Package 'FieldSpec'

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kage
actions for processing field spectroscopy data
0.89
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on A field spectroscopy package containing functions to process ctral files collected with an Analytical Spectral Devices (ASD) or extral Evolution instrument. The goal is to streamline and standardize ctral data processing for use in remote sensing, ecology, and plant ecophysiological research.
XML, Hmisc
testthat, Rprospect, signal
equirements
Windows, unix, mac
FreeBSD + file LICENSE
nt Authors
d yes
a FALSE
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average.spec

average replicate spectra within a directory of spectra files

Description

Function to average replicate spectra within a directory of spectra files

Usage

```
average.spec(file.dir = NULL, out.dir = NULL,
    start.wave = NULL, end.wave = NULL, step.size = NULL,
    bias.threshold = NULL, suffix.length = NULL,
    output.file.ext = NULL, spec.dataframe = FALSE,
    settings.file = NULL)
```

Arguments

file.dir	directory of spectra files to process
out.dir	output directory for processed spectra files
start.wave	starting wavelength of spectra files. Not needed if specified in XML settings file.
end.wave	ending wavelength of spectra files. Not needed if specified in XML settings file.
step.size	resolution of spectra files. E.g. 1 for 1nm, 5 for 5nm. Not needed if specified in XML settings file.
bias.threshold	reflectance/transmittance cutoff to remove spectra with anartificial bias (shift) due to improper spectral collection
suffix.length	length of auto numbering attached to ASD file names. This number of characters will be removed from the filename when averaged.
output.file.ex	t
	optional setting to set file extension of output files. Defaults to .csv
spec.dataframe	option to return a data frame with the converted spectra files
settings.file	settings file used for spectral processing options (OPTIONAL). Contains information related to the spectra collection instrument, output directories, and processing options such as applying a jump correction to the spectra files. Options in the settings file take precedent over options selected in the function call.

Author(s)

concat.spectra 3

Examples

Description

Concatenate a directory of spectra files into a single .csv file. Works on a single directory or a series of directories

Usage

```
concat.spectra(file.dir = NULL, out.dir = NULL,
  out.filename = NULL, in.file.ext = ".csv",
  out.file.ext = ".csv", transpose = FALSE,
  spec.dataframe = FALSE)
```

Arguments

file.dir	directory of spectra files to process. Currently works on spectra files formatted where each row is a wavelength and associated spectral observation.
out.dir	output directory for concatenated spectra files. If not set then the output directory defaults to file.dir (input directory)
out.filename	filename for concatenated spectra files
in.file.ext	file extension for individual spectra files. Defaults to ".csv"
out.file.ext	option to set the output extension. Defaults to ".csv"
transpose	option to transpose the spectra to/from row/column major output. This will be applied to both the returned dataframe (optional) and output file. TRUE/FALSE. Default FALSE
spec.dataframe	option to return a R dataframe containing the concatenated spectral data. TRUE/FALSE

Author(s)

Shawn P. Serbin

Examples

```
## Not run:
concat.spectra()
spectra <- concat.spectra(file.dir=file.dir,out.dir=out.dir,out.filename="concat.spectra",
out.file.ext=".csv",transpose=FALSE,spec.dataframe=FALSE)
## End(Not run)</pre>
```

4 extract.metadata

extract.metadata	Extract metadata for each spectra sample. or a directory.	Works on a single spectra
	or a arrectory.	

Description

Extract metadata for each spectra sample. Works on a single spectra or a directory.

Usage

```
extract.metadata(file.dir = NULL, out.dir = NULL,
  instrument = NULL, spec.file.ext = NULL,
  output.file.ext = ".csv", tz = NULL, intern = FALSE,
  settings.file = NULL)
```

Arguments

file.dir	File directory or filename of single spectra for processing
out.dir	Output directory for metadata information file
instrument	What instrument was used to collect spectra. Current options: ASD, SE
spec.file.ext	[Optional] Input spectra file extension. E.gasd (ASD) or .sed (Spectral Evolution). Default for ASD instruments is .asd. Default for Spectral Evolution instruments is .sed
output.file.ext	
	[Optional] Output file extension of metadata information file. Default .csv
tz	[Optional] Set the timezone of the spectra file collection. Used to covert spectra collection time to UTC. If unused it is assumed that the correct timezone is the current system timezone.
intern	logical. [Optional] Keep meta-data output as an internal object (TRUE) or write to file (FALSE) $$
settings.file	[Optional] Spectral settings file

Author(s)

