

Phenopix

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This vignette aims at illustrating the main features of the package `phenopix`. This package was designed for processing digital images of the vegetation cover in order to compute vegetation indexes that can be in turn used to track the seasonal development of the vegetation. The analysis can be run on one or more portions of the image (so called regions of interest, ROIs). Regions of interest can be of any polygonal shape. For data processing, two approaches are available: ROI-averaged analysis or pixel based analysis. ROI-averaged analysis is based on the computation of vegetation indexes as the average of the entire ROI, whereas pixel based analysis allows to treat separately each pixel of the image. Data used to show phenopix package are from imagery archive of Torgnon Grassland and Larch sites, belonging to the phenocam network. The rationale and the objectives that motivate the processing chain that will be described here are established in the scientific literature since a quite long time. See References for a sample of the most relevant publications.

1 Install the package

The package `phenopix` is hosted in the r-forge repository and can be installed via the following command:

```
> install.packages("phenopix", repos="http://R-Forge.R-project.org")
```

Note that by running this command you will likely be asked to install the dependencies, which are available via the usual command:

```
> install.packages('package.name')
```

Once the package is properly installed you will be able to open this vignette by running:

```
> vignette('phenopix')
```

2 The steps

The first step is to give a well defined structure to a folder with the function `structureFolder()`.

The second step of the analysis is to choose a region of interest in an image. The functions useful for this step include:

- `DrawROI()` to draw a region of interest in your pictures
- `PrintROI()` to plot your ROI into an image
- `updateROI()` to apply ROI coordinates to an image of different size

Once the ROI is chosen, drawn and the underlying coordinates properly saved, color digital numbers are extracted and vegetation indexes (VIs) are calculated, using one main function `extractVIs()`.

Afterwards, raw VIs must be filtered out to get a reliable seasonal trajectory. This is the job of the function `autoFilter()`.

Then, several options are available to process the resulting data, ranging from fitting a curve to extracting break points on a seasonal trajectory, including several methods to extract relevant moments in the season (aka thresholds).

Functions useful for this step include:

- `greenProcess()` to fit a curve to the data (ROI-averaged approach)
- `greenExplore()` to fit all curves and thresholds with no uncertainty estimation, this function is coupled with
 - `plotExplore()`, which plots all fittings and thresholds in the object in output from `greenExplore()`
 - `spatialGreen()` to fit a curve to the data (pixel-based approach)
 - `PhenoBP()` to extract break points on a seasonal trajectory of data

A number of facilities are then built to plot, summarize, post process and render the results. These include:

- generic `plot()`, `print()`, `update()` and `summary()` functions with dedicated methods
- `plotSpatial()` to plot results from the pixel-based analysis
- `extractParameters()` to extract thresholds and curve parameters after the pixel-based analysis.

In the following paragraphs each step will be discussed and illustrated in detail.

3 Structuring a folder tree useful for the analysis

Giving a good structure to your analysis can make all subsequent steps simple and straightforward. If you are running a site that records images you will be dealing with quite heavy folders (with likely multiple years of data, hence some thousand files of images) that you need to handle with care. We suggest separate folders for each site (of course) but also year of analysis. Each year folder should contain a subfolder with all images to be processed (/IMG), one folder containing the reference image, i.e. the image you will use to draw your ROI (/REF), one folder containing data for the region of interest (/ROI) and one folder containing extracted vegetation indexes (/VI). The function `structureFolder()` provides a facility to create appropriate subfolders:

```
> library(phenopix)
> my.path <- structureFolder(path = getwd(), showWarnings = FALSE)

Put all your images in /home/gian/sweave/IMG/
Put your reference image in /home/gian/sweave/REF/
Draw your ROI with DrawROI():
  set path_img_ref to /home/gian/sweave/REF/
  set path_ROIs to /home/gian/sweave/ROI/
Then you can extractVIs():
  set img.path as /home/gian/sweave/IMG/
  set roi.path as /home/gian/sweave/ROI/
  set vi.path to /home/gian/sweave/VI/
-----
Alternatively, assign this function to an object and use named elements of the returned li

> str(my.path)

List of 4
$ img: chr "/home/gian/sweave/IMG/"
$ ref: chr "/home/gian/sweave/REF/"
$ roi: chr "/home/gian/sweave/ROI/"
$ VI : chr "/home/gian/sweave/VI/"

structureFolder() creates subfolder at a given path (in this example, the working directory) and stores all path in a named list. You can easily access all needed paths by simply pointing to the right object in your path object.
```

Note that if one folder already exists the function do not overwrite existing folders, but gives a warning. Note that the suggested structure is absolutely not mandatory. It is just a suggestion that can make easier the next steps. Once the folder structure is done, you have to:

- manually put your series of images to be processed into the `/IMG` folder
- manually put one of such images in the `/REF` folder, this is the image that will be printed on screen to draw your ROI.

4 Drawing a region of interest (ROI)

Apart from structuring folders, drawing a ROI is the first, hence most important step of the analysis. The procedure is based on two steps: first, a reference image (chosen by the user) is plotted by calling function `readJPEG()` from package `jpeg` and `rasterImage()`. In Fig. 1 is the reference image from one of our sites, Torgnon (NW Italy, 2100 m of elevation) and the code used to plot the image.

```

> img <- jpeg::readJPEG('REF/20130630T1000.jpg')
> ratio <- dim(img)[1]/dim(img)[2]
> par(mar=c(rep(1,4)))
> plot(0, type = "n", xlim = c(0, 1),
+ ylim = c(0, 1), axes = FALSE,
+ ylab='', xlab='')
> rasterImage(img, xleft = 0, ybottom = 0,
+ xright = 1,
+ ytop = ratio)

```



Figure 1: A jpeg image printed on a graphic device using `readJPEG()` and `rasterImage()`

