale_y_continuous

### Note

This function only alters two default arguments, please, see documentation for [scale\\_continuous](#)

### Examples

```

Tfr_as_default()

ggplot(yellow_gel.spct) +

```

```

geom_line() +
scale_y_Tfr_continuous(Tfr.type = getTfrType(yellow_gel.spct)) +
scale_x_wl_continuous()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Tfr_continuous(unit.exponent = -2,
                        Tfr.type = getTfrType(yellow_gel.spct)) +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Tfr_continuous(unit.exponent = -3,
                        Tfr.type = getTfrType(yellow_gel.spct)) +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Tfr_total_continuous() +
  scale_x_wl_continuous()

ggplot(yellow_gel.spct) +
  geom_line() +
  scale_y_Tfr_total_continuous(axis.symbols = FALSE) +
  scale_x_wl_continuous(axis.symbols = FALSE)

ggplot(normalize(yellow_gel.spct)) +
  geom_line() +
  scale_y_Tfr_total_continuous(normalized = "max") +
  scale_x_wl_continuous()

unset_filter_qty_default()

```

---

sec\_axis\_w\_number      *Secondary axes for wavelengths*

---

### Description

Secondary axes for wavelength data in nanometres. With suitable scaling and name (axis label) for frequency, wave number, photon energy and wavelength.

### Usage

```

sec_axis_w_number(
  unit.exponent = -6,
  label.text = axis_labels()[["w.number"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

```

```

sec_axis_w_frequency(
  unit.exponent = 12,
  label.text = axis_labels()[["freq"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

sec_axis_energy_eV(
  unit.exponent = 0,
  label.text = axis_labels()[["energy"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

sec_axis_energy_J(
  unit.exponent = -18,
  label.text = axis_labels()[["energy"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

sec_axis_wl(
  unit.exponent = -9,
  label.text = axis_labels()[["w.length"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

```

### Arguments

<code>unit.exponent</code>	integer The exponent on base 10 of the scale multiplier used for the axis labels, e.g., 3 for $10^3$ or $k$ .
<code>label.text</code>	character Textual portion of the labels.
<code>axis.symbols</code>	logical If TRUE symbols of the quantities are added to the name. Supported only by <code>format = "R.expression"</code> .

### Details

These secondary axis functions can be used only when the  $x$  aesthetic is mapped to a numerical variable containing wavelength values expressed in nanometres. They can be used to add a secondary  $x$  axis to plots created using `ggplot()` or `autoplot()`.

### See Also

the default text used for quantity names are most easily changed by resetting all the defaults once as explained in [axis\\_labels\\_uk](#), even if it is possible to override them also in each call.

### Examples

```

ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(sec.axis = sec_axis_w_number())

```

```

# Secondary axes can be added to plots built with autoplot() methods
autoplot(sun.spct) +
  scale_x_wl_continuous(sec.axis = sec_axis_w_number())

# Using 'ggplot2' scale
ggplot(sun.spct) +
  geom_line() +
  scale_x_continuous(name = w_length_label(),
                    sec.axis = sec_axis_w_number())

# change scale multipliers, SI defined
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(-6, sec.axis = sec_axis_w_number(-3))

# change scale multipliers, not SI defined (best avoided)
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(-8, sec.axis = sec_axis_w_number(-4))

# Change quantity name to Spanish
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(label.text = "Longitud de onda,",
                      sec.axis = sec_axis_w_frequency(label.text = "Frecuencia,"))

# Frequency in secondary axis
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(sec.axis = sec_axis_w_frequency())

# Energy (per photon) in atto joules
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(sec.axis = sec_axis_energy_J())

# Energy (per photon) in electron volts
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(sec.axis = sec_axis_energy_eV())

# Secondary axis with wavelength using a different scale factor
ggplot(sun.spct) +
  geom_line() +
  scale_x_wl_continuous(sec.axis = sec_axis_wl(-6))

# Secondary axes can be added to plots built with autoplot() methods
autoplot(sun.spct) +
  scale_x_wl_continuous(sec.axis = sec_axis_wl(-6))

```

---

`set_annotations_default`*Set defaults for autoplot annotations*

---

## Description

Set R options used when plotting spectra. Option "photobiology.plot.annotations" is used as default argument to formal parameter annotations and option "photobiology.plot.bands" is used as default argument to formal parameter w.band in all the autoplot() methods exported from package 'ggspectra'. These convenience functions make it easier to edit these two option which are stored as a vector of characters strings and a list of waveband objects, respectively.

## Usage

```
set_annotations_default(annotations = NULL)
```

```
set_w.band_default(w.band = NULL)
```

## Arguments

annotations	character vector Annotations to add or remove from defaults used by the autoplot() methods defined in this package..
w.band	a single waveband object or a list of waveband objects.

## Value

Previous value of option "photobiology.plot.annotations", returned invisibly.

## Plot Annotations

The recognized annotation names are: "summaries", "peaks", "peak.labels", "valleys", "valley.labels", "wls", "wls.labels", "colour.guide", "color.guide", "boxes", "segments", "labels". In addition, "+" is interpreted as a request to add to the already present default annotations, "-" as request to remove annotations and "=" or missing "+" and "-" as a request to reset annotations to those requested. If used, "+", "-" or "=" must be the first member of a character vector, and followed by one or more of the names given above. To simultaneously add and remove annotations one can pass a list containing character vectors each assembled as described. The vectors are applied in the order they appear in the list. To disable all annotations pass "" or c("=", "") as argument. Adding a variation of an annotation already present, replaces the existing one automatically: e.g., adding "peak.labels" replaces "peaks" if present.

## Title Annotations

metadata retrieved from object object is passed to ggplot2::ggtitle() as arguments for title, subtitle and caption. The specification for the title is passed as argument to annotations, and consists in the keyword title with optional modifiers selecting the kind of metadata to use, separated by colons. Up to three keywords separated by colons are accepted, and correspond to title,

subtitle and caption. The recognized keywords are: "objt", "class", "what", "when", "where", "how", "inst.name", "inst.sn", "comment" and "none" are recognized as modifiers to "title"; "none" is a placeholder. Default is "title:objt" or no title depending on the context.

### Note

The syntax used and behaviour are the same as for the annotations parameter of the `autoplot()` methods for spectra, but instead of affecting a single plot, `set_annotations_default()` changes the default used for subsequent calls to `autoplot()`.

### See Also

Other autoplot methods: `autoplot.calibration_spct()`, `autoplot.cps_spct()`, `autoplot.filter_spct()`, `autoplot.object_spct()`, `autoplot.raw_spct()`, `autoplot.reflector_spct()`, `autoplot.response_spct()`, `autoplot.source_spct()`, `autoplot.waveband()`

---

SI\_pl\_format

*Formatter for plain labels discounting for SI multipliers*

---

### Description

The labels generated represent numbers rescaled to compensate for a change in unit's by a factor of ten or by a power of ten.

### Usage

```
SI_pl_format(exponent = 0, digits = 3, ...)
```

```
SI_plain(x, exponent = 0, digits = 3, ...)
```

### Arguments

exponent	numeric Power of 10 to use as multiplier
digits	number of significant digits to show
...	other arguments passed on to <code>format</code>
x	a numeric vector to format

### Value

a function with single parameter x, a numeric vector, that returns a character vector

### Examples

```
SI_pl_format()(1:10)
SI_pl_format()(runif(10))
SI_pl_format(exponent = 2)(runif(10))
SI_plain(1:10)
SI_plain(runif(10))
SI_plain(runif(10), digits = 2)
```

---

SI_tg_format	<i>Formatter for tagged labels using SI multipliers</i>
--------------	---

---

### Description

The labels generated represent the same numbers, but with trailing zeros removed/added and compensated by attaching to each label an SI multiplier "prefix".

### Usage

```
SI_tg_format(exponent = 0, digits = 3, ...)
```

```
SI_tagged(x, exponent = 0, digits = 3, ...)
```

### Arguments

exponent	numeric Power of 10 to use as multiplier
digits	number of significant digits to show
...	other arguments passed on to <a href="#">format</a>
x	a numeric vector to format

### Value

a function with single parameter x, a numeric vector, that returns a character vector

### Note

If the exponent passed has no SI prefix defined, the exponent will be adjusted to match one.

### Examples

```
SI_tg_format()(1:10)
SI_tg_format()(runif(10))
SI_tg_format(exponent = 2)(runif(10))
SI_tagged(1:10)
SI_tagged(runif(10))
SI_tagged(runif(10), digits = 2)
```

---

stat\_color

*Calculate colours from wavelength.*


---

### Description

stat\_color computes colour definitions according to human vision.

### Usage

```
stat_color(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  ...,
  chroma.type = "CMF",
  x.colour.transform = I,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)
```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
x.colour.transform	function Applied to x values before computing matching colours.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

**Details**

For each row in data a colour definition is computed assuming that after transformation with `x.colour.transform()` the values in `x` are wavelengths expressed in nanometres.

**Value**

The original data frame with variable `wl.color` containing colour definitions added.

**Computed variable**

**wl.color** color corresponding to `x`-value giving wavelength in nanometres.

**Default aesthetics**

Set by the statistic and available to geoms.