Shawn P. Serbin

Examples

```
# ASD
file <- system.file("extdata/PM01_TIAM_B_LC_REFL00005.asd",package="FieldSpec")
extract.metadata(file,instrument="ASD",intern=TRUE)

# Spectral Evolution
file <- system.file("extdata/cvars_grape_leaf1_lc_rg_01236.sed",package="FieldSpec")
extract.metadata(file,instrument="SE",intern=TRUE)</pre>
```

extract.metadata.asd 5

 ${\it extract.metadata.asd} \quad {\it Extract\ metadata\ from\ raw\ binary\ ASD\ files.} \quad {\it Called\ from\ extract.metadata}$

Description

Extract metadata from raw binary ASD files. Called from extract.metadata

Usage

```
extract.metadata.asd(file.dir, out.dir, spec.file.ext,
  output.file.ext, tz, intern)
```

Arguments

file.dir File directory or filename of single spectra for processing

out.dir Output directory for meta-data information file

spec.file.ext [Optional] Input spectra file extension. E.g. .asd (ASD) or .sed (Spectral Evo-

lution). Default for ASD instruments is .asd. Default for Spectral Evolution

instruments is .sed

output.file.ext

[Optional] Output file extension of meta-data information file. Default .csv

tz [Optional] Set the timezone of the spectra file collection. Used to covert spectra

collection time to UTC. If unused it is assumed that the correct timezone is the

current system timezone.

intern logical. [Optional] Keep meta-data output as an internal object (TRUE) or write

to file (FALSE)

Author(s)

Shawn P. Serbin

extract.metadata.se $\it Extract$ metadata from Spectral Evolution files. Called from extract.metadata

Description

Extract metadata from Spectral Evolution files. Called from extract.metadata

Usage

```
extract.metadata.se(file.dir, out.dir, spec.file.ext,
  output.file.ext, tz, intern)
```

Author(s)

6 import.asd

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Import raw binary ASD spectra files

Description

A wrapper function to import raw binary ASD spectra files, apply jump correction to the imported files and average replicate files into a single spectrum. Calls read.asd(), jump.correction(), and average.spec() functions.

Usage

```
import.asd(asd.dir = NULL, output.dir = NULL,
 jump.correction = FALSE, average = FALSE,
 start.wave = NULL, end.wave = NULL, step.size = NULL,
 jumploc1 = NULL, jumploc2 = NULL, asd.file.ext = NULL,
 output.file.ext = NULL, settings.file = NULL)
```

Arguments

asd.dir	directory containing raw ASD files or sub-directories to be processed. Not needed if specified in settings file.
output.dir	main ouput directory. Not needed if specified in settings file.
jump.correction	1
	option to apply a jump correction to the imported spectra files. Not needed if specified in settings file.
average	option to average imported spectra files. Will use jump corrected files if exist. Not needed if specified in settings file.
start.wave	starting wavelength of spectra files. Not needed if specified in settings file.
end.wave	ending wavelength of spectra files. Not needed if specified in settings file.
step.size	resolution of spectra files. E.g. 1 for 1nm, 5 for 5nm. Not needed if specified in settings file.
jumploc1	location of the first jump in the spectra to correct. Not needed if jump.correction=FALSE or if specified in settings file.
jumploc2	location of the second jump in the spectra to correct. Not needed if jump.correction=FALSE or if specified in settings file.

asd.file.ext file extension for input binary asd spectra files

output.file.ext

optional setting to set file extension to output files. Defaults to .csv

settings file used for spectral processing options (OPTIONAL). Contains inforsettings.file mation related to the spectra collection instrument, output directories, and processing options such as applying a jump correction to the spectra files. Options

in the settings file take precedent over options selected in the function call.

Author(s)

jump.correction 7

jump.correction	apply a jump (splice) correction to imported ASD spectra files. This splice or jump occurs at the boundaries between detectors

Description

A function to apply a jump (splice) correction to imported spectra files

Usage

```
jump.correction(file.dir = NULL, out.dir = NULL,
    start.wave = NULL, end.wave = NULL, step.size = NULL,
    jumploc1 = NULL, jumploc2 = NULL, firstJumpMax = NULL,
    secondJumpMax = NULL, output.file.ext = NULL,
    settings.file = NULL)
```

Arguments

file.dir	directory of spectra files to process			
out.dir	output directory for processed spectra files			
start.wave	starting wavelength of spectra files. Not needed if specified in XML settings file.			
end.wave	ending wavelength of spectra files. Not needed if specified in XML settings file.			
step.size	resolution of spectra files. E.g. 1 for 1nm, 5 for 5nm. Not needed if specified in settings file.			
jumploc1	location of the first jump in the spectra to correct. Not needed if jump.correction=FALSE or if specified in XML settings file.			
jumploc2	location of the second jump in the spectra to correct. Not needed if jump.correction=FALSE or if specified in XML settings file.			
firstJumpMax	maximum jump threshold for the first jump location. Determines whether spectra will be corrected or flaged as bad. (Optional. Default is 0.02)			
secondJumpMax	maximum jump threshold for the second jump location. Determines whether spectra will be corrected or flaged as bad. (Optional. Default is 0.02)			
output.file.ext				
	option to set file extension of the output files. Defaults to .csv			
settings.file	settings file used for spectral processing options (OPTIONAL). Contains information related to the spectra collection instrument, output directories, and processing options such as applying a jump correction to the spectra files.			