This chunk of code is automatically included in the `DrawROI()` function.
The usage is:

```
> DrawROI(path_img_ref, path_ROIs, nroi = 2, roi.names=c("fg", "bg"))
```

where `path_img_ref` is the folder of your reference image, `path_ROIs` is the path in your computer where to store RData with ROI properties, number of ROIs and their names. A call to the function opens a graphic device and allows the use of `locator()` to define your ROI(s). Note that the use of `locator` is

somewhat system specific. Check out the help file `?locator` for more details. Locator allows to draw a polygon by left-clicking vertices and then right-clicking (or press ESC on MacOS) to close the polygon. If you have chosen more than one ROI, after closing your first polygon, the image will appear again unmodified to draw the second ROI, and so on. Note that the plot title helps you to remember which of your ROIs you are actually drawing. When you are done, in your `path_ROIs` an RData called `roi.data.RData` will be stored. This is actually a list with the following structure:

```
> load('ROI/roi.data.Rdata')
> str(roi.data)

List of 2
$ fg:List of 2
..$ pixels.in.roi:'data.frame':      273920 obs. of  3 variables:
...$ rowpos: num [1:273920] 0.00156 0.00313 0.00469 0.00625 0.00781 ...
...$ colpos: num [1:273920] 0.00156 0.00156 0.00156 0.00156 0.00156 ...
...$ pip   : int [1:273920] 0 0 0 0 0 0 0 0 0 ...
..$ vertices    :List of 2
...$ x: num [1:9] 0.0176 0.0193 0.2443 0.5051 0.6551 ...
...$ y: num [1:9] 0.2666 0.0288 0.0194 0.0138 0.0232 ...
$ bg:List of 2
..$ pixels.in.roi:'data.frame':      273920 obs. of  3 variables:
...$ rowpos: num [1:273920] 0.00156 0.00313 0.00469 0.00625 0.00781 ...
...$ colpos: num [1:273920] 0.00156 0.00156 0.00156 0.00156 0.00156 ...
...$ pip   : int [1:273920] 0 0 0 0 0 0 0 0 0 ...
..$ vertices    :List of 2
...$ x: num [1:8] 0.0278 0.0244 0.3346 0.5699 0.633 ...
...$ y: num [1:8] 0.416 0.364 0.332 0.327 0.388 ...
```

A two elements list (one for each ROI) with ROI names. Each element is again a list containing two elements. One is a data.frame containing coordinates of all image pixels, together with a code indicating whether the given pixel belongs to the ROI or not. The second is a list with the coordinates of ROI margins as in output from `locator()`.

Additionally, in `path_ROIs` separate jpeg files for each of your regions of interest are stored. A call to the function `printROI()` allows to plot in the same graph all existing ROIs for a picture. In the example from Torgnon, two ROIs were drawn, one corresponding to the foreground of the image and one to

the background (`fg` and `bg` respectively. Here is the code to generate the plot in fig. 2:

```
> PrintROI(path_img_ref = 'REF/20130630T1000.jpg',
+ path_ROIs = 'ROI/',
+ which = 'all',
+ col = palette())
```

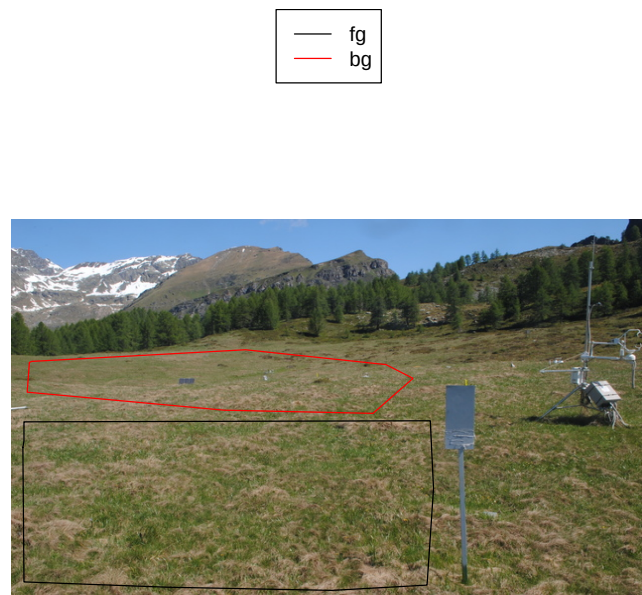


Figure 2: A plot of your regions of interest (ROIs), in output from `PrintROI()`

When you draw a ROI on your best quality image (say 640 x 428 pixels, as the REF image for Torgnon) you will probably need to identify the same ROI in smaller size images. This will be the case, for example, if you want to conduct a pixel-based analysis, illustrated later on. Pixel based analysis is computationally intense and therefore it is suggested to run it on rather small size images. The function `updateROI()` allows to recalculate pixels falling within a given ROI in images of different size compared to the one where the ROI was first drawn. Usage is:

```
> updateROI(old.roi.path, new.path.img.ref, new.roi.path)
```

`old.roi.path` is the path to the folder containing your old `roi.data.Rdata`,

`new.path.img.ref` is the path to the folder containing your re-sized image, and `new.roi.path` is the folder where your new `roi.data.Rdata` will be stored.

5 Extraction of vegetation indexes

At this point, you have an r object stored as `roi.data.Rdata` in your ROI path that defines which pixels fall into one or more ROIs. The next step will be to extract information on those pixels from each of your images. The function that performs this task is `extractVIs()` and the usage is as follows:

```
> extractVIs(img.path, roi.path, vi.path = NULL, roi.name = NULL,  
+   plot = TRUE, begin = NULL, spatial = FALSE)
```

`img.path` is the path where one year of images are stored. It is not mandatory to have only one year of images in your folder. However it is suggested to structure your data into separate folders for each year because nearly all the functions we will see later are designed to work on a single season of data. `roi.path` is the path to your `roi.data.Rdata`, `VI.path` is the path where extracted vegetation indexes will be saved. Hence, this function can be assigned to an object to have your vegetation indexes returned into R, or alternatively loaded later if not assigned. The argument `begin` allows to set a beginning date to update an existing time series without reprocessing the whole year of data. For example, if you run `extractVIs` in mid June to have a first look at your time series, once your season will be completed you do not want to re-run the analysis on the already processed images. Hence, you set the argument `begin` to the first unprocessed date. A new `roi.data.Rdata` will be saved in your path, with the beginning date incorporated in the filename. It is up to the user to bind the old and new time series.