**color** `..wl.color..`

**fill** `..wl.color..`

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**See Also**

[color\\_of](#), which is used internally.

Other stats functions: [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
ggplot(sun.spct) +
  geom_line() +
  stat_color() +
  scale_color_identity()
```

```
ggplot(sun.spct) +
  geom_line() +
  stat_color(x.colour.transform = function(x) {-x}) +
  scale_color_identity() +
  scale_x_reverse()
```

```
ggplot(sun.spct) +
  geom_line() +
  stat_color(x.colour.transform = function(x) {10^x}) +
```

```
scale_color_identity() +
scale_x_log10()
```

---

stat_find_qtys	<i>Find quantity value for target wavelength value.</i>
----------------	---

---

### Description

stat\_find\_qtys finds at which y positions values equal to an x target are located. **Axis flipping is currently not supported.**

### Usage

```
stat_find_qtys(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  ...,
  target = "half.maximum",
  interpolate = TRUE,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  x.label.fmt = label.fmt,
  y.label.fmt = label.fmt,
  x.label.transform = I,
  y.label.transform = I,
  x.colour.transform = x.label.transform,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)
```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
target	numeric value indicating the spectral quantity value for which wavelengths are to be searched and interpolated if need. The character string "half.maximum" is also accepted as argument.

<code>interpolate</code>	logical Indicating whether the nearest wavelength value in <code>x</code> should be returned or a value calculated by linear interpolation between wavelength values straddling the target.
<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
<code>label.format</code> , <code>x.label.format</code> , <code>y.label.format</code>	character strings giving a format definition for construction of character strings labels with function <code>sprintf</code> from <code>x</code> and/or <code>y</code> values.
<code>x.label.transform</code> , <code>y.label.transform</code> , <code>x.colour.transform</code>	function Applied to <code>x</code> or <code>y</code> values when constructing the character labels or computing matching colours.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Details

These stats use `geom_point` by default as it is the geom most likely to work well in almost any situation without need of tweaking. The default aesthetics set by these stats allow their direct use with `geom_text`, `geom_label`, `geom_line`, `geom_rug`, `geom_hline` and `geom_vline`. The formatting of the labels returned can be controlled by the user.

### Value

A data frame with one row for each match to the target subset from the data or linearly interpolated between the two nearest values available. As spectra are monotonic in wavelength, this statistic will never return more than one row in data per target value.

### Computed variables

**x** x-value at or nearest to the match to the target as numeric  
**y** target value or y-value nearest to the target as numeric  
**x.label** x-value at or nearest to the match formatted as character  
**y.label** target value or y-value nearest to the target formatted as character  
**color** color definition calculated by assuming that x-values are wavelengths expressed in nanometres.

### Default aesthetics

Set by the statistic and available to geoms.

**label** `..x.label..`

**xintercept** `..x..`

**yintercept** ..y..

**fill** ..color..

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

### Note

These stats work nicely together with geoms `geom_text_repel` and `geom_label_repel` from package [ggrepel](#) to solve the problem of overlapping labels by displacing them. To discard overlapping labels use `check_overlap = TRUE` as argument to `geom_text`. By default the labels are character values suitable to be plotted as is, but with a suitable `label.fmt` labels suitable for parsing by the geoms (e.g. into expressions containing greek letters or super or subscripts) can be also easily obtained.

### See Also

[find\\_peaks](#).

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(yellow_gel.spct) +
  geom_line() +
  stat_find_qtys(target = "half.range")

ggplot(yellow_gel.spct) +
  geom_line() +
  stat_find_qtys(target = c(490, 500, 510))

ggplot(yellow_gel.spct) +
  geom_line() +
  stat_find_qtys(target = 500, geom = "point", colour = "red") +
  stat_find_qtys(target = 500, geom = "text", colour = "red",
    hjust = 1.1, label.fmt = "Tfr = %1.2f")
```

---

stat_find_wls	<i>Find wavelength for target quantity value.</i>
---------------	---

---

### Description

stat\_find\_wls finds at which x positions values equal to a target are located. **Axis flipping is currently not supported.**

### Usage

```
stat_find_wls(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  ...,
  target = "half.maximum",
  interpolate = TRUE,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  x.label.fmt = label.fmt,
  y.label.fmt = label.fmt,
  x.label.transform = I,
  y.label.transform = I,
  x.colour.transform = x.label.transform,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)
```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
target	numeric vector indicating the spectral quantity values for which wavelengths are to be searched and interpolated if need. The character strings "half.maximum" and "half.range" are also accepted as arguments. A list with numeric and/or character values is also accepted.
interpolate	logical Indicating whether the nearest wavelength value in x should be returned or a value calculated by linear interpolation between wavelength values straddling the target.

<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
<code>label.fmt</code> , <code>x.label.fmt</code> , <code>y.label.fmt</code>	character strings giving a format definition for construction of character strings labels with function <code>sprintf</code> from x and/or y values.
<code>x.label.transform</code> , <code>y.label.transform</code> , <code>x.colour.transform</code>	function Applied to x or y values when constructing the character labels or computing matching colours.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

## Details

For each row in the subset of data matching `target` a colour definition is computed assuming that after transformation with `x.colour.transform()` the values in `x` are wavelengths expressed in nanometres. Labels are constructed from `x` and `y` values after applying to them `x.label.transform` and `y.label.transform`, respectively. In most cases the `x.label.transform` is used to back-transform the values in data to make them agree with those displayed on the axis guides.

These stats use `geom_point` by default as it is a geometry likely to work well in almost any situation. The additional default aesthetic mappings set by these statistics allow their direct use with `geom_text`, `geom_label`, `geom_line`, `geom_rug`, `geom_hline` and `geom_vline`. The format of the labels returned can be controlled by the user.

## Value

A data frame with one row for each match to the target subset from the data or linearly interpolated between the two nearest values available. As spectra are not monotonic in the spectral quantity, this statistic can return more than one row in data per target value.

## Computed variables

**x** x-value at or nearest to the match to the target as numeric

**y** target value or y-value nearest to the target as numeric

**x.label** x-value at or nearest to the match formatted as character

**y.label** target value or y-value nearest to the target formatted as character

**wl.color** color definition calculated by assuming that x-values are wavelengths expressed in nanometres.

**Default aesthetics**

Set by the statistic and available to geoms.

**label** ..x.label..

**xintercept** ..x..

**yintercept** ..y..

**fill** ..wl.color..

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**Note**

These stats work nicely together with geoms `geom_text_repel` and `geom_label_repel` from package `ggrepel` to solve the problem of overlapping labels by displacing them. To discard overlapping labels use `check_overlap = TRUE` as argument to `geom_text`. By default the labels are character values suitable to be plotted as is, but with a suitable `label.fmt` labels suitable for parsing by the geoms (e.g. into expressions containing greek letters or super or subscripts) can be also easily obtained.

**See Also**

[find\\_peaks](#).

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(yellow_gel.spct) +
  geom_line() +
  stat_find_wls(target = c(0.25, 0.5, 0.75))

ggplot(yellow_gel.spct) +
  geom_line() +
  stat_find_wls(target = "half.maximum", geom = "point", colour = "red") +
  stat_find_wls(target = "half.maximum", geom = "text", colour = "red",
               hjust = 1.1, label.fmt = "%3.0f nm")
```

---

stat_label_peaks	<i>Label peaks and valleys.</i>
------------------	---------------------------------

---

### Description

stat\_labels\_peaks finds at which x positions local maxima are located, and adds labels and colors to the data without subsetting. To find local minima, you can use stat\_labels\_valleys instead. The variable mapped to the x aesthetic is expected to contain wavelength values expressed in nanometres. **Axis flipping is currently not supported.**

### Usage

```
stat_label_peaks(  
  mapping = NULL,  
  data = NULL,  
  geom = "text",  
  position = "identity",  
  ...,  
  span = 5,  
  ignore_threshold = 0,  
  strict = TRUE,  
  chroma.type = "CMF",  
  label.fmt = "%.3g",  
  x.label.fmt = label.fmt,  
  y.label.fmt = label.fmt,  
  x.label.transform = I,  
  y.label.transform = I,  
  x.colour.transform = x.label.transform,  
  label.fill = "",  
  na.rm = TRUE,  
  show.legend = FALSE,  
  inherit.aes = TRUE  
)
```

```
stat_label_valleys(  
  mapping = NULL,  
  data = NULL,  
  geom = "text",  
  position = "identity",  
  ...,  
  span = 5,  
  ignore_threshold = 0,  
  strict = TRUE,  
  chroma.type = "CMF",  
  label.fmt = "%.3g",  
  x.label.fmt = label.fmt,  
  y.label.fmt = label.fmt,
```

