

# photobiologyWavebands Version 0.3.4

## User Guide

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January 2, 2016

## 1 Introduction

We have developed a set of packages to facilitate the calculation of many different quantities that can be derived from spectral irradiance data. The basic package is called `photobiology`, and the package described here is an extension of the basic facilities for quantification of ultraviolet radiation. It will be submitted to CRAN (Comprehensive R archive network), it is meanwhile available from <https://bitbucket.org/aphalo/photobiology/downloads> and <https://bitbucket.org/aphalo/photobiologyuv/downloads>. There are also a public Git repositories at <https://bitbucket.org/aphalo/photobiology> and <https://bitbucket.org/aphalo/photobiologyuv>. Functions are included for calculating weighted and unweighted UV doses, irradiances and related quantities.

## 2 Calculating irradiances

Functions for several colour bands, in some cases according to different optional definitions, are listed in Table 1.

A new waveband definition can be created with `new_waveband()`<sup>1</sup>. When using data stored in vectors integrated irradiance or dose is calculated with functions `energy_irradiance()` and `photon_irradiance`, which take as one argument the waveband descriptor list as an argument. When working with data stored in `source_spct` objects functions and `e_irrad` and `q_irrad` are used instead.

The functions used for calculating the irradiances from vectors have additional arguments, that permit to indicate the type of scale used for the input spectrum ("photon" or "energy"). In the case of `irradiance` and `irrad`, output units ("photon" or "energy") can also be supplied.

An example using `sun_spct` included in package `photobiology`. As the input spectral irradiance is units of  $\text{W m}^{-2} \text{nm}^{-1}$  the output is in  $\text{mol m}^{-2} \text{s}^{-1}$

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<sup>1</sup>See package `photobiology` and its documentation.

Table 1: Functions in R package photobiologyWavebands used for constructing descriptors of wavebands used for calculation of irradiances or exposures. The boundaries of the band given as wavelengths in nm ( $\lambda$ ). Definition according to ISO-21348 "ISO" is the default for all functions except PAR() for which there is only one definition in common use "Plant" which is the default.

Waveband	Source	Dose or irrad.	Waveband (nm)
UV	ISO-21348	UV("ISO")	$100 \leq \lambda < 400$
UV-C	ISO-21348	UVC("ISO")	$100 \leq \lambda < 280$
UV-C	n.a.	UVC("medical")	$220 \leq \lambda < 290$
UV-C	n.a.	UVC("none")	$200 \leq \lambda < 280$
UV-B	ISO-21348	UVB("ISO")	$280 \leq \lambda < 315$
UV-B	n.a.	UVB("none")	$280 \leq \lambda < 320$
UV-A	ISO-21348	UVA("ISO")	$315 \leq \lambda < 400$
UV-A	n.a.	UVA("none")	$320 \leq \lambda < 400$
Visible	ISO-21348	VIS("ISO")	$380 \leq \lambda < 760$
Photosynthesis	n.a.	PAR("Plant")	$400 \leq \lambda < 700$
Purple	ISO-21348	Purple("ISO")	$360 \leq \lambda < 450$
Blue	Sellaro	Blue("Sellaro")	$420 \leq \lambda < 490$
Blue	ISO-21348	Blue("ISO")	$450 \leq \lambda < 500$
Green	Sellaro	Green("Sellaro")	$500 \leq \lambda < 570$
Green	ISO-21348	Green("ISO")	$500 \leq \lambda < 570$
Yellow	ISO-21348	Yellow("ISO")	$570 \leq \lambda < 591$
Orange	ISO-21348	Orange("ISO")	$591 \leq \lambda < 610$
Red	ISO-21348	Red("ISO")	$610 \leq \lambda < 760$
Red	Smith	Red("Smith10")	$655 \leq \lambda < 665$
Red	Smith ?	Red("Smith20")	$650 \leq \lambda < 670$
Red	Inada	Red("Inada")	$600 \leq \lambda < 700$
Red	Warrington	Red("Warrington")	$625 \leq \lambda < 675$
Red	Sellaro	Red("Sellaro")	$620 \leq \lambda < 680$
Far-red	ISO	Far_red("ISO")	not defined
Far-red	Smith10	Far_red("Smith")	$725 \leq \lambda < 735$
Far-red	Smith20	Far_red("Smith")	$720 \leq \lambda < 740$
Far-Red	Inada	Red("Inada")	$700 \leq \lambda < 800$
Far-Red	Warrington	Red("Warrington")	$700 \leq \lambda < 850$
Far-red	Sellaro	Far_red("Sellaro")	$700 \leq \lambda < 750$
Far-red	BT	Far_red("BT")	$700 \leq \lambda < 760$
Arbitrary	n.a.	new_waveband(lo,hi)	$lo \leq \lambda < hi$