The argument `spatial` allows to perform pixel-based analysis. This is a topic that will be discussed later.

The construction of the time series implies that R recognizes a time vector, typically retrieved from the file name of each picture. The function responsible for this conversion is `ExtractDateFilename()`. It is a rather internal function but it is worth to look how it works to properly set the filenames of your imagery archive. Arguments to the function are `filename` and `date.code`. `filename` must be a character string with an underscore '`_`' that separates site name and date (e.g. '`torgnon_20140728.jpg`'). One and only one underscore must appear before the date stamp. The format of your date must be provided in `date.code`. In the example above, `date.code` will be: '`yyyymmdd`'. Let's look at some

examples, but before doing so, it is worth to remember the the file naming system is under your responsibility when you set up the storage process for your images, or by some renaming routines set up later.

```
> filename <- 'torgnon_20140728.jpg' ## correct, with no hour
> ## if hour is missing it is defaulted to 12 pm
> extractDateFilename(filename, date.code='yyyymmdd')

[1] "2014-07-28 12:00:00 CEST"

> filename <- 'torgnon_201407281100.jpg' ## correct, with hour
> ## hours and minutes to upper letters, in R POSIX style
> extractDateFilename(filename, date.code='yyyymddHHMM')

[1] "2014-07-28 11:00:00 CEST"

> filename <- 'torgnon_ND_201407281100.jpg' ## wrong, with two
> ## underscores before date, the function returns NA
> extractDateFilename(filename, date.code='yyyymddHHMM')

[1] NA

> filename <- 'torgnon_1407281100.jpg' ## correct, with 2 numbers for the year
> extractDateFilename(filename, date.code='yyymddHHMM')

[1] "2014-07-28 11:00:00 CEST"

> ## any separator for date elements is allowed
> ## including underscore
> filename <- 'torgnon_2014.07_28-11.00.jpg'
> extractDateFilename(filename, date.code='yyyy.mm_dd-HH.MM')

[1] "2014-07-28 11:00:00 CEST"
```

Now let's look from closer at the structure of the object `VI.data` saved in your `/VI` directory.

```
> load('VI/VI.data.Rdata')
> summary(VI.data) ## a list with two data.frames, one for each ROI

      Length Class      Mode
fg    18   data.frame  list
bg    18   data.frame  list
```

```

> names(VI.data[[1]]) ## check which vegetation indexes are extracted

[1] "date"    "doy"     "r.av"    "g.av"    "b.av"    "r.sd"    "g.sd"    "b.sd"
[9] "bri.av"  "bri.sd"  "gi.av"   "gi.sd"   "gei.av"  "gei.sd"  "ri.av"   "ri.sd"
[17] "bi.av"   "bi.sd"

```

The processing of each ROI produces a data.frame object with date in POSIX format, numeric day of year (doy), and the vegetation indexes. Green, red and blue digital numbers (range [0,255]) averaged over the ROI (g.av, r.av and b.av, respectively), their standard deviations (g.sd, r.sd and b.sd). bri.av is the ROI averaged brightness, calculated as the sum of red green and blue digital numbers for each pixel and then averaged. From the digital numbers (dn) of each color, relative indexes (rel.i) are calculated as follows:

$$\text{rel.i} = \text{dn color} / (\text{dn red} + \text{dn green} + \text{dn blue})$$

These values are calculated for each pixel and then averaged over the entire ROI (columns gi.av, ri.av, bi.av), and the standard deviation is calculated as well. In fig.3 you can see how a seasonal course of raw color digital numbers of a subalpine grassland site looks like:

```

> with(VI.data$fg, plot(date, r.av, pch=20, col='red',
+   ylim=c(0,255), ylab='DN [0,255]'))
> with(VI.data$fg, points(date, g.av, col='green', pch=20))
> with(VI.data$fg, points(date, b.av, col='blue', pch=20))

```

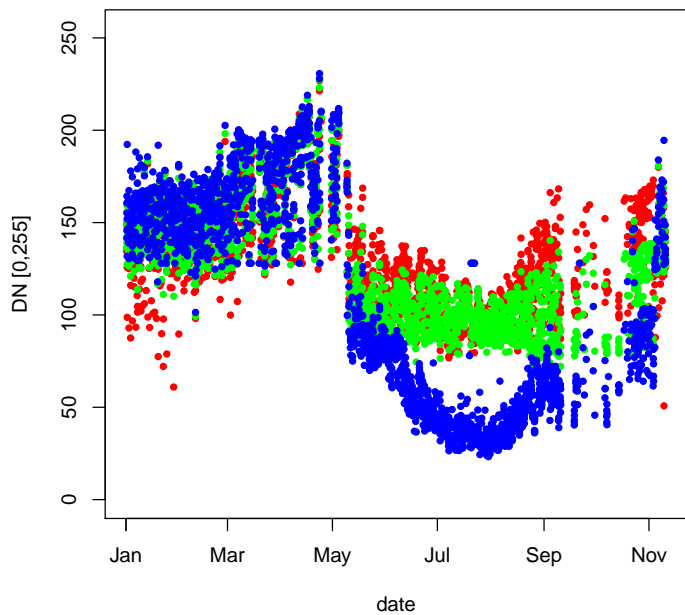


Figure 3: Seasonal course of raw digital numbers, Torgnon, year 2012

More interesting is the plot of relative indexes (fig. 4):

```

> with(VI.data$fg, plot(date, ri.av, pch=20, col='red',
+   ylim=c(0.1,0.6), ylab='Relative indexes'))
> with(VI.data$fg, points(date, gi.av, col='green', pch=20))
> with(VI.data$fg, points(date, bi.av, col='blue', pch=20))