```

x.label.transform = I,
y.label.transform = I,
x.colour.transform = x.label.transform,
label.fill = "",
na.rm = TRUE,
show.legend = FALSE,
inherit.aes = TRUE
)

```

## Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
span	a peak is defined as an element in a sequence which is greater than all other elements within a window of width span centered at that element. The default value is 5, meaning that a peak is bigger than two consecutive neighbors on each side. Default: 5.
ignore_threshold	numeric value between 0.0 and 1.0 indicating the size threshold below which peaks will be ignored.
strict	logical flag: if TRUE, an element must be strictly greater than all other values in its window to be considered a peak. Default: FALSE.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spect</a> object.
label.fmt, x.label.fmt, y.label.fmt	character strings giving a format definition for construction of character strings labels with function <a href="#">sprintf</a> from x and/or y values.
x.label.transform, y.label.transform, x.colour.transform	function Applied to x or y values when constructing the character labels or computing matching colours.
label.fill	character string to use for labels not at peaks or valleys being highlighted.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

## Details

These statistics assemble text labels for each peak or valley and compute the colour corresponding to the wavelength of the peaks and valleys. Defaults work as long as the variable mapped to the `x` aesthetic contains wavelengths expressed in nanometres and the plot has an `x`-scale that does not apply a transformation. The three `transform` parameters can be used to back-transform the values when scales apply transformations so that peak/valley labels and axis labels match. Of course, `x.label.transform` and `y.label.transform` make also possible to scale the values in the labels.

Both statistics use `geom_text` by default as it is the geom most likely to work well in almost any situation without need of tweaking. These statistics work best with `geom_text_repel` and `geom_label_repel` from package 'ggrepel' as they are designed so that peak or valley labels will not overlap any observation in the whole data set. Default aesthetics set by these statistics allow their direct use with `geom_text`, `geom_label`, `geom_line`, `geom_rug`, `geom_hline` and `geom_vline`. The formatting of the labels returned can be controlled by the user.

## Value

The original data with additional computed variables added.

## Computed variables

**x.label** x-value at a peak (or valley) formatted as character or otherwise the value passed to `label.fill` which defaults to an empty string ("").

**y.label** y-value at the peak (or valley) formatted as character or otherwise the value passed to `label.fill` which defaults to an empty string ("").

**wl.color** At peaks and valleys, color definition calculated by assuming that x-values are wavelengths expressed in nanometres, otherwise, `rgb(1, 1, 1, 0)` (transparent white).

## Default aesthetics

Set by the statistic and available to geoms.

**label** `..x.label..`

**xintercept** `..x..`

**yintercept** `..y..`

**color** `black_or_white(..wl.color..)`

**fill** `..wl.color..`

## Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**Note**

These stats work nicely together with geoms `geom_text_repel` and `geom_label_repel` from package `ggrepel` to solve the problem of overlapping labels by displacing them. To discard overlapping labels use `check_overlap = TRUE` as argument to `geom_text`. By default the labels are character values suitable to be plotted as is, but with a suitable `label.fmt` labels suitable for parsing by the geoms (e.g. into expressions containing greek letters or super or subscripts) can be also easily obtained.

**See Also**

[stat\\_peaks](#), [stat\\_valleys](#) and [find\\_peaks](#), which is used internally.

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_label_peaks(hjust = "left", span = 31, angle = 90, color = "red")

ggplot(sun.spct) +
  geom_line() +
  stat_label_valleys(hjust = "right", span = 21, angle = 90, color = "blue")

# using transformed scales requires the user to pass functions as arguments
ggplot(sun.spct) +
  geom_line() +
  stat_label_peaks(hjust = "left", span = 31, angle = 90, color = "red",
                  x.label.transform = abs) +
  scale_x_reverse()

ggplot(sun.spct) +
  geom_line() +
  stat_label_peaks(hjust = "left", span = 31, angle = 90, color = "red",
                  x.label.transform = function(x) {10^x}) +
  scale_x_log10()

# geom_label
ggplot(sun.spct) +
  geom_line() +
  stat_peaks(span = 41, shape = 21, size = 3) +
  stat_label_peaks(span = 41, geom = "label", label.fmt = "%3.0f nm") +
  scale_fill_identity() +
  scale_color_identity() +
  expand_limits(y = c(NA, 1))

# using 'ggrepel' to avoid overlaps
# too slow for CRAN checks
```

```
## Not run:
library(ggrepel)

ggplot(sun.spct) + geom_line() +
  stat_peaks(span = 41, shape = 21, size = 3) +
  stat_label_peaks(span = 41, geom = "label_repel", segment.colour = "red",
                  nudge_y = 0.12, label.fmt = "%3.0f nm",
                  max.overlaps = Inf, min.segment.length = 0) +
  scale_fill_identity() +
  scale_color_identity() +
  expand_limits(y = c(NA, 1))

## End(Not run)
```

---

stat\_peaks

*Find peaks and valleys.*


---

### Description

stat\_peaks finds at which x positions local maxima are located. If you want find local minima, you can use stat\_valleys instead. **Axis flipping is currently not supported.**

### Usage

```
stat_peaks(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  ...,
  span = 5,
  ignore_threshold = 0.01,
  strict = is.null(span),
  refine.wl = FALSE,
  method = "spline",
  chroma.type = "CMF",
  label.fmt = "%.3g",
  x.label.fmt = label.fmt,
  y.label.fmt = label.fmt,
  x.label.transform = I,
  y.label.transform = I,
  x.colour.transform = x.label.transform,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)
```

```

stat_valleys(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  ...,
  span = 5,
  ignore_threshold = -0.01,
  strict = is.null(span),
  refine.wl = FALSE,
  method = "spline",
  chroma.type = "CMF",
  label.fmt = "%.3g",
  x.label.fmt = label.fmt,
  y.label.fmt = label.fmt,
  x.label.transform = I,
  y.label.transform = I,
  x.colour.transform = x.label.transform,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)

```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
span	integer A peak is defined as an element in a sequence which is greater than all other elements within a window of width span centered at that element. Use NULL for the global peak. Valleys are the reverse.
ignore_threshold	numeric For peaks, value between 0.0 and 1.0 indicating the relative size of peaks compared to tallest peak threshold below which peaks will be ignored, while negative values between 0.0 and -1.0 set a threshold so that the tallest peaks are ignored, instead of the shortest. For valleys, value between 0.0 and 1.0 indicating the relative depth of valleys below which valleys will be ignored, while negative values between 0.0 and -1.0 set a threshold so that the deeper valleys are ignored, instead of the shallower ones.
strict	logical If TRUE, an element must be strictly greater than all other values in its window to be considered a peak.
refine.wl	logical Flag indicating if peak or valleys locations should be refined by fitting a function.

<code>method</code>	character String with the name of a method used for peak fitting. Currently only spline interpolation is implemented.
<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spect</code> object.
<code>label.fmt</code> , <code>x.label.fmt</code> , <code>y.label.fmt</code>	character strings giving a format definition for construction of character strings labels with function <code>sprintf</code> from x and/or y values.
<code>x.label.transform</code> , <code>y.label.transform</code> , <code>x.colour.transform</code>	function Applied to x or y values when constructing the character labels or computing matching colours.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Details

These stats use `geom_point` by default as it is the geom most likely to work well in almost any situation without need of tweaking. The default aesthetics set by these stats allow their direct use with `geom_text`, `geom_label`, `geom_line`, `geom_rug`, `geom_hline` and `geom_vline`. The formatting of the labels returned can be controlled by the user.

### Value

A data frame with one row for each peak (or valley) found in the data.

### Computed variables

**x** x-value at the peak (or valley) as numeric

**y** y-value at the peak (or valley) as numeric

**x.label** x-value at the peak (or valley) formatted as character

**y.label** y-value at the peak (or valley) formatted as character

**wl.color** color definition calculated by assuming that x-values are wavelengths expressed in nanometres.

**BW.color** color definition, either "black" or "white", as needed to ensure high contrast to `wl.color`.

### Default aesthetics

Set by the statistic and available to geoms.

**label** `stat(x.label)`

**xintercept** `stat(x)`

**yintercept** `stat(y)`

**fill** `stat(wl.color)`

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**Note**

These stats work nicely together with geoms `geom_text_repel` and `geom_label_repel` from package `ggrepel` to solve the problem of overlapping labels by displacing them. To discard overlapping labels use `check_overlap = TRUE` as argument to `geom_text`. By default the labels are character values suitable to be plotted as is, but with a suitable `label.fmt` labels suitable for parsing by the geoms (e.g. into expressions containing greek letters or super or subscripts) can be also easily obtained.

**See Also**

[find\\_peaks](#), which is used internally.