Value

output list containing processed spectra and associated diagnostic information

Author(s)

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Examples

```
## Not run:
jump.correction()
jump.correction(file.dir,out.dir, start.wave=350,end.wave=2500,step.size=1,jumploc1=651,jumploc2=1451,
output.file.ext=".csv",settings.file=NULL)
## End(Not run)
```

read.asd

A function to import raw ASD binary files

Description

A function to import raw ASD binary files to ascii flat file format

Usage

```
read.asd(file.dir = NULL, out.dir = NULL,
  start.wave = NULL, end.wave = NULL, step.size = NULL,
  image = FALSE, asd.file.ext = ".asd",
  output.file.ext = ".csv", settings.file = NULL)
```

Arguments

file.dir	A single ASD binary file or directory of ASD files to import. Currently only supports single directory processing (i.e. no nested dir structures)				
out.dir	main output directory for processed spectra files.				
start.wave	starting wavelength of ASD binary spectra files. Depends on instrument.				
end.wave	ending wavelength of ASD binary spectra files. Depends on instrument.				
step.size	wavelength step size for ASD files. E.g. 1nm, 5nm, 10nm				
image	logical. Whether to produce png images of each spectrum				
asd.file.ext	file extension of ASD files. Usually ".asd" (Default)				
output.file.ext					
	optional setting to set file extension to output files. Defaults to .csv				
settings.file	settings file used for spectral processing options (OPTIONAL). Contains information related to the spectra collection instrument, output directories, and processing options such as applying a jump correction to the spectra files. Options in the settings file take precedent over options selected in the function call.				

Value

output for a single file returns a list with wavelengths and measured reflectance or transmittance values. For a directory, outputs individual ascii text files with wavelength and spectra values for each input ASD file.

Author(s)

settings 9

settings

parse settings file used for spectra file import and processing

Description

Read settings file for spectra import and processing

Usage

```
settings(input.file = NULL)
```

Arguments

input.file

settings file containing information needed for spectra processing

Author(s)

Shawn P. Serbin

Examples

```
## Not run:
settings <- settings()
settings <- settings('/home/$USER/settings.xml')
## End(Not run)</pre>
```

smooth.spectra

Smooth spectra file with Savitsky-Golay smoothing filter. Works with either a single spectrum or array of spectral observations in row-major format.

Description

Smooth spectra file with Savitsky-Golay smoothing filter. Works with either a single spectrum or array of spectral observations in row-major format.

Usage

```
smooth.spectra(file.dir = NULL, input.file = NULL,
  out.dir = NULL, out.filename = NULL, header = TRUE,
  p = NULL, n = NULL, length = "full", file.ext = ".csv")
```

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Arguments

file.dir	Directory of spectra files to apply sgolay smoothing filter.
input.file	Spectra file to apply sgolay smoothing filter. If not set then all files in file.dir are processed.
out.dir	Output directory for smoothed spectra files. If not set then processed spectra will be output to file.dir
out.filename	Output filename for processed spectra file. If not set then the original filename(s) will be modified with the .sg suffix.
header	logical. Does the spectra file(s) have a header line? Default = TRUE
р	SG filter order. Default 1.
n	SG filter length. Needs to be an odd value. Default 21
length	Apply sgolay smoothing filter to the entire (default=full) or subset of the spectrum. Define subset limits as (wavelength1,wavelength2), e.g. (2000,2500) in nanometers [NOT YET IMPLEMENTED]
file.ext	File extension for spectra files. Default '.csv'

Author(s)

Shawn P. Serbin

Examples

transpose.spectra

Transpose spectra file(s) from/to row or column major format. Works with either a single spectrum or directory of spectral observations.

Description

Transpose spectra file(s) from/to row or column major format. Works with either a single spectrum or directory of spectral observations.

Usage

```
transpose.spectra(file.dir = NULL, input.file = NULL,
  out.dir = NULL, out.filename = NULL,
  in.file.ext = ".csv", out.file.ext = ".csv",
  header = FALSE)
```

transpose.spectra 11

Arguments

file.dir directory of spectra files to process input.file name of input spectra file. If ommitted then the function transposes all files in the input file.dir out.dir output directory for transposed spectra file(s) output filename of transposed spectra file(s). If using a directory then the origiout.filename nal filenames are appended with the .t suffix file extension for individual spectra files. Defaults to ".csv" in.file.ext out.file.ext option to set the output extension. Defaults to ".csv" header logical. Do/does the file(s) have a header line?

Author(s)

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