```

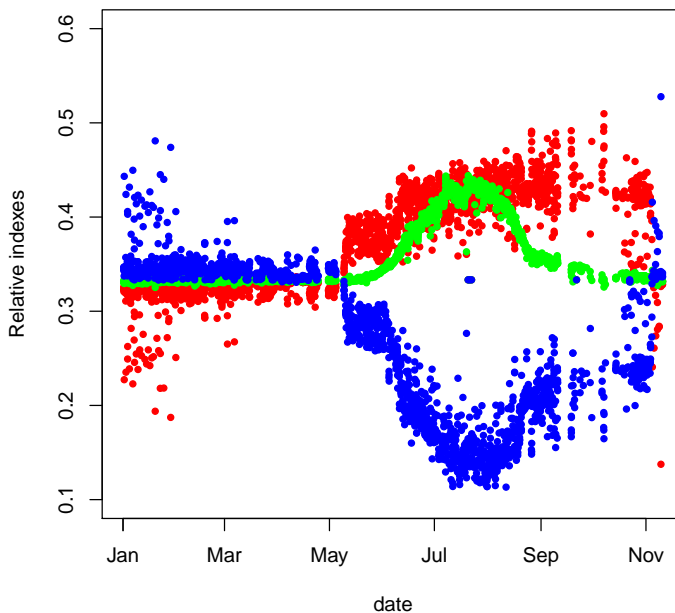


Figure 4: Seasonal course of relative green red and blue indexes, Torgnon grass-land, year 2012

- Several patterns are interesting in the seasonal course of fig.4:
- Snow disappearance (mid May) lead to an increase in relative red and a sharp decrease in relative blue
 - The green signal follows a bell shaped pattern throughout the growing season, with a maximum in late July. This signal is somewhat mirrored by an inverse behavior of relative blue, whereas relative red gradually increases throughout the season.

6 Filter out data

Data retrieved from images often need robust methods for polishing the time series. Bad weather conditions, low illumination, dirty lenses are among the

most common issues that determine noise in the time series of vegetation indexes. Accordingly we designed a function `autoFilter()` based on 4 different approaches, see the examples in `?autoFilter` for details in the filtering procedure. The function is designed to receive in input a data.frame structured as in output from `extractVIs`, hence its default expression may appear rather complicate:

```
> autoFilter(data, dn=c('ri.av', 'gi.av', 'bi.av'), raw.dn = FALSE,  
+ brt = 'bri.av', na.fill = TRUE,  
+ filter = c("night", "spline", "max"),  
+ filter.options = NULL, plot = TRUE, ...)
```

But when applied to the `VI.data` object generated before it is quite straightforward as you see in the code below. Note also that `autoFilter()` returns by default a diagnostic plot shown in fig.5:

```

> filtered.data <- autoFilter(VI.data$fg)
> str(filtered.data)

âzoo series from 1 to 314
Data: num [1:277, 1:7] 0.323 0.324 0.318 0.324 0.316 ...
- attr(*, "dimnames")=List of 2
..$ : NULL
..$ : chr [1:7] "rcc" "gcc" "bcc" "brt" ...
Index: num [1:277] 1 2 3 4 5 6 7 8 9 10 ...

> names(filtered.data)

[1] "rcc"           "gcc"           "bcc"           "brt"
[5] "night.filtered" "spline.filtered" "max.filtered"

```

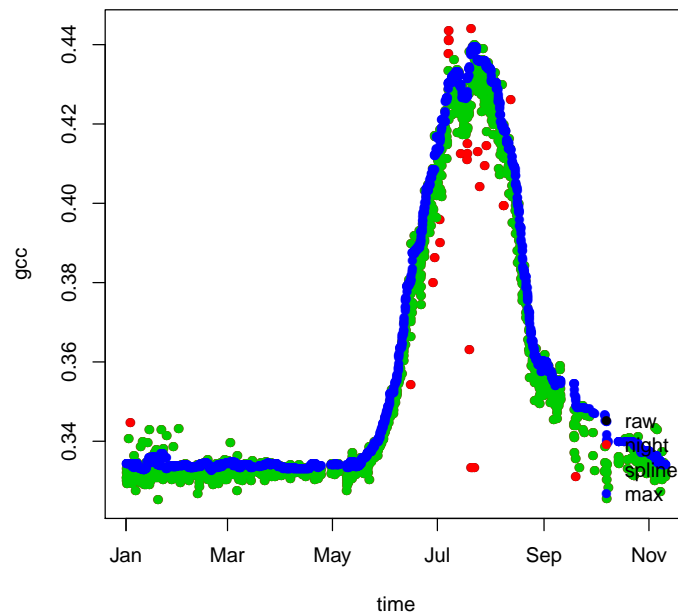


Figure 5: Raw and filtered relative greenness index, default plot of function `autoFilter()`

In the structure of the output `data.frame` there are two important points:

- We introduce here a new class of R objects (`zoo`). From here on all further analyses are based on `zoo` (or, to a lesser extent `ts`) time series. The time index of the data is numeric day of year (doy). As a consequence, the attribute `year`

is lost at this step of the analysis (i.e. we suggest to include it in the object name);

- The function `autoFilter` aggregates the data at a daily time step by default. The returned `data.frame` contains unfiltered (but still daily aggregated) color indexes (here called `gcc`, `rcc` and `bcc`, `cc` standing for chromatic coordinate) and a column of data for each filtering step. The name of the filter applied is reported in the column name.

For those unfamiliar with the `zoo` structure we have created a function `convert` to convert from `zoo` to a normal `data.frame`

```
> dataframed <- convert(filtered.data, year='2012')
> str(dataframed)

'data.frame':      277 obs. of  9 variables:
 $ rcc           : num  0.323 0.324 0.318 0.324 0.316 ...
 $ gcc           : num  0.331 0.332 0.334 0.331 0.332 ...
 $ bcc           : num  0.346 0.344 0.35 0.343 0.354 ...
 $ brt           : num  442 409 447 409 435 ...
 $ night.filtered: num  0.331 0.332 0.334 0.331 0.332 ...
 $ spline.filtered: num  0.331 0.332 0.333 0.331 0.332 ...
 $ max.filtered   : num  0.334 0.334 0.334 0.334 0.334 ...
 $ doy            : num  1 2 3 4 5 6 7 8 9 10 ...
 $ time           : POSIXct, format: "2012-01-01" "2012-01-02" ...
```

However, we strongly recommend to get familiar with the `zoo` package since it has wonderful facilities for plotting, aggregating and filling time series.