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_peaks()

ggplot(sun.spct) +
  geom_line() +
  stat_valleys()

ggplot(sun.spct) +
  geom_line() +
  stat_peaks(span = 51, geom = "point", colour = "red") +
  stat_peaks(span = 51, geom = "text", colour = "red",
            vjust = -0.4, label.fmt = "%3.2f nm")

ggplot(sun.spct) +
  geom_line() +
  stat_peaks(span = 51, geom = "point", colour = "red", refine.wl = TRUE) +
  stat_peaks(span = 51, geom = "text", colour = "red",
            vjust = -0.4, label.fmt = "%3.2f nm",
            refine.wl = TRUE)

ggplot(sun.spct) +
  geom_line() +
  stat_peaks(span = 51, geom = "point", colour = "red", refine.wl = TRUE) +
```

```

stat_peaks(mapping = aes(fill = after_stat(wl.colour), color = after_stat(BW.colour)),
           span = 51, geom = "label",
           size = 3, vjust = -0.2, label.fmt = "%.3g nm",
           refine.wl = TRUE) +
stat_valleys(span = 71, geom = "point", colour = "blue", refine.wl = TRUE) +
stat_valleys(mapping = aes(fill = after_stat(wl.colour), color = after_stat(BW.colour)),
             span = 71, geom = "label",
             size = 3, vjust = 1.2, label.fmt = "%.3g nm",
             refine.wl = TRUE) +
expand_limits(y = 0.85) + # make room for label
scale_fill_identity() +
scale_color_identity()

```

---

stat\_spikes

*Find spikes*


---

## Description

stat\_spikes finds at which x positions spikes are located. Spikes can be either upwards or downwards from the baseline. **Axis flipping is currently not supported.**

## Usage

```

stat_spikes(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  ...,
  z.threshold = 9,
  max.spike.width = 8,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  x.label.fmt = label.fmt,
  y.label.fmt = label.fmt,
  x.label.transform = I,
  y.label.transform = I,
  x.colour.transform = x.label.transform,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)

```

## Arguments

mapping      The aesthetic mapping, usually constructed with `aes` or `aes_`. Only needs to be set at the layer level if you are overriding the plot defaults.

<code>data</code>	A layer specific dataset - only needed if you want to override the plot defaults.
<code>geom</code>	The geometric object to use display the data
<code>position</code>	The position adjustment to use for overlapping points on this layer
<code>...</code>	other arguments passed on to <code>layer</code> . This can include aesthetics whose values you want to set, not map. See <code>layer</code> for more details.
<code>z.threshold</code>	numeric Modified Z values larger than <code>z.threshold</code> are considered to be spikes.
<code>max.spike.width</code>	integer Wider regions with high Z values are not detected as spikes.
<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
<code>label.fmt, x.label.fmt, y.label.fmt</code>	character strings giving a format definition for construction of character strings labels with function <code>sprintf</code> from x and/or y values.
<code>x.label.transform, y.label.transform, x.colour.transform</code>	function Applied to x or y values when constructing the character labels or computing matching colours.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Details

This stat uses `geom_point` by default as it is the geom most likely to work well in almost any situation without need of tweaking. The default aesthetics set by this stat allows its direct use with `geom_text`, `geom_label`, `geom_line`, `geom_rug`, `geom_hline` and `geom_vline`. The formatting of the labels returned can be controlled by the user.

### Value

A data frame with one row for each peak (or valley) found in the data.

### Computed variables

**x** x-value at the peak (or valley) as numeric

**y** y-value at the peak (or valley) as numeric

**x.label** x-value at the peak (or valley) formatted as character

**y.label** y-value at the peak (or valley) formatted as character

**wl.color** color definition calculated by assuming that x-values are wavelengths expressed in nanometres.

**BW.color** color definition that either "black" or "white", to ensure high contrast to `wl.color`.

**Default aesthetics**

Set by the statistic and available to geoms.

**label** stat(x.label)

**xintercept** stat(x)

**yintercept** stat(y)

**fill** stat(wl.color)

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**Note**

This stat works nicely together with geoms `geom_text_repel` and `geom_label_repel` from package `ggrepel` to solve the problem of overlapping labels by displacing them. To discard overlapping labels use `check_overlap = TRUE` as argument to `geom_text`. By default the labels are character values suitable to be plotted as is, but with a suitable `label.fmt` labels suitable for parsing by the geoms (e.g. into expressions containing greek letters or super or subscripts) can be also easily obtained.

**See Also**

`find_spikes`, which is used internally, for a description of the algorithm used.

Other stats functions: `stat_color()`, `stat_find_qtys()`, `stat_find_wls()`, `stat_label_peaks()`, `stat_peaks()`, `stat_wb_box()`, `stat_wb_column()`, `stat_wb_contribution()`, `stat_wb_hbar()`, `stat_wb_irrad()`, `stat_wb_label()`, `stat_wb_mean()`, `stat_wb_relative()`, `stat_wb_sirrad()`, `stat_wb_total()`, `stat_wl_strip()`, `stat_wl_summary()`

**Examples**

```
# ggplot() methods for spectral objects set a default mapping for x and y.

# two spurious(?) spikes
ggplot(sun.spct) +
  geom_line() +
  stat_spikes(colour = "red", alpha = 0.3)

# no spikes detected
ggplot(sun.spct) +
  geom_line() +
  stat_spikes(colour = "red", alpha = 0.3,
             max.spike.width = 3,
             z.threshold = 12)

# small noise spikes detected
```

```

ggplot(white_led.raw_spct) +
  geom_line() +
  stat_spikes(colour = "red", alpha = 0.3)

ggplot(white_led.raw_spct) +
  geom_line() +
  stat_spikes(colour = "red", alpha = 0.3) +
  stat_spikes(geom = "text", colour = "red", check_overlap = TRUE,
             vjust = -0.5, label.fmt = "%3.0f nm")

ggplot(white_led.raw_spct, aes(w.length, counts_2)) +
  geom_line() +
  stat_spikes(colour = "red", alpha = 0.3,
             max.spike.width = 3,
             z.threshold = 12)

```

---

stat\_wb\_box

*Draw colour boxes for wavebands*


---

### Description

stat\_wb\_box plots boxes corresponding to wavebands, by default located slightly above the peak of the spectrum. Sets suitable default aesthetics for geom\_rect(). **x-scale transformations and axis flipping are currently not supported.**

### Usage

```

stat_wb_box(
  mapping = NULL,
  data = NULL,
  geom = "rect",
  position = "identity",
  ...,
  w.band = NULL,
  chroma.type = "CMF",
  ypos.mult = 1.07,
  ypos.fixed = NULL,
  box.height = 0.06,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)

```

### Arguments

mapping            The aesthetic mapping, usually constructed with [aes](#) or [aes\\_](#). Only needs to be set at the layer level if you are overriding the plot defaults.

<code>data</code>	A layer specific dataset - only needed if you want to override the plot defaults.
<code>geom</code>	The geometric object to use display the data
<code>position</code>	The position adjustment to use for overlapping points on this layer
<code>...</code>	other arguments passed on to <code>layer</code> . This can include aesthetics whose values you want to set, not map. See <code>layer</code> for more details.
<code>w.band</code>	a waveband object or a list of waveband objects or numeric vector of at least length two.
<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
<code>ypos.mult</code>	numeric Multiplier constant used to compute returned y values. This is numerically similar to using <code>npc</code> units, but values larger than one expand the plotting area.
<code>ypos.fixed</code>	numeric If not NULL used a constant value returned in y.
<code>box.height</code>	numeric The height of the box as a fraction of the range of <code>\$\$</code> . This is similar to using <code>npc</code> units.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named `integral` below is the result of applying `integral.fun` to the data, with default `integrate_xy`.

**x** `w.band-midpoint`

**wb.xmin** `w.band` minimum

**wb.xmax** `w.band` maximum

**wb.ymin** `data$y` minimum

**wb.ymax** `data$y` maximum

**ymin** box bottom

**ymax** box top

**y** `ypos.fixed` or top of data, adjusted by `ypos.mult`

**wb.color** color of the `w.band`

**wb.name** label of `w.band`

**BW.color** `black_or_white(wb.color)`

### Default aesthetics

Set by the statistic and available to geoms.

**xmin** stat(wb.xmin)

**xmax** stat(wb.xmax)

**ymin** stat(ymin)

**ymax** stat(ymax)

**fill** ..wb.color..

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

### Note

This stat uses a panel function and ignores grouping as it is meant to be used for annotations. The value returned as default value for `y` is based on the `y`-range of spectral values for the whole data set.

### See Also

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```
library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  stat_wb_box(w.band = VIS_bands()) +
  geom_line() +
  scale_fill_identity()
ggplot(sun.spct) +
  stat_wb_box(w.band = VIS_bands(), color = "white") +
  geom_line() +
  scale_fill_identity()
```

---

stat_wb_column	<i>Integrate ranges under curve.</i>
----------------	--------------------------------------

---

### Description

stat\_wb\_column computes means under a curve. It first integrates the area under a spectral curve and also the mean expressed per nanometre of wavelength for each waveband in the input. Sets suitable default aesthetics for geom\_rect(). **x-scale transformations and axis flipping are currently not supported.**

### Usage

```
stat_wb_column(
  mapping = NULL,
  data = NULL,
  geom = "rect",
  position = "identity",
  ...,
  w.band = NULL,
  integral.fun = integrate_xy,
  chroma.type = "CMF",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)
```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
integral.fun	function on \$x\$ and \$y\$.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.

`inherit.aes` If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. `borders`.

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named `integral` below is the result of applying `integral.fun`, with default `integrate_xy`.

`x` `w.band`-midpoint

`wb.xmin` `w.band` minimum

`wb.xmax` `w.band` maximum

`wb.ymin` `data$y` minimum

`wb.ymax` `data$y` maximum

`wb.ymean` `yint` divided by `wl_expance(w.band)`

`y` `wb.ymean`

`wb.color` color of the `w.band`

`wb.name` label of `w.band`

`BW.color` `black_or_white(wb.color)`

### Default aesthetics

Set by the statistic and available to geoms.