or  $\text{W m}^{-2}$ . We multiply by  $10^6$  to obtain photon irradiance expressed in  $\mu\text{mol m}^{-2} \text{s}^{-1}$ .

```
e_irrad(sun.spct, PAR()) # W m-2

##      PAR
## 196.6343
## attr(,"time.unit")
## [1] "second"
## attr(,"radiation.unit")
## [1] "energy irradiance total"

q_irrad(sun.spct, PAR()) * 1e6 # umol s-1 m-2

##      PAR
## 894.1352
## attr(,"time.unit")
## [1] "second"
## attr(,"radiation.unit")
## [1] "photon irradiance total"
```

To integrate the whole spectrum, without selecting a waveband or applying any weighting function, we simply omit the `waveband` in the function call.

```
e_irrad(sun.spct)

##      Total
## 269.1249
## attr(,"time.unit")
## [1] "second"
## attr(,"radiation.unit")
## [1] "energy irradiance total"
```

It is also very easy to define your own waveband as described in the photobiology manual. Here we give a very simple example.

```
e_irrad(sun.spct, PAR())

##      PAR
## 196.6343
## attr(,"time.unit")
## [1] "second"
## attr(,"radiation.unit")
## [1] "energy irradiance total"

e_irrad(sun.spct, waveband(c(400, 700))) # Same as PAR()

##  range.400.700
##          196.6343
## attr(,"time.unit")
## [1] "second"
## attr(,"radiation.unit")
## [1] "energy irradiance total"
```

Irradiances for different wavebands can be grouped into a list of any length. If the list has named members, then these names are used instead of the default ones.

```

e_irrad(sun.spct, list(Blue(), VIS()))

## Blue.ISO    VIS.ISO
## 37.55207 231.86345
## attr(),"time.unit"
## [1] "second"
## attr(),"radiation.unit"
## [1] "energy irradiance total"

e_irrad(sun.spct, list(B = Blue(), VIS()))

## Blue.ISO    VIS.ISO
## 37.55207 231.86345
## attr(),"time.unit"
## [1] "second"
## attr(),"radiation.unit"
## [1] "energy irradiance total"

```

A few functions for generating coherent lists of wavebands are also defined (Table 2).

```

e_irrad(sun.spct, VIS_bands())

## Purple.ISO    Blue.ISO    Green.ISO    Yellow.ISO
## 47.75529    37.55207    49.26860    13.67971
## Orange.ISO   Red.ISO
## 12.00432    79.38159
## attr(),"time.unit"
## [1] "second"
## attr(),"radiation.unit"
## [1] "energy irradiance total"

```

Function `irrad` behaves either as `e_irrad` or `q_irrad` depending on a global option, and in some circumstances this may help when needing to switch between bases of expression. By default energy irradiance are calculated.

```

e_irrad(sun.spct)

## Total
## 269.1249
## attr(),"time.unit"
## [1] "second"
## attr(),"radiation.unit"
## [1] "energy irradiance total"

q_irrad(sun.spct)

## Total
## 0.001255336
## attr(),"time.unit"
## [1] "second"
## attr(),"radiation.unit"
## [1] "photon irradiance total"

irrad(sun.spct, PAR())

```

Table 2: Functions in R package photobiologyWavebands used for constructing lists descriptors of wavebands used for calculation of irradiances or exposures.