7 Fit a curve to the data

The seasonal trajectory of greenness index of a vegetation canopy provides per se important information, but to turn qualitative information into quantitative data we need to make some more computation. Traditionally, data similar to these (e.g. satellite-based NDVI trajectories) are processed in two main ways:

- extract time thresholds based on a percentage of development (e.g. the day when half of the maximum value of the index is reached);
- fit a curve and extract relevant thresholds based on curve properties.

In the package `phenopix` both possibilities are available. The core function for data fitting and threshold extraction is `greenProcess()`. This function calls and is related to several rather internal functions that perform the different fittings. Available fittings include:

- the fit of a cubic spline
- the fit of an equation proposed by Beck et al. (2006)
- the fit of an equation proposed by Elmore et al. (2012)
- the fit of an equation proposed by Klosterman et al. (2014) with two implementations
 - the fit of an equation proposed by Gu et al. (2009)

All fits are based on a double - logistic function with a different number of parameters.

After curve fitting, relevant dates in the seasonal trajectory (aka thresholds) are extracted with different methods:

- A method called **spline** (the name will soon change) which splits the seasonal course into increasing and decreasing trajectory based on the sign of the first derivative and then identifies the 50% of both the increasing and decreasing trajectory. It allows to determine start of season (sos), end of season (eos) and length of season (los) as the difference between the two.
- A method called **derivatives** which extends **spline** in that it also calculates maximum growing and decreasing rates
- A method based on Klosterman approach which individuates 4 moments in the seasonal trajectory. Greenup represents the beginning of growth, maturity represents the reaching of some summer plateau, senescence represents the beginning of green decrease (or yellowing increase) and dormancy represents the end of the growing season.
- A method based on Gu approach which individuates 4 moments and some other curve parameters. The 4 relevant moments do not differ in their meaning compared to Klosterman phases, and are called upturn date (UD), stabilization date (SD), downturn date (DD) and recession date (RD).

Detail on curve fitting and threshold extraction is provided in the help function of **?greenProcess** as well as in the help files of other more internal functions such as **?KlostermanFit**, **?GuFit**, **?PhenoExtract**. In fig.6 we show 4 different fitting methods applied to the same data (Torgnon grassland).

```
> ## spline curve + spline thresholds
> fit1 <- greenProcess(filtered.data$max.filtered,
+   'spline',
+   'spline',
+   plot=FALSE
+ )
> summary(fit1)
> ## check the plot
```

```

> plot(fit1, type='p', pch=20, col='grey')
> ## Beck fitting + derivatives
> fit2 <- greenProcess(filtered.data$max.filtered,
+ 'beck',
+ 'derivatives',
+ plot=FALSE)
> summary(fit2)
> plot(fit2, type='p', pch=20, col='grey')
> ## klosterman fitting + klosterman thresholds
> fit3 <- greenProcess(filtered.data$max.filtered,
+ 'klosterman',
+ 'klosterman',
+ plot=FALSE)
> summary(fit3)
> ## plot(fit3, type='p', pch=20, col='grey')
>
> ## gu fitting and thresholds
> fit4 <- greenProcess(filtered.data$max.filtered,
+ 'gu',
+ 'gu',
+ plot=FALSE)
> summary(fit4)
> plot(fit4, type='p', pch=20, col='grey')

```

```

> ## show all together
> par(lwd=3)
> plot(filtered.data$max.filtered, type='p', pch=20,
+       ylab='Green chromatic coordinate', xlab='DOYs')
> lines(fitted(fit1), col='blue')
> lines(fitted(fit2), col='red')
> lines(fitted(fit3), col='green')
> lines(fitted(fit4), col='violet')
> legend('topleft', col=c('blue', 'red', 'green', 'violet'),
+        lty=1, legend=c('Spline', 'Beck', 'Klosterman', 'Gu'),
+        bty='n')

```

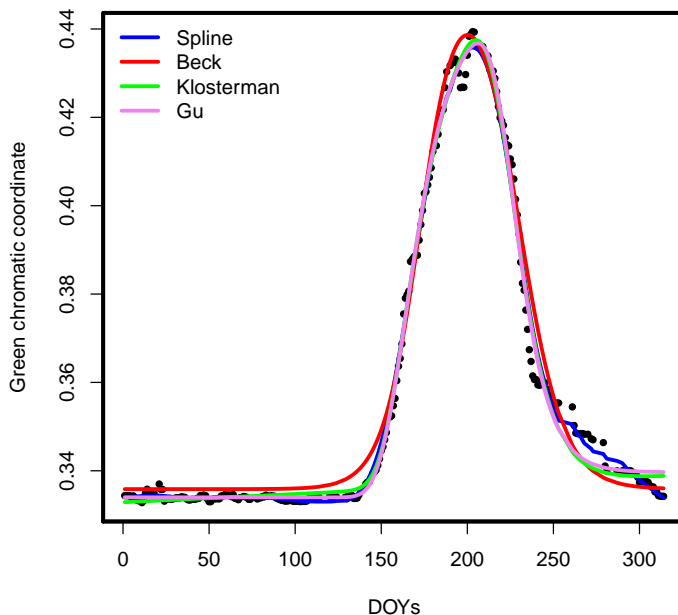


Figure 6: Comparison of 4 different fittings from `phenopix` package

The function `greenProcess` creates an object of class `phenopix` with dedicated methods. The `summary` function displays a summary of the input data and of the predicted points. It then reports the formula of the fitting equation, if pertinent, see e.g. summary of `fit1` which is not based on an equation. Thresholds are printed as well. Note also the `fitted` function applied to `phenopix` object that returns a `zoo` time series of fitted values that can be directly `lined` to the

plot.