`xmin` `..wb.xmin..`

`xmax` `..wb.xmax..`

`ymin` 0

`ymax` `..wb.ymean..`

`fill` `..wb.color..`

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

`x` numeric, wavelength in nanometres

`y` numeric, a spectral quantity

### Note

If the argument passed to `w.band` is a BSWF it is silently converted to a wavelength range and the average of spectral values without weighting is returned as default value for `ymax` while the default value for `ymin` is zero.

**See Also**

Other stats functions: `stat_color()`, `stat_find_qtys()`, `stat_find_wls()`, `stat_label_peaks()`, `stat_peaks()`, `stat_spikes()`, `stat_wb_box()`, `stat_wb_contribution()`, `stat_wb_hbar()`, `stat_wb_irrad()`, `stat_wb_label()`, `stat_wb_mean()`, `stat_wb_relative()`, `stat_wb_sirrad()`, `stat_wb_total()`, `stat_wl_strip()`, `stat_wl_summary()`

**Examples**

```
library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands()) +
  geom_line() +
  scale_fill_identity()

ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands(), alpha = 0.5) +
  geom_line() +
  scale_fill_identity()
```

---

stat\_wb\_contribution *Integrate ranges under spectral curve.*

---

**Description**

`stat_wb_contribution` integrates the area under a spectral curve. It first integrates the area under the curve for each waveband and for the whole curve and then expresses the integral for each band as a relative contribution to the area under the whole spectral curve. Sets suitable default aesthetics for "rect", "hline", "vline", "text" and "label" geoms displaying "contributions" per waveband to the total of the spectral integral. **x-scale transformations and axis flipping are currently not supported.**

**Usage**

```
stat_wb_contribution(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  integral.fun = integrate_xy,
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%1.2f",
  ypos.mult = 1.07,
  ypos.fixed = NULL,
```

```

na.rm = FALSE,
show.legend = NA,
inherit.aes = TRUE
)

```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
integral.fun	function on $\$x\$$ and $\$y\$$ .
label.mult	numeric Scaling factor applied to y-integral values before conversion into character strings.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
label.fmt	character string giving a format definition for converting y-integral values into character strings by means of function <a href="#">sprintf</a> .
ypos.mult	numeric Multiplier constant used to scale returned y values.
ypos.fixed	numeric If not NULL used a constant value returned in y.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named `integral` below is the result of applying `integral.fun` to the data, with default `integrate_xy`.

**y.label** yint multiplied by `label.mult` and formatted according to `label.fmt`

**x** w.band-midpoint

**xmin** w.band minimum  
**xmax** w.band maximum  
**ymin** data\$y minimum  
**ymax** data\$y maximum  
**yint** data\$y integral for w.band / data\$y integral for whole range of data\$x  
**xmean** yint divided by wl\_expanse(w.band)  
**y** ypos.fixed or top of data, adjusted by ypos.mult  
**wb.color** color of the w.band  
**wb.name** label of w.band

### Default aesthetics

Set by the statistic and available to geoms.

**label** ..y.label..  
**x** ..x..  
**xmin** ..xmin..  
**xmax** ..xmax..  
**ymin** ..y.. - (..ymax.. - ..ymin..) \* 0.03  
**ymax** ..y.. + (..ymax.. - ..ymin..) \* 0.03  
**yintercept** ..ymean..  
**fill** ..wb.color..

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres  
**y** numeric, a spectral quantity

### See Also

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```

library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.

# Using defaults
ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS()) +

```

```

stat_wb_contribution(w.band = VIS()) +
scale_fill_identity() + scale_color_identity()

# Setting position and angle of the text
ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS_bands()) +
  stat_wb_contribution(w.band = VIS_bands(), angle = 90, size = 2.5) +
  scale_fill_identity() + scale_color_identity()

# Showing percentages, i.e., using a different format for numbers
ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS_bands()) +
  stat_wb_contribution(w.band = VIS_bands(), size = 2.5,
                      label.mult = 100, label.fmt = "%3.0f%") +
  scale_fill_identity() + scale_color_identity()

# Including the name of the waveband, i.e., changing the mapping for label
ggplot(sun.spct, range = c(NA, 410)) +
  geom_line() +
  stat_wb_box(w.band = UV_bands(), color = "white") +
  stat_wb_contribution(w.band = UV_bands(), size = 2.5,
                      label.mult = 100, label.fmt = "%3.0f%",
                      mapping = aes(label = after_stat(paste(wb.name, y.label)))) +
  scale_fill_identity() + scale_color_identity()

```

---

stat\_wb\_hbar

*Integrate ranges under curve.*


---

## Description

stat\_wb\_hbar computes means under a curve. It first integrates the area under a spectral curve and also the mean expressed per nanometre of wavelength for each waveband in the input. Sets suitable default aesthetics for geoms "errorbarh" and "hline" from 'ggplot', and "linerrangeh", and "errorbarh" from 'ggstance'. **x-scale transformations and axis flipping are currently not supported.**

## Usage

```

stat_wb_hbar(
  mapping = NULL,
  data = NULL,
  geom = "linerrange",
  position = "identity",
  ...,
  w.band = NULL,
  integral.fun = integrate_xy,

```

```

  chroma.type = "CMF",
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)

```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
integral.fun	function on $x$ and $y$ .
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
ypos.fixed	numeric If not NULL used a constant value returned in $y$ .
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named `integral` below is the result of applying `integral.fun`, with default `integrate_xy`.

**x** w.band-midpoint

**xmin** w.band minimum

**xmax** w.band maximum

**ymin** data\$y minimum

**ymax** data\$y maximum

**yint** data\$y integral for the range of w.band

**ymean** yint divided by wl\_expanse(w.band)

**y** ypos.fixed or mean of data

**wb.color** color of the w.band

**wb.name** label of w.band

### Default aesthetics

Set by the statistic and available to geoms.

**xmin** ..xmin..

**xmax** ..xmax..

**yintercept** ..ymean..

**height** (..ymax.. - ..ymin..) \* 2e-2

**color** ..wb.color..

### Required aesthetics

Required by the statistic and need to be set with aes().

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

### Note

If the argument passed to w.band is a BSWF it is silently converted to a wavelength range and the average of spectral values without any weighting is returned as default value for y.

### See Also

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```
library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_wb_hbar(w.band = VIS_bands(), size = 1) +
  scale_color_identity() +
  theme_bw()

ggplot(sun.spct) +
  geom_line() +
  stat_wb_hbar(w.band = PAR(), size = 1) +
  scale_color_identity() +
  theme_bw()
```

```

ggplot(sun.spct) +
  geom_line() +
  stat_wb_hbar(w.band = PAR(), size = 1, ypos.fixed = 0) +
  scale_color_identity() +
  theme_bw()

ggplot(sun.spct) +
  geom_line() +
  stat_wb_hbar(w.band = CIE(), size = 1) +
  scale_color_identity() +
  theme_bw()

```

---

stat_wb_irrad	<i>Integrate irradiance for wavebands.</i>
---------------	--

---

### Description

stat\_wb\_irrad integrates the area under a spectral irradiance curve, yielding energy or photon irradiance. The range(s) of wavelengths to integrate are set with a list of waveband objects. **x-scale transformations and axis flipping are currently not supported.**

### Usage

```

stat_wb_irrad(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  time.unit,
  unit.in,
  label.qty = "total",
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  ypos.mult = 1.07,
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)

stat_wb_e_irrad(
  mapping = NULL,
  data = NULL,

```

```

    geom = "text",
    position = "identity",
    ...,
    w.band = NULL,
    time.unit = "second",
    unit.in = "energy",
    label.qty = "total",
    label.mult = 1,
    chroma.type = "CMF",
    label.fmt = "%.3g",
    ypos.mult = 1.07,
    ypos.fixed = NULL,
    na.rm = FALSE,
    show.legend = NA,
    inherit.aes = TRUE
  )

stat_wb_q_irrad(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  time.unit = "second",
  unit.in = "photon",
  label.qty = "total",
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  ypos.mult = 1.07,
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)

```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.

<code>time.unit</code>	character or lubridate::duration
<code>unit.in</code>	character One of "photon","quantum" or "energy"
<code>label.qty</code>	character
<code>label.mult</code>	numeric Scaling factor applied to y-integral values before conversion into character strings.
<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
<code>label.fmt</code>	character string giving a format definition for converting y-integral values into character strings by means of function <code>sprintf</code> .
<code>ypos.mult</code>	numeric Multiplier constant used to scale returned y values.
<code>ypos.fixed</code>	numeric If not NULL used a constant value returned in y.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named integral below is the result of applying `irrad`, `e_irrad` or `q_irrad` to the data.

**y.label** `yeff` multiplied by `label.mult` and formatted according to `label.fmt`

**x** `w.band-midpoint`

**wb.xmin** `w.band` minimum

**wb.xmax** `w.band` maximum

**wb.ymin** `data$y` minimum

**wb.ymax** `data$y` maximum

**wb.yeff** weighted irradiance if `w.band` describes a BSWF

**wb.yint** not weighted irradiance for the range of `w.band`

**wb.xmean** `yint` divided by `wl_expanse(w.band)`

**y** `ypos.fixed` or top of data, adjusted by `ypos.mult`

**wb.color** color of the `w.band`

**wb.name** label of `w.band`

**BW.color** `black_or_white(wb.color)`

**Default aesthetics**

Set by the statistic and available to geoms.