Waveband	Source	Function	std values used
VIS defs.	ISO-21348	VIS_bands("ISO")	"ISO"
UV defs.	ISO-21348	UV_bands("ISO")	"ISO"
UV defs.	n.a.	UV_bands("none")	"none"
Plant sens.	n.a.	Plant_bands("sensory20")	"ISO", "Sellaro" and "Smith20"
Plant sens.	n.a.	Plant_bands("sensory")	"ISO", "Sellaro" and "Smith20"
Plant sens.	n.a.	Plant_bands("sensory10")	"ISO", "Sellaro" and "Smith10"
Plant sens.	n.a.	Plant_bands("energy")	"ISO" and "McCree"

```
##      PAR
## 196.6343
## attr(*,"time.unit")
## [1] "second"
## attr(*,"radiation.unit")
## [1] "energy irradiance total"
```

Low level functions, allow doing the same calculations using numeric vectors with arbitrary names, with a more complicated syntax and weaker error diagnosis. These functions do also allow more detailed user control of speed optimizations.

```
with(sun.spct,
     photon_irradiance(w.length, s.e.irrad, PAR())
   )

##      PAR
## 0.0008941352
```

### 3 Calculating photon ratios

Photon ratios can be calculated from any pair of waveband objects. This a convenient and very flexible way of doing this type of calculations.

```
q_ratio(sun.spct, Blue(), VIS())

##  Blue.ISO: VIS.ISO(q:q)
##            0.1371157
## attr(*,"radiation.unit")
## [1] "q:q ratio"
```

Although not so frequently used, enery ratios can be also calculated (please see the documentation of packae photobiology for a description of the available functions.

```
e_ratio(sun.spct, Blue(), VIS())

##  Blue.ISO: VIS.ISO(q:q)
##                0.1371157
## attr(,"radiation.unit")
## [1] "q:q ratio"
```

Photon ratios can be calculated from pairs of lists of waveband objects. As can be seen in the example recycling applies.

```
q_ratio(sun.spct, VIS_bands(), VIS())

##  Purple.ISO: VIS.ISO(q:q)      Blue.ISO: VIS.ISO(q:q)
##                0.15087813          0.13711571
##  Green.ISO: VIS.ISO(q:q)      Yellow.ISO: VIS.ISO(q:q)
##                0.20259364          0.06106049
##  Orange.ISO: VIS.ISO(q:q)     Red.ISO: VIS.ISO(q:q)
##                0.05545498          0.41504754
## attr(,"radiation.unit")
## [1] "q:q ratio"
```

Low-level functions, including convenience functions are provided for calculating photon ratios from data stored in numeric vectors. These functions are listed in Table 3. We follow the most frequently used wavelength ranges for the different colours, but also provide generic functions that can be used when other limits are needed. Here we calculate Blue:PAR photon ratio.

```
with(sun.spct,
  B_PAR_ratio(w.length, s.e.irrad)
)

## [1] 0.2117127
```

Please, be aware that following common practice in the literature, the wavelength range used for red light is different for the different photon ratios.

## 4 Calculating effective irradiances and exposures

The waveband definitions and SWFs are stored in `waveband` objects, that can be created with functions `waveband` or `new_waveband()`. The same functions described above for unweighted irradiances are used to calculate effective irradiances and doses.

Currently functions for constructing `waveband` objects describing several BSWFs are implemented (see Table 4). These functions take three arguments in most cases as they have been used and continue to be used inconsistently in the scientific literature. By supplying these arguments different variations of the BSWFs can be obtained. The defaults used are those values which we consider best, usually the most frequently used ones, except in cases when we consider the use of those values problematic for the reliability of the calculations.

Table 3: Functions in R package photobiologyWavebands for calculation of photon ratios from spectra in spectral energy units. `w.length` = vector of wavelengths (nm); `s.e.irrad` = vector of spectral energy irradiances.