To complete the overview on display generic methods applied to the objects of class `phenopix` here is the application of generic `plot` (fig.7) and `print` functions:

```
> plot(fit4, pch=20, col='grey', type='p',
+      xlab='DOYs', ylab='Green chromatic coordinates')
```

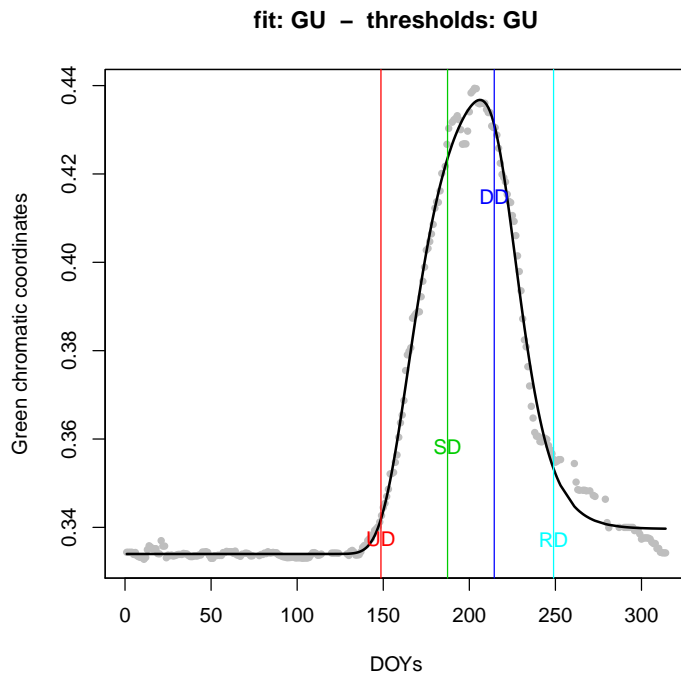


Figure 7: Generic plot function applied to `phenopix` objects

```
> print(fit4)

##### phenopix time series processing #####
FITTING: GU

PREDICTED VALUES:
  Index      x$fit$fit$predicted
  Min.   : 1.0   Min.   :0.3340
  1st Qu.: 71.0  1st Qu.:0.3340
  Median :152.0  Median :0.3397
  Mean   :150.6  Mean   :0.3582
```

```

3rd Qu.:223.0   3rd Qu.:0.3750
Max.     :314.0   Max.    :0.4368

```

FITTING EQUATION:

```

expression(y0 + (a1/(1 + exp(-(t - t01)/b1))^c1) - (a2/(1 + exp(-(t -
t02)/b2))^c2))

```

FITTING PARAMETERS:

y0	a1	a2	t01	t02	b1
0.33339788	0.1104471	0.1047788	129.3082257	200.3364257	14.6352282
b2	c1	c2			
11.7331508	11.1601187	9.0723878			

THRESHOLDS: GU

UD	SD	DD	RD	maxline
148.661704355	187.342880427	214.467153725	248.926022679	0.436761779
baseline	prr	psr	plateau.slope	
0.3333978802	0.002657183	-0.003004145	0.000327951	

UNCERTAINTY: FALSE

N of replications = 0

The `print` function returns information similar to `summary` but it also reports which fitting and threshold methods were used, and if the uncertainty was estimated. The `plot` function returns a plot similar to the one constructed above, except that extracted thresholds are also shown as vertical colored lines. Fig.5 shows that different fitting equation lead to very similar fitted values on the example from Torgnon data. For the sake of robustness, in such situation it is preferable to choose a fitted equation rather than a spline fit. Let's decide to choose the fitting from Gu. Now let's look from closer how do the different threshold extraction methods impact when applied to the same fitted curve in fig.8 (and note the use of `update` generic function with method `phenopix`):

```

> fit4.spline <- update(fit4, 'spline', plot=FALSE)
> fit4.klosterman <- update(fit4, 'klosterman', plot=FALSE)
> fit4.gu <- update(fit4, 'gu', plot=FALSE)
> par(mfrow=c(2,2), oma=rep(5,4,4,2), mar=rep(0,4))
> plot(fit4.spline, type='n', main='', xaxt='n')
> mtext('spline', 3, adj=0.1, line=-2)
> plot(fit4.klosterman, type='n', main='', xaxt='n', yaxt='n')
> mtext('klosterman', 3, adj=0.1, line=-2)
> plot(0, type='n', axes=FALSE, xlab='', ylab='')
> plot(fit4.gu, type='n', main='', yaxt='n')
> axis(4)
> mtext('gu', 3, adj=0.1, line=-2)

```

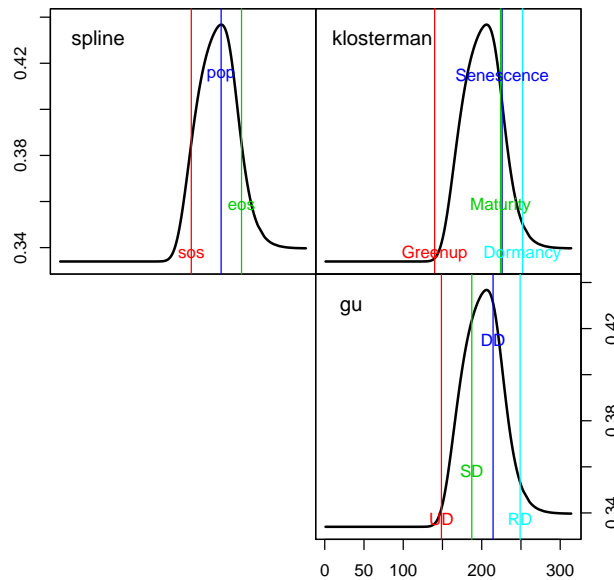


Figure 8: Three threshold methods applied to the Gu fitting

The **spline** thresholds (50% of increasing and decreasing trajectory) hold a different meaning compared to Klosterman and Gu thresholds. The latter two show good correspondence except that the Klosterman's beginning of senescence occurs later compared to correspondent phase in Gu thresholds (i.e DD, downturn date).