**label** ..y.label..

**x** ..x..

**xmin** ..wb.xmin..

**xmax** ..wb.xmax..

**ymin** ..y.. - (..wb.ymax.. - ..wb.ymin..) \* 0.03

**ymax** ..y.. + (..wb.ymax.. - ..wb.ymin..) \* 0.03

**yintercept** ..wb.ymean..

**fill** ..wb.color..

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**See Also**

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.

# using defaults for energy irradiance in W m-2
ggplot(sun.spct) +
  stat_wb_column(w.band = PAR(), alpha = 0.5) +
  stat_wb_e_irrad(w.band = PAR(), ypos.fixed = 0.32) +
  geom_line() +
  scale_fill_identity() + scale_color_identity()

# using defaults for photon irradiance in umol m-2 s-1
ggplot(sun.spct, unit.out = "photon") +
  stat_wb_column(w.band = PAR(), alpha = 0.5) +
  stat_wb_q_irrad(w.band = PAR(), ypos.fixed = 1.5e-6, label.mult = 1e6) +
  geom_line() +
  scale_fill_identity() + scale_color_identity()

# modify label format and position
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands(), alpha = 0.7) +
  stat_wb_e_irrad(w.band = VIS_bands(),
```

```

      angle = 90, size = 3, hjust = "left",
      label.fmt = "%2.0f~W~m^{-2}", parse = TRUE,
      ypos.fixed = 0.1) +
  geom_line() +
  scale_fill_identity() + scale_color_identity()

# Changing label mapping
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands(), alpha = 0.5) +
  stat_wb_e_irrad(w.band = VIS_bands(),
    label.fmt = "%.2f",
    angle = 90, color = "black", ypos.fixed = 0.1,
    hjust = "left", size = 3,
    mapping = aes(label = after_stat(paste(wb.name, ": ",
      signif(wb.yint, 3),
      sep = "")))) +
  geom_line() +
  scale_fill_identity() + scale_color_identity() +
  theme_bw()

```

---

stat\_wb\_label

*Label ranges under spectral curve.*


---

## Description

stat\_wb\_label computes the center of a waveband. Sets suitable default aesthetics for "text" and "label" geoms displaying "boundaries" and "names" of wavebands. **x-scale transformations and axis flipping are currently not supported.**

## Usage

```

stat_wb_label(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  chroma.type = "CMF",
  label.fmt = "%s",
  ypos.fixed = 0,
  na.rm = TRUE,
  show.legend = NA,
  inherit.aes = TRUE
)

```

**Arguments**

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
label.fmt	character string giving a format definition for formatting the name of the waveband. <a href="#">sprintf</a> .
ypos.fixed	numeric If not NULL used a constant value returned in y.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

**Value**

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

**Computed variables**

<b>x</b>	w.band-midpoint
<b>wb.xmin</b>	w.band minimum
<b>wb.xmax</b>	w.band maximum
<b>y</b>	ypos.fixed or zero
<b>wb.color</b>	color of the w.band
<b>wb.name</b>	label of w.band
<b>wb.label</b>	formatted wb.name

**Default aesthetics**

Set by the statistic and available to geoms.

**label** ..wb.label..

```

x ..x..
xmin ..wb.xmin..
xmax ..wb.xmax..
fill ..wb.color..

```

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

### Note

This stat uses a panel function and ignores grouping as it is meant to be used for annotations.

### See Also

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```

library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS(), ymin = -0.04, ymax = 0,
             color = "black", fill = "white") +
  stat_wb_label(w.band = VIS(), ypos.fixed = -0.02, color = "black")

ggplot(sun.spct) +
  geom_line() +
  stat_wb_hbar(w.band = PAR(), ypos.fixed = 0, size = 1) +
  stat_wb_label(aes(color = ..wb.color..),
               w.band = PAR(), ypos.fixed = +0.025) +
  scale_color_identity()

```

---

stat\_wb\_mean

*Integrate ranges under curve.*

---

### Description

`stat_wb_mean` computes mean spectral irradiance under a curve for each waveband in the input. Sets suitable default aesthetics for "rect", "hline", "vline", "text" and "label" geoms. **x-scale transformations and axis flipping are currently not supported.**

**Usage**

```
stat_wb_mean(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  integral.fun = integrate_xy,
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  ypos.mult = 1.07,
  xpos.fixed = NULL,
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)
```

**Arguments**

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
integral.fun	function on $\$x\$$ and $\$y\$$ .
label.mult	numeric Scaling factor applied to y-integral values before conversion into character strings.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
label.fmt	character string giving a format definition for converting y-integral values into character strings by means of function <a href="#">sprintf</a> .
ypos.mult	numeric Multiplier constant used to scale returned y values.
xpos.fixed, ypos.fixed	numeric If not NULL used as constant value returned in x or y.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.

**inherit.aes** If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. [borders](#).

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named `integral` below is the result of applying `integral.fun`, with default `integrate_xy`.

**y.label** `y.mean` multiplied by `label.mult` and formatted according to `label.fmt`

**x** `w.band`-midpoint

**wb.xmin** `w.band` minimum

**wb.xmax** `w.band` maximum

**wb.ymin** `data$y` minimum

**wb.ymax** `data$y` maximum

**wb.yint** `data$y` integral for the range of `w.band`

**wb.xmean** `yint` divided by `wl_expance(w.band)`

**y** `ypos.fixed` or top of data, adjusted by `ypos.mult`

**wb.color** color of the `w.band`

**wb.name** label of `w.band`

**BW.color** `black_or_white(wb.color)`

### Default aesthetics

Set by the statistic and available to geoms.

**label** `..y.label..`

**x** `..x..`

**xmin** `..wb.xmin..`

**xmax** `..wb.xmax..`

**ymin** 0

**ymax** `..wb.y.mean..`

**yintercept** `..wb.y.mean..`

**fill** `..wb.color..`

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**See Also**

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.

# Using defaults
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands()) +
  stat_wb_mean(w.band = VIS_bands(),
              color = "black") +
  scale_fill_identity() + scale_color_identity()

# Setting format for numbers, position, angle, and color
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands(), alpha = 0.5) +
  stat_wb_mean(w.band = VIS_bands(),
              label.fmt = "%.2f",
              angle = 90, color = "black", ypos.fixed = 0.1) +
  geom_line() +
  scale_fill_identity() + scale_color_identity() +
  theme_bw()

# Changing label mapping
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands(), alpha = 0.5) +
  stat_wb_mean(w.band = VIS_bands(),
              label.fmt = "%.2f",
              angle = 90, color = "black", ypos.fixed = 0.1,
              hjust = "left", size = 3,
              mapping = aes(label = after_stat(paste(wb.name, ": ", y.label, sep = "")))) +
  geom_line() +
  scale_fill_identity() + scale_color_identity() +
  theme_bw()

# example using repulsion
library(ggrepel)
ggplot(sun.spct) +
  geom_line() +
  stat_wb_hbar(w.band = VIS_bands(), size = 1.5) +
  stat_wb_mean(w.band = VIS_bands(),
              geom = "label_repel", nudge_y = +0.04, size = 3,
              segment.colour = NA, label.size = NA) +
  expand_limits(y = 0.9) +
  scale_fill_identity() + scale_color_identity() +
  theme_bw()
```

---

stat\_wb\_relative      *Integrate ranges under spectral curve.*

---

## Description

stat\_wb\_relative computes relative-irradiances under a curve. It first integrates the area under the spectral curve for each waveband in the input, and expresses these irradiances relative to their sum. Sets suitable default aesthetics for "rect", "hline", "vline", "text" and "label" geoms. **x-scale transformations and axis flipping are currently not supported.**

## Usage

```
stat_wb_relative(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  integral.fun = integrate_xy,
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%1.2f",
  ypos.mult = 1.07,
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)
```

## Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
integral.fun	function on $\$x$ and $\$y$ .
label.mult	numeric Scaling factor applied to y-integral values before conversion into character strings.

<code>chroma.type</code>	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <code>chroma_spct</code> object.
<code>label.fmt</code>	character string giving a format definition for converting y-integral values into character strings by means of function <code>sprintf</code> .
<code>ypos.mult</code>	numeric Multiplier constant used to scale returned y values.
<code>ypos.fixed</code>	numeric If not NULL used a constant value returned in y.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named `integral` below is the result of applying `integral.fun` to the data, with default `integrate_xy`.

**y.label** `yint` multiplied by `label.mult` and formatted according to `label.fmt`

**x** `w.band`-midpoint

**wb.xmin** `w.band` minimum

**wb.xmax** `w.band` maximum

**wb.ymin** `data$y` minimum

**wb.ymax** `data$y` maximum

**wb.yint** `data$y` integral for each member of `w.band` / sum of `data$y` integrals for all wavebands in `w.band`

**wb.xmean** `yint` divided by `wl_expance(w.band)`

**y** `ypos.fixed` or top of data, adjusted by `ypos.mult`

**wb.color** color of the `w.band`

**wb.name** label of `w.band`

**BW.color** `black_or_white(wb.color)`

### Default aesthetics

Set by the statistic and available to geoms.

**label** `..y.label..`

**x** `..x..`