Ratio	R function	wavelength ranges (nm)
UV:PAR	UV_PAR_ratio(w.length, s.e.irrad)	$100 \leq \lambda < 400, 400 \leq \lambda < 700$
UV-C:PAR	UVC_PAR_ratio(w.length, s.e.irrad)	$100 \leq \lambda < 280, 400 \leq \lambda < 700$
UV-B:PAR	UVB_PAR_ratio(w.length, s.e.irrad)	$280 \leq \lambda < 315, 400 \leq \lambda < 700$
UV-A:PAR	UVA_PAR_ratio(w.length, s.e.irrad)	$315 \leq \lambda < 400, 400 \leq \lambda < 700$
Blue:PAR	B_PAR_ratio(w.length, s.e.irrad)	$420 \leq \lambda < 490, 400 \leq \lambda < 700$
Green:PAR	G_PAR_ratio(w.length, s.e.irrad)	$500 \leq \lambda < 570, 400 \leq \lambda < 700$
Red:Far-red	R_FR_ratio(w.length, s.e.irrad)	$650 \leq \lambda < 670, 720 \leq \lambda < 740$
Blue:Green	B_G_ratio(w.length, s.e.irrad)	$420 \leq \lambda < 490, 500 \leq \lambda < 570$

```
e_irrad(sun.spct, CIE())
##   CIE98.298.tr.lo
##   0.08181583
## attr(,"time.unit")
## [1] "second"
## attr(,"radiation.unit")
## [1] "energy irradiance total"
```

## 5 Calculating an action spectrum at given wavelengths

The functions available for calculating action spectra take as argument a vector of wavelengths, and return a vector of effectiveness (either quatum/photon or energy based) depending on how the original source describes them. These functions are listed in Table 5, and an example of their use follows. In these examples we generate the wavelengths vectors in R, but they can be also read from a file.

```
# at 1 nm intervals
wavelengths1 <- 285:400
action.spectrum1 <- CIE_e_fun(wavelengths1)
# plot(wavelengths1, action.spectrum1, type = "p")
```

All functions accept a wavelengths vector with variable and arbitrary step sizes, with the condition that the wavelengths are sorted in strictly increasing order.

In practice these functions are mainly used internally by the package, and very rarely in user code, as the same output can be obtained by multiplication of `source_spct` objects by `waveband` objects.

Table 4: Functions in R package `photobiologyWavebands` used for constructing `waveband` objects describing BSWFs used for calculation of effective irradiances or doses. The functions for BSWFs available in this package, are by default as in the original source. Optionally they can be normalized to any wavelength within their non-zero range by providing the `norm` argument with a wavelength in nm. The range of wavelengths included when calculating integrals is given by `w.low` and `w.high`. The values in the table below are the defaults.

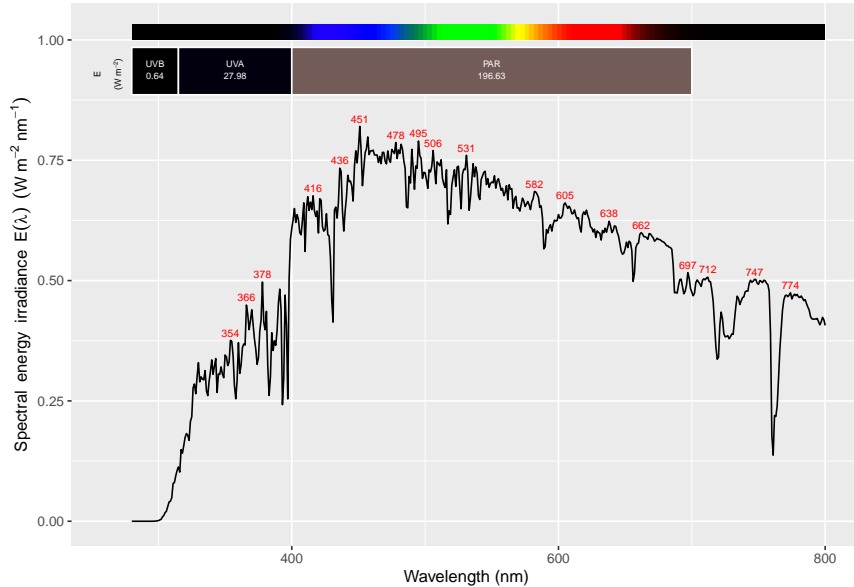
Action spectrum	Formulation	Constructor function	norm $\lambda$ (nm)	w.low $\lambda$ (nm)	w.high $\lambda$ (nm)
Gen. plant action	Green	GEN.G(norm, w.low, w.high)	300	275	313.3
Gen. plant action	Thimijan	GEN.T(norm, w.low, w.high)	300	275	345.0
Gen. plant action	Michelletti	GEN.M(norm, w.low, w.high)	300	275	313.3
Plant growth	Flint & Caldwell	PG.(norm, w.low, w.high)	300	275	390.0
Erythemal	CIE98	CIE.(norm, w.low, w.high)	298	250	400.0
ICNIRP	ICNIRP2004	ICNIRP.(norm, w.low, w.high)	270	210	400.0
'Naked' DNA	TUV, from Setlow	DNA.N(norm, w.low, w.high)	300	250	400.0
'Naked' DNA	Green & Miller	DNA.GM(norm, w.low, w.high)	300	250	400.0
'Plant' DNA	Musil, from Quaite	DNA.P(norm, w.low, w.high)	300	250	400.0
Flavonoid	Ibdah	FLAV.(norm, w.low, w.high)	300	275	346.0

