photobiology Version 0.9.2 Upgrade Guide

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1 Changes in versions 0.6.0 and higher

I have renamed classes and functions as I have learnt in the hard way that using dots is S3 class names and methods causes lots of problems. The name changes will break old code and stored/saved spectral objects need to be *upgraded* before they can be used with the current version of the suite. Of course, they continue to be valid data.table and data.frame objects.

Several frequently used classes like data.frame and data.table include a dot in their name, but as Hadley Wickham emphasizes in his new book R packages it is preferable to consistently use underscores instead of dots.

Below we describe new naming rules used, and the resulting changes in class, method and function names compared to versions previous to 0.6.0.

Class names include only letters, or letter plus underscore characters, leading to these name changes:

```
generic.spct → generic_spct
source.spct → source_spct
response.spct → response_spct
filter.spct → filter_spct
reflector.spct → reflector_spct
object.spct → object_spct
chroma.spct → chroma_spct
Constructors have been renamed accordingly:
generic.spct() → generic_spct()
source.spct() → source_spct()
etc.
```

The class-test and copy-and-set class functions were renamed to match, but following 'normal' R style the first dot, following is or is, has been maintained:

```
\begin{split} &\texttt{is.any.spct()} \rightarrow \texttt{is.any.spct()} \\ &\texttt{is.source.spct()} \rightarrow \texttt{is.source\_spct()} \\ &\texttt{as.source.spct()} \rightarrow \texttt{as.source\_spct()} \\ &\textit{etc.} \end{split}
```

With exception of 'is. ()' and 'as. ()' functions shown above, all function names no longer contain dots. In the new names dots have been replaced by underscores. This also affected those is. () functions which are not S3 methods and which are used to query other attributes or properties of spectral objects.

```
\begin{split} & \texttt{is.effective()} \rightarrow \texttt{is\_effective()} \\ & \texttt{is.normalized()} \rightarrow \texttt{is\_normalized()} \\ & \texttt{is.rescaled()} \rightarrow \texttt{is\_scaled()} \\ & \textit{etc.} \end{split}
```

BSWF-waveband creation functions were also renamed to achieve consistency:

```
GEN.G() \rightarrow GEN_G()
DNA.N() \rightarrow DNA_N()
etc.
```

In the case of functions used to automate the creation of BSWF-based wavebands the old names are no longer indexed in the documentation, but the functions themselves will remain available. However, **in new scripts the new names should be used**, as they are *deprecated* and may be no longer available in future versions of the suite.

There is one further name change, required by conflicting names with other existing R functions: $Rescale() \rightarrow fscale()$, the f coming from function, as rescaling is done based on a summary function supplied through formal parameter f.

2 Changes in versions 0.7.0 and higher

generic_spct objects now have an atribute "spct.version" which help automate reading of *old* objects in future updates. It will also allow the detection of lost attributes.

The new generic_mspct objects have an attribute "mspct.version" which allows the detection of lost attributes.

3 Changes in versions 0.8.0 and higher

Although data.table is no longer used internally, objects from versions 6.0 or later do not need to be upgraded. They are fully compatible with the new version. Of course, data table syntax is not longer accepted, although this should make no difference as its use converted the spectral objects into data table objects.

However, use of copy() is no longer needed, as the default behaviour is no longer to use references. The assignment <- operator, can be used instead. A dummy copy() function can be defined if needed as:

```
copy <- function(x) {x}
```

4 Checking if an object is an old spectrum

Function is.old_spct can be used to query if an R object is a spectral object created with a version of package photobiology with version; 0.6.0. Function getObjectVersion can be used to query the version. Current version is 1, for older (or corrupted) objects the returned value is 0.

We need to only consider generic.spct as objects of derived classes like source.spct are by inheritance also generic.spct objects.

5 Upgrading old spectra

To upgrade the spectral objects what needs to be done is to change the names stored in the class attribute of spectral objects, as only the name of the class has changed rather than its definition.

Function upgrade_spct updates the class of old objects, adds the spct.version attribute and calls check() on the resulting object. The *upgrade* is done by reference (in place) in the object supplied as argument.

To demonstrate this we fake an spectral object defined by and earlier version of the package by changing the class attribute.

```
library(photobiology)

my.old.spct <- source_spct(w.length = 400:450, s.e.irrad = 1)
class(my.old.spct) <- gsub("_spct", ".spct", class(my.old.spct), fixed = TRUE)
class(my.old.spct)

## [1] "source.spct" "generic.spct" "tbl_df" "tbl"

## [5] "data.frame"
another.old.spct <- my.old.spct</pre>
```

We can use method upgrade_spct() to upgrade a single spectrum.

```
is.old_spct(my.old.spct)
## [1] TRUE

upgrade_spct(my.old.spct)
is.old_spct(my.old.spct)
## [1] TRUE

is.source_spct(my.old.spct)
## [1] FALSE
```

Another function upgrade_spectra takes as argument a list of R objects. Objects which are not *old spectra* are not altered, while *old spectra* are upgraded. The default is to upgrade all objects in the current environment and enclosing environments.

```
is.old_spct(another.old.spct)
## [1] TRUE

upgrade_spectra()

## Upgrading: another.old.spct
## Skipping: copy
## Upgrading: my.old.spct
## Skipping: my_version

is.old_spct(another.old.spct)

## [1] TRUE

is.source_spct(another.old.spct)

## [1] FALSE
```

We hope that these functions will make the transition to the new versions less painful.

6 Updating old scripts

Updating old scripts consists in searching and replacing the changed names. This can be done with a text editor or IDE like RStudio or 'blindly' with a shell script or batch file.

A command line tool like grep or the GUI program grepWin make it possible to do the substitutions on several script files in one job. It is wise when doing mass batch updating, to keep a backup of the old scripts at least until the updated scripts have been carefully tested.