```

xmin ..wb.xmin..
xmax ..wb.xmax..
ymin ..y.. - (..wb.ymax.. - ..wb.ymin..) * 0.03
ymax ..y.. + (..wb.ymax.. - ..wb.ymin..) * 0.03
yintercept ..wb.ymean..
fill ..wb.color..

```

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

### See Also

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```

library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS()) +
  stat_wb_relative(w.band = VIS()) +
  scale_fill_identity() + scale_color_identity()

ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS_bands()) +
  stat_wb_relative(w.band = VIS_bands(), angle = 90, size = 2.5) +
  scale_fill_identity() + scale_color_identity()

ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS_bands()) +
  stat_wb_relative(w.band = VIS_bands(), angle = 90, size = 2.5,
                  label.mult = 100, label.fmt = "%3.0f%%") +
  scale_fill_identity() + scale_color_identity()

```

---

stat_wb_sirrad	<i>Integrate spectral irradiance for wavebands.</i>
----------------	---

---

### Description

stat\_wb\_sirrad computes the mean spectral irradiance under a curve, yielding energy or photon spectral irradiance. The range(s) of wavelengths to integrate are set with a list of waveband objects. **x-scale transformations and axis flipping are currently not supported.**

### Usage

```
stat_wb_sirrad(  
  mapping = NULL,  
  data = NULL,  
  geom = "text",  
  position = "identity",  
  ...,  
  w.band = NULL,  
  time.unit,  
  unit.in,  
  label.qty = "mean",  
  label.mult = 1,  
  chroma.type = "CMF",  
  label.fmt = "%.3g",  
  ypos.mult = 0.55,  
  xpos.fixed = NULL,  
  ypos.fixed = NULL,  
  na.rm = FALSE,  
  show.legend = NA,  
  inherit.aes = TRUE  
)
```

```
stat_wb_e_sirrad(  
  mapping = NULL,  
  data = NULL,  
  geom = "text",  
  position = "identity",  
  ...,  
  w.band = NULL,  
  time.unit = "second",  
  unit.in = "energy",  
  label.qty = "mean",  
  label.mult = 1,  
  chroma.type = "CMF",  
  label.fmt = "%.3g",  
  ypos.mult = 0.55,  
  xpos.fixed = NULL,
```

```

ypos.fixed = NULL,
na.rm = FALSE,
show.legend = NA,
inherit.aes = TRUE
)

stat_wb_q_sirrad(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  time.unit = "second",
  unit.in = "photon",
  label.qty = "mean",
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  ypos.mult = 1.07,
  xpos.fixed = NULL,
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)

```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
time.unit	character or <code>lubridate::duration</code>
unit.in	character One of "photon", "quantum" or "energy"
label.qty	character
label.mult	numeric Scaling factor applied to y-integral values before conversion into character strings.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.

<code>label.fmt</code>	character string giving a format definition for converting y-integral values into character strings by means of function <code>sprintf</code> .
<code>ypos.mult</code>	numeric Multiplier constant used to scale returned y values.
<code>xpos.fixed</code> , <code>ypos.fixed</code>	numeric If not NULL used a constant value returned in x or y.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .

### Value

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

### Computed variables

What it is named integral below is the result of applying `irrad`, `e_irrad` or `q_irrad` to the data.

**y.label** `yeff` multiplied by `label.mult` and formatted according to `label.fmt`

**x** `w.band-midpoint`

**wb.xmin** `w.band` minimum

**wb.xmax** `w.band` maximum

**wb.ymin** `data$y` minimum

**wb.ymax** `data$y` maximum

**wb.yeff** weighted irradiance if `w.band` describes a BSWF

**wb.yint** not weighted irradiance for the range of `w.band`

**wb.xmean** `yint` divided by `wl_expanse(w.band)`

**y** `ypos.fixed` or top of data, adjusted by `ypos.mult`

**wb.color** color of the `w.band`

**wb.name** label of `w.band`

**BW.color** `black_or_white(wb.color)`

### Default aesthetics

Set by the statistic and available to geoms.

**label** `..y.label..`

**x** `..x..`

**xmin** `..wb.xmin..`

**xmax** `..wb.xmax..`

```

ymin 0
ymax ..wb.ymean..
yintercept ..wb.ymean..
fill ..wb.color..

```

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

### See Also

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```

library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  stat_wb_column(w.band = VIS_bands()) +
  stat_wb_e_sirrad(w.band = VIS_bands(), angle = 90, size = 4,
                  label.fmt = "%1.2f", ypos.fixed = 0.1) +
  geom_line() +
  scale_fill_identity() + scale_color_identity()

ggplot(sun.spct, unit.out = "photon") +
  geom_line() +
  stat_wb_hbar(w.band = PAR(), size = 1) +
  stat_wb_q_sirrad(aes(color = ..wb.color..),
                  w.band = PAR(), label.fmt = "mean = %.3g",
                  ypos.mult = 1, xpos.fixed = 390, hjust = 1) +
  scale_color_identity()

```

---

stat\_wb\_total

*Integrate ranges under spectral curve.*

---

### Description

`stat_wb_total` computes integral under a curve. Sets suitable default aesthetics for "rect", "hline", "vline", "text" and "label" geoms displaying "totals" per waveband. **x-scale transformations and axis flipping are currently not supported.**

**Usage**

```

stat_wb_total(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  w.band = NULL,
  integral.fun = integrate_xy,
  label.mult = 1,
  chroma.type = "CMF",
  label.fmt = "%.3g",
  ypos.mult = 1.07,
  ypos.fixed = NULL,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)

```

**Arguments**

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	a waveband object or a list of waveband objects or numeric vector of at least length two.
integral.fun	function on $x$ and $y$ .
label.mult	numeric Scaling factor applied to y-integral values before conversion into character strings.
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
label.fmt	character string giving a format definition for converting y-integral values into character strings by means of function <a href="#">sprintf</a> .
ypos.mult	numeric Multiplier constant used to scale returned y values.
ypos.fixed	numeric If not NULL used a constant value returned in y.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

**Value**

A data frame with one row for each waveband object in the argument to `w.band`. Waveband outside the range of the spectral data are trimmed or discarded.

**Computed variables**

What it is named `integral` below is the result of applying `integral.fun`, with default `integrate_xy`.

**y.label** `y.mean` multiplied by `label.mult` and formatted according to `label.fmt`

**x** `w.band`-midpoint

**wb.xmin** `w.band` minimum

**wb.xmax** `w.band` maximum

**wb.ymin** `data$y` minimum

**wb.ymax** `data$y` maximum

**wb.yint** `data$y` integral for the range of `w.band`

**wb.xmean** `yint` divided by `wl_expance(w.band)`

**y** `ypos.fixed` or top of data, adjusted by `ypos.mult`

**wb.color** color of the `w.band`

**wb.name** label of `w.band`

**BW.color** `black_or_white(wb.color)`

**Default aesthetics**

Set by the statistic and available to geoms.

**label** `..y.label..`

**x** `..x..`

**xmin** `..wb.xmin..`

**xmax** `..wb.xmax..`

**ymin** `..y.. - (..wb.ymax.. - ..wb.ymin..) * 0.03`

**ymax** `..y.. + (..wb.ymax.. - ..wb.ymin..) * 0.03`

**yintercept** `..wb.ymean..`

**fill** `..wb.color..`

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**See Also**

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wl\\_strip\(\)](#), [stat\\_wl\\_summary\(\)](#)

**Examples**

```
library(photobiologyWavebands)
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = VIS()) +
  stat_wb_total(w.band = VIS()) +
  scale_fill_identity() + scale_color_identity()

ggplot(sun.spct) +
  geom_line() +
  stat_wb_box(w.band = UV_bands(), color = "white") +
  stat_wb_total(w.band = UV_bands()) +
  scale_fill_identity() + scale_color_identity()
```

---

stat\_wl\_strip

*Calculate colours from wavelength.*


---

**Description**

stat\_wl\_strip computes color definitions according to human vision and by default plots a narrow, guide-like colour gradient strip based on wavelength. **x-scale transformations and axis flipping are currently not supported.**

**Usage**

```
stat_wl_strip(
  mapping = NULL,
  data = NULL,
  geom = "rect",
  position = "identity",
  ...,
  w.band = NULL,
  length.out = 150,
  chroma.type = "CMF",
  na.rm = TRUE,
  show.legend = FALSE,
  inherit.aes = TRUE
)
```