Table 5: Biological spectral weighting functions predefined in R package `photobiologyWavebands`. The functions for BSWFs available in this package, implement the functions as defined in the original publications. When the original ‘definition’ is available as tabulated data, or we have tabulated it by digitizing a figure, the values returned are calculated by spline interpolation.

Action spectrum	Formulation	Function	Norm. $\lambda$ (nm)
Gen. plant action	Green	GEN_G_q_fun(w.length)	280
Gen. plant action	Thimijan	GEN_T_q_fun(w.length)	300
Gen. plant action	Micheletti	GEN_M_q_fun(w.length)	300
Plant growth	Flint & Caldwell	PG_q_fun(w.length)	300
Erythemal	CIE98	CIE_e_fun(w.length)	298
ICNIRP	ICNIRP2004	ICNIRP_e_fun(w.length)	270
‘Naked’ DNA	TUV, from Setlow	DNA_N_q_fun(w.length)	n.a.
‘Naked’ DNA	Green & Miller	DNA_GM_q_fun(w.length)	n.a.
‘Plant’ DNA	Musil, from Quaite	DNA_P_q_fun(w.length)	290
Flavonoid	Ibdah	FLAV_q_fun(w.length)	300

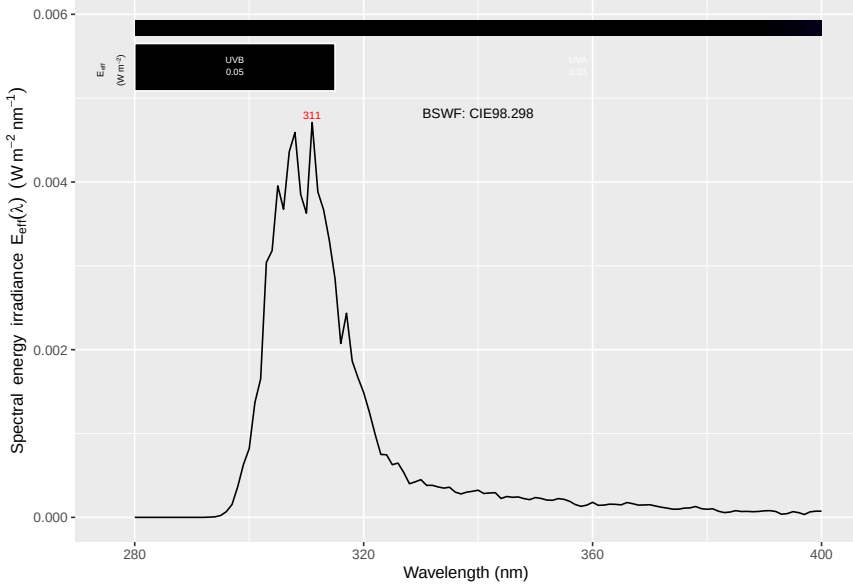
Compare the following two plots,

```
plot(sun.spct)
```



```
plot(sun.spct * CIE())
```