In this paragraph we have shown 4 different approaches to mathematically describe the seasonal trajectory of greenness, with additionally 5 methods to extract thresholds on the obtained curves. The combination of curves and threshold methods leads to as many as 20 possible approaches to describe a seasonal trajectory. Sometimes it could be useful to make a decision on which curves and thresholds to use, without computing the uncertainty on all of them. To do so we have designed two functions that provide a quick overview on what would be the best fit and threshold method for your actual trajectory. Here is how to compute the 20 combinations of fit and uncertainty in a single function:

```
> explored <- greenExplore(filtered.data$max.filtered)
```

explored is a list with $20 + 1$ elements, i.e. the 20 combinations + a vector containing the RMSEs from each of the 4 fittings. This object will only be used as argument of the `plotExplore()` function (fig.9):

```
> plotExplore(explored)
```

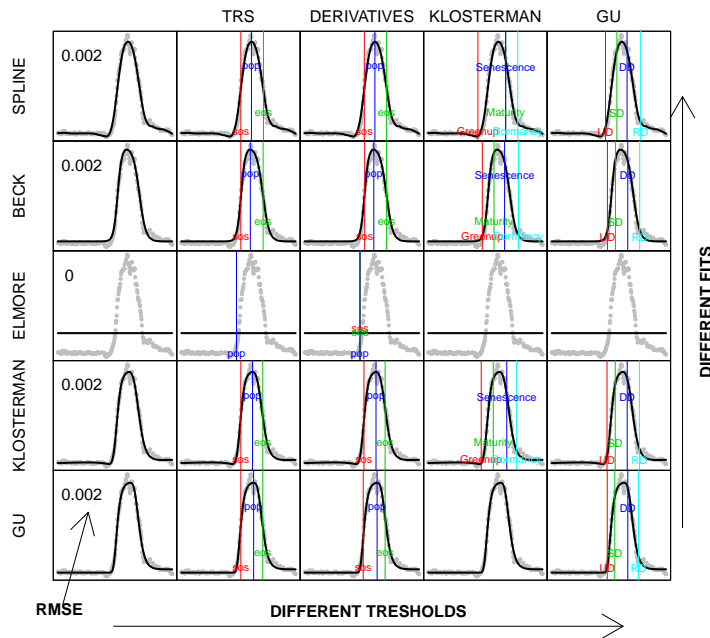


Figure 9: Overview of all combinations of curves and fits as obtained by the `plotExplore` function

The plot in fig.9 shows the impact of different fittings (moving up-downwards) and different thresholds (from left to right) on the same data (Torgnon grass-

land). The RMSE for each of the four fitting methods is also annotated in the first column. This plot might be useful to choose the most appropriate fitting and thresholding methods on your data.

8 The uncertainty estimation

One main functionality of the package is the uncertainty estimation. This is performed in different ways depending on the fitting equation. The basic idea behind the uncertainty estimation is how good the smoothing curve fits to the data. The residuals between fitted and observed is therefore used to generate random noise to the data and fitting is applied recursively to randomly - noised original data. This procedure results in an ensemble of curves, curve parameters and extracted thresholds that represent the uncertainty estimate. The uncertainty on curve parameters is automatically propagated to threshold extraction. In the following example the uncertainty estimation is performed on Torgnon grassland data fitted with the approach of Klosterman et al. (2014), with 100 replications. Here is the code:

```
> fit.complete <- greenProcess(ts = filtered.data$max.filtered,
+                               fit = 'gu',
+                               threshold= 'gu',
+                               plot = FALSE,
+                               uncert = TRUE,
+                               nrep = 50)
```

And here is `fit.complete` printed:

```
> print(fit.complete)

##### phenopix time series processing #####
FITTING: KLOSTERMAN

PREDICTED VALUES:
  Index      x$fit$fit$predicted
Min.   : 1.0   Min.   :0.3329
1st Qu.: 71.0  1st Qu.:0.3342
Median :152.0  Median :0.3388
Mean   :150.6  Mean   :0.3582
3rd Qu.:223.0  3rd Qu.:0.3758
```

```
Max. :314.0 Max. :0.4375
```

FITTING EQUATION:

```
expression((a1 * t + b1) + (a2 * t^2 + b2 * t + c) * (1/(1 +
q1 * exp(-B1 * (t - m1)))^v1 - 1/(1 + q2 * exp(-B2 * (t -
m2)))^v2))
```

FITTING PARAMETERS:

a1	a2	b1	b2	c
1.867151e-05	5.090923e-06	3.328690e-01	-1.245649e-03	1.519304e-01
B1	B2	m1	m2	q1
8.736767e-02	8.705025e-02	1.299899e+02	2.056079e+02	3.967244e+00
q2	v1	v2		
1.995802e+00	4.198100e+00	2.418243e+00		

THRESHOLDS: GU ENVELOPE:QUANTILES

	UD	SD	DD	RD	maxline	baseline	prr
10%	146.9326	188.5265	213.0008	251.0400	0.4374889	0.3328877	0.002446034
50%	147.2705	189.1760	213.1443	251.4086	0.4374889	0.3328877	0.002498147
90%	147.6557	189.7483	213.3274	251.6832	0.4374889	0.3328877	0.002559312
	psr	plateau.slope					
10%	-0.002779570	0.0003013841					
50%	-0.002737603	0.0003013841					
90%	-0.002707150	0.0003013841					

UNCERTAINTY: TRUE

```
N of replications = 100
```

As you can see from the output, the default behavior of `greenProcess()` for the computation of uncertainty is to provide the median, 10th and 90th percentile of the uncertainty ensemble. This may be changed by modifying the `envelope` argument. The other possible option is `min-max` to get minimum mean and maximum. In addition, the quantiles to be chosen with `envelope = quantiles` can be changed by modifying the `quantile` argument. Here is the example:

```
> print(update(fit.complete, 'gu', envelope='min-max', plot = FALSE))

#### phenopix time series processing ####
```