```

wl_guide(
  mapping = NULL,
  data = NULL,
  position = "identity",
  ...,
  chroma.type = "CMF",
  w.band = NULL,
  length.out = 150,
  ymin = -Inf,
  ymax = Inf,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE
)

```

### Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data.
position	The position adjustment to use for overlapping points on this layer.
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
w.band	waveband object or a list of such objects or NULL.
length.out	The number of steps to use to simulate a continuous range of colours when <code>w.band == NULL</code> .
chroma.type	character one of "CMF" (color matching function) or "CC" (color coordinates) or a <a href="#">chroma_spct</a> object.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .
ymin, ymax	numeric used as aesthetics for plotting the guide.

### Value

generic\_spect object with new x values plus other computed variables described below.

### Computed variables

**x**  $(w.low + wl.high) / 2$

**wl.low** boundary of waveband

**wl.high** boundary of waveband  
**wl.color** color corresponding to wavelength  
**wb.color** color corresponding to waveband  
**wb.name** label of w.band

### Default aesthetics

Set by the statistic and available to geoms.

**x** ..x..  
**label** as.character(..wb.f..)
   
**xmin** ..wl.low..
   
**xmax** ..wl.high..
   
**fill** ..wb.color..

### Required aesthetics

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

### Note

This stat uses a panel function and ignores grouping as it is meant to be used for annotations.

### See Also

`color_of`, which is used internally.

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_summary\(\)](#)

### Examples

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) +
  geom_line() +
  stat_wl_strip(ymax = -0.02, ymin = -0.04) +
  scale_fill_identity()

# on some graphic devices the output may show spurious vertical lines
ggplot(sun.spct) +
  wl_guide(alpha = 0.33, color = NA) +
  geom_line()
```

---

stat_wl_summary	<i>Average area under curve for regions.</i>
-----------------	--

---

## Description

stat\_wl\_summary computes the area under a curve.

## Usage

```
stat_wl_summary(
  mapping = NULL,
  data = NULL,
  geom = "text",
  position = "identity",
  ...,
  range = NULL,
  integral.fun = integrate_xy,
  label.fmt = "%.3g",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)
```

## Arguments

mapping	The aesthetic mapping, usually constructed with <a href="#">aes</a> or <a href="#">aes_</a> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
geom	The geometric object to use display the data
position	The position adjustment to use for overlapping points on this layer
...	other arguments passed on to <a href="#">layer</a> . This can include aesthetics whose values you want to set, not map. See <a href="#">layer</a> for more details.
range	a numeric vector of at least length two.
integral.fun	function on $\$x\$$ and $\$y\$$ .
label.fmt	character string giving a format definition for converting y-integral values into character strings by means of function <a href="#">sprintf</a> .
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

**Value**

A data frame with one row.

**Computed variables**

What it is named `integral` below is the result of applying `integral.fun`, with default `integrate_xy`.

**y.label** `y` formatted according to `label.fmt`

**x** range-midpoint

**wb.xmin** range minimum

**wb.xmax** range maximum

**y** `data$y` integral for the range by the expanse of the range

**Default aesthetics**

Set by the statistic and available to geoms.

**label** `..label..`

**x** `..x..`

**xmin** `..wb.xmin..`

**xmax** `..wb.xmax..`

**y** `..y..`

**ymin** `0`

**ymax** `..y..`

**yintercept** `..y..`

**Required aesthetics**

Required by the statistic and need to be set with `aes()`.

**x** numeric, wavelength in nanometres

**y** numeric, a spectral quantity

**See Also**

Other stats functions: [stat\\_color\(\)](#), [stat\\_find\\_qtys\(\)](#), [stat\\_find\\_wls\(\)](#), [stat\\_label\\_peaks\(\)](#), [stat\\_peaks\(\)](#), [stat\\_spikes\(\)](#), [stat\\_wb\\_box\(\)](#), [stat\\_wb\\_column\(\)](#), [stat\\_wb\\_contribution\(\)](#), [stat\\_wb\\_hbar\(\)](#), [stat\\_wb\\_irrad\(\)](#), [stat\\_wb\\_label\(\)](#), [stat\\_wb\\_mean\(\)](#), [stat\\_wb\\_relative\(\)](#), [stat\\_wb\\_sirrad\(\)](#), [stat\\_wb\\_total\(\)](#), [stat\\_wl\\_strip\(\)](#)

**Examples**

```
# ggplot() methods for spectral objects set a default mapping for x and y.
ggplot(sun.spct) + geom_line() +
  stat_wl_summary(geom = "hline")
ggplot(sun.spct) + geom_line() +
  stat_wl_summary(label.fmt = "mean = %.3f", color = "red", vjust = -0.3) +
  stat_wl_summary(geom = "hline", color = "red")
```

---

Tfr\_label                      *Transmittance axis labels*

---

### Description

Generate cps axis labels in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

### Usage

```
Tfr_label(
  unit.exponent = ifelse(pc.out, -2, 0),
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE),
  pc.out = getOption("ggspectra.pc.out", default = FALSE),
  Tfr.type
)

Tfr_internal_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)

Tfr_total_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = NULL,
  scaled = FALSE,
  normalized = FALSE,
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

### Arguments

unit.exponent	integer
format	character string, "R", "R.expresion", "R.character", or "LaTeX".
label.text	character Textual portion of the labels.
scaled	logical If TRUE relative units are assumed.
normalized	logical (FALSE) or numeric Normalization wavelength in manometers (nm).

axis.symbols	logical If TRUE symbols of the quantities are added to the name. Supported only by format = "R.expression".
pc.out	logical, if TRUE use percent as default instead of fraction of one.
Tfr.type	character, either "total" or "internal".

**Value**

a character string or an R expression.

**Note**

Default for label.text depends on the value passed as argument to Tfr.type.

**Examples**

```
Tfr_label(Tfr.type = "internal")
Tfr_label(Tfr.type = "total")
Tfr_label(Tfr.type = "internal", axis.symbols = FALSE)

Tfr_internal_label()
Tfr_internal_label(format = "R.expression", axis.symbols = FALSE)
Tfr_internal_label(-2)
Tfr_internal_label(-3)
Tfr_internal_label(format = "R.expression")
Tfr_internal_label(format = "LaTeX")
Tfr_internal_label(-3, format = "LaTeX")

Tfr_total_label()
Tfr_total_label(format = "R.expression", axis.symbols = FALSE)
Tfr_total_label(-2)
Tfr_total_label(-3)
Tfr_total_label(format = "R.expression")
Tfr_total_label(format = "LaTeX")
Tfr_total_label(-3, format = "LaTeX")
```

---

w\_length\_label

*Wave- axis labels*


---

**Description**

Generate wavelength, wavenumber, wave frequency, and energy per photon axis labels in SI units, using SI scale factors. Output can be selected as character, expression (R default devices) or LaTeX (for tikz device).

**Usage**

```
w_length_label(
  unit.exponent = -9,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["w.length"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
w_number_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["w.number"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
w_frequency_label(
  unit.exponent = 9,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["freq"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
w_energy_eV_label(
  unit.exponent = 0,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["energy"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

```
w_energy_J_label(
  unit.exponent = -18,
  format = getOption("photobiology.math", default = "R.expression"),
  label.text = axis_labels()[["energy"]],
  axis.symbols = getOption("ggspectra.axis.symbols", default = TRUE)
)
```

**Arguments**

<code>unit.exponent</code>	integer The exponent in base 10 of the scale multiplier to use.
<code>format</code>	character string, "R", "R.expression", "R.character", or "LaTeX".
<code>label.text</code>	character Textual portion of the labels.
<code>axis.symbols</code>	logical If TRUE symbols of the quantities are added to the name. Supported only by <code>format = "R.expression"</code> .

**Details**

By default labels consist in a textual name for the quantity, a symbol separated by a comma and units with scale factor in parenthesis. The textual names are by default in English but this default can be

overridden for example with translations to a different language. To change or translate the default texts please see [axis\\_labels\\_uk](#). The markup language used for the labels can be selected through a parameter argument, with character strings ready to be parsed into R expressions as default.

Wavelengths are assumed to be expressed in nanometres in the data. The `unit.exponent` corresponds to that desired for the tick labels with the corresponding axis label automatically set to an SI scale factor if possible, and otherwise shown as a power of 10.

These functions are used internally by *x* scales; see [sec\\_axis\\_w\\_number](#) and [scale\\_x\\_wl\\_continuous](#). The scales and secondary axis functions should be used except when defining new scale functions.

### Value

a character string or an R expression.

### Examples

```
w_length_label()
w_length_label(axis.symbols = FALSE)
w_length_label(format = "R.expression")
w_length_label(format = "LaTeX")
w_number_label()
w_number_label(format = "R.expression")
w_frequency_label()
w_frequency_label(format = "R.expression")
w_energy_J_label()
w_energy_eV_label()
```

---

w\_number

*Deprecated functions*

---

### Description

To convert wavelength into wavenumber or into frequency, please, use the conversion functions from package 'photobiology' in place of the deprecated functions `w_number()` and `w_frequency()` from this package.

### Usage

```
w_number(w.length, unit.exponent = 0)

w_frequency(w.length, unit.exponent = 0)
```

### Arguments

`w.length`        numeric wavelength (nm)  
`unit.exponent`   integer Exponent of the scale multiplier implicit in result, e.g., use 3 for kJ.

**Deprecated**

These functions will be removed from package 'ggpmisc' in the near future.

**See Also**

See [w12wavenumber](#) for the functions to be used in all new code.

**Examples**

```
library(photobiology)
```

```
w12wavenumber(600)
```

```
w12frequency(600)
```

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