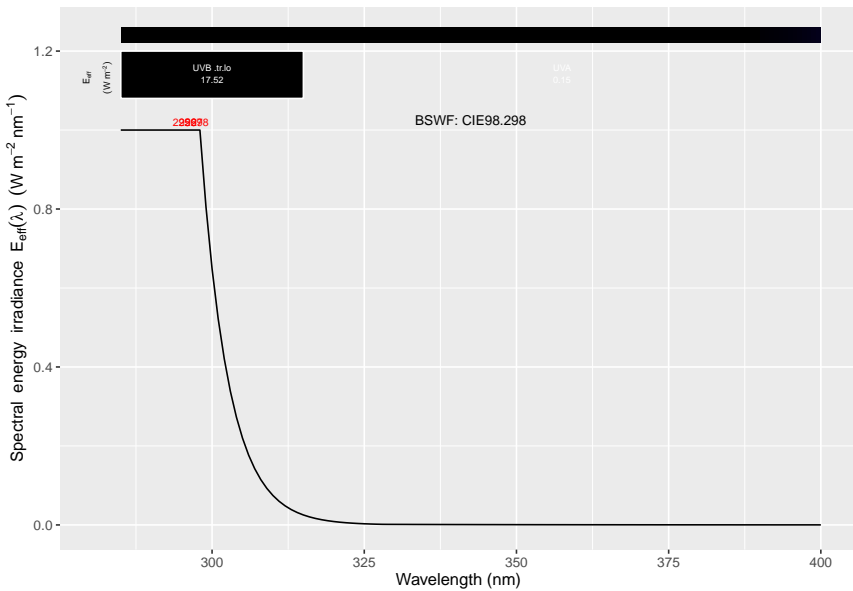
```
## Warning in trim.tails(data$x, data$y, low.limit = range[1], high.limit =
range[2]): Ignoring high.limit as it is too high.
## Warning: Removed 1 rows containing missing values
## (geom_rect).
```



One can use operators and by multiplying spectral irardiance equal to one by a waveband obtain a plot,

```
plot(source_spct(285:400, 1) * CIE())

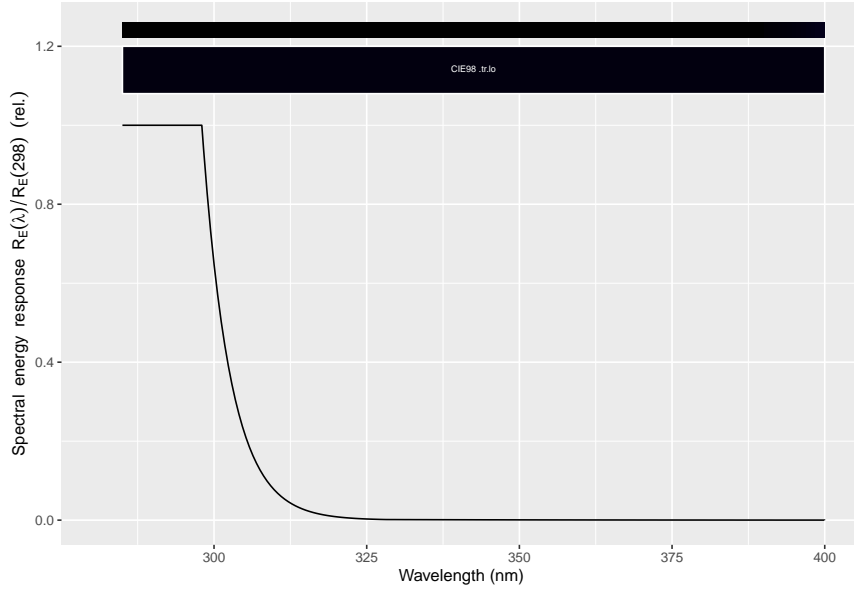
## Warning in trim.tails(data$x, data$y, low.limit = range[1], high.limit =
range[2]): Ignoring high.limit as it is too high.
## Warning: Removed 1 rows containing missing values
## (geom_rect).
```



However, it is simpler and clearer to directly plot the wavband.