FITTING: KLOSTERMAN

PREDICTED VALUES:

```
Index      x$fit$fit$predicted
Min.    : 1.0   Min.   :0.3329
1st Qu.: 71.0  1st Qu.:0.3342
Median  :152.0  Median  :0.3388
Mean    :150.6  Mean   :0.3582
3rd Qu.:223.0  3rd Qu.:0.3758
Max.    :314.0  Max.   :0.4375
```

FITTING EQUATION:

```
expression((a1 * t + b1) + (a2 * t^2 + b2 * t + c) * (1/(1 +
q1 * exp(-B1 * (t - m1)))^v1 - 1/(1 + q2 * exp(-B2 * (t -
m2)))^v2))
```

FITTING PARAMETERS:

a1	a2	b1	b2	c
1.867151e-05	5.090923e-06	3.328690e-01	-1.245649e-03	1.519304e-01
B1	B2	m1	m2	q1
8.736767e-02	8.705025e-02	1.299899e+02	2.056079e+02	3.967244e+00
q2	v1	v2		
1.995802e+00	4.198100e+00	2.418243e+00		

THRESHOLDS: GU ENVELOPE:MIN-MAX

	UD	SD	DD	RD	maxline	baseline	prr
min	146.1843	186.1598	212.4913	250.4383	0.4374889	0.3328877	0.002368027
mean	147.2843	189.1266	213.1333	251.3941	0.4374889	0.3328877	0.002501166
max	149.1719	190.3566	213.5794	252.0516	0.4374889	0.3328877	0.002827987
	psr	plateau.slope					
min	-0.002850601	0.0002539504					
mean	-0.002739581	0.0003043358					
max	-0.002667329	0.0004128567					

UNCERTAINTY: TRUE

N of replications = 100

Beside the few options available by default and described above, the uncertainty data.frame is accessible via the `extract` command, by running:

```
> extract(fit.complete, 'metrics.uncert') ## get threshold uncertainty data`  
> extract(fit.complete, 'params.uncert') ## get parameters of each fitting curve`
```

For example, if you want to use thresholds extracted from the true model and construct uncertainty envelope on them, you can access the uncertainty data.frame by the commands given above. Note than when the uncertainty is computed, also the `plot` function changes its behavior, in that it also shows the uncertainty curve ensemble and an error bar on extracted thresholds (fig.8).

```
> plot(fit.complete, type='p', pch=20)
```

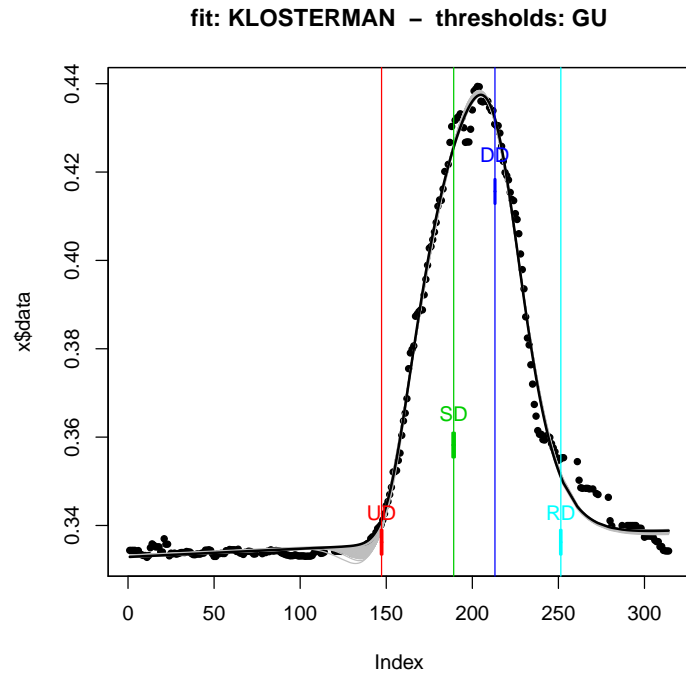


Figure 10: The Uncertainty Estimation (100 rep) on Klosterman fit and Gu thresholds

The distribution of uncertainty parameters (thresholds + curve parameters) can also be evaluated by means of box-plots with an extra option to the default `plot` method:

```
> plot(fit.complete, what='thresholds')
```

By using the `update` function you can also extract thresholds according to a different method, without refitting the data. Here is the code:

```
> update(fit.complete, 'klosterman', plot=FALSE)
##### phenopix time series processing #####

```

FITTING: KLOSTERMAN

PREDICTED VALUES:

Index	x\$fit\$fit\$predicted
Min.	: 1.0 Min. :0.3329
1st Qu.:	71.0 1st Qu.:0.3342
Median :	152.0 Median :0.3388
Mean :	150.6 Mean :0.3582
3rd Qu.:	223.0 3rd Qu.:0.3758
Max. :	314.0 Max. :0.4375

FITTING EQUATION:

```
expression((a1 * t + b1) + (a2 * t^2 + b2 * t + c) * (1/(1 +
q1 * exp(-B1 * (t - m1)))^v1 - 1/(1 + q2 * exp(-B2 * (t -
m2)))^v2))
```

FITTING PARAMETERS:

a1	a2	b1	b2	c
1.867151e-05	5.090923e-06	3.328690e-01	-1.245649e-03	1.519304e-01
B1	B2	m1	m2	q1
8.736767e-02	8.705025e-02	1.299899e+02	2.056079e+02	3.967244e+00
q2	v1	v2		
1.995802e+00	4.198100e+00	2.418243e+00		

THRESHOLDS: KLOSTERMAN ENVELOPE:QUANTILES

	Greenup	Maturity	Senescence	Dormancy
10%	136	223	225	253
50%	138	224	225	253
90%	140	224	225	253

UNCERTAINTY: TRUE

N of replications = 100

There is a last method to define thresholds on a time series that does not need a fitting. It implements the use of break points from the package **strucchange** and works as follows:

```

> print(PhenoBP(x = filtered.data$max.filtered,
+                 breaks = 3, plot = FALSE,
+                 confidence= 0.99))

  bp1 bp2 bp3
0.5% 141 203