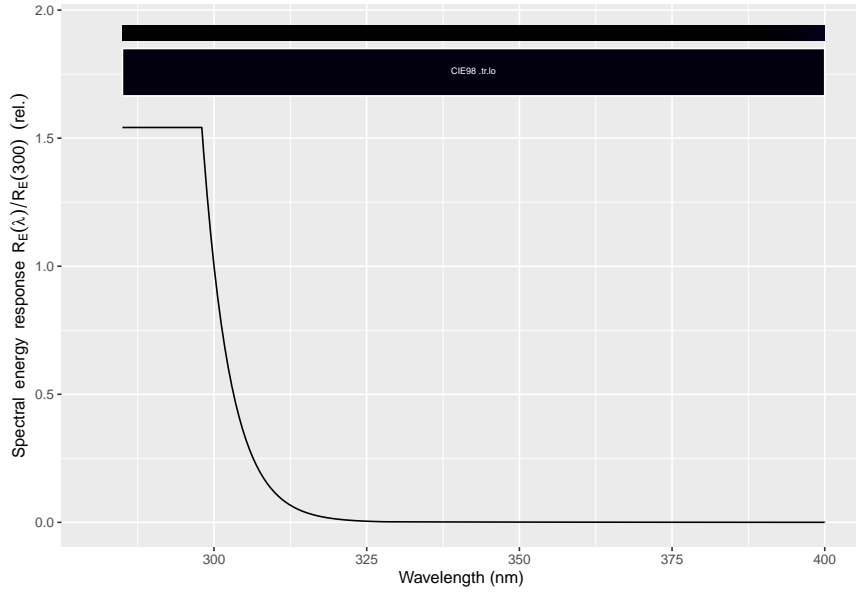
```
plot(CIE(), range = c(285,400))

## Warning in f(...): BSWFs not yet supported: skipping summary for 'CIE98
.tr.lo'.
```



```
plot(CIE(), range = c(285,400), norm = 300)

## Warning in f(...): BSWFs not yet supported: skipping summary for 'CIE98
.tr.lo'.
```



## 6 Luminous flux

The luminous flux per unit area in lux can be calculated as follows using the original luminous efficiency function for the human eye using for defining the lumen. As we start with spectral irradiance we obtain luminous flux per unit area expressed in lux.

```
e_response(sun.spct * CIE1924_lef.spct) * photopic_sensitivity

##      Total
## 49579.93
## attr(),"time.unit")
## [1] "second"
## attr(),"radiation.unit")
## [1] "energy response total"
```

The luminous flux per unit area in lux can be calculated as follows using the latest luminous efficiency function for the human eye.

```
e_response(sun.spct * CIE2008_lef2deg.spct) * photopic_sensitivity

##      Total
## 53057.78
## attr(),"time.unit")
## [1] "second"
## attr(),"radiation.unit")
## [1] "energy response total"
```

As the luminous efficiency functions vary slightly in the wavelength at which the maximum is located, and the wavelength used for the sensitivity constant is fixed by the definition of the Lumen, a small correction is need for exact results.

```
e_response(sun.spct * CIE2008_lef2deg.spct) * photopic_sensitivity *
    interpolate_spct(CIE2008_lef2deg.spct, 555)$s.e.response

##     Total
## 53910.01
## attr("time.unit")
## [1] "second"
## attr("radiation.unit")
## [1] "energy response total"
```

An equivalent quantity can be calculated for scotopic vision, using the corresponding function and constant.

```
moon.spct <- sun.spct / 1000 # a placeholder for now!
e_response(moon.spct * CIE1951_scotopic_lef.spct) * scotopic_sensitivity

##     Total
## 118.6256
## attr("time.unit")
## [1] "second"
## attr("radiation.unit")
## [1] "energy response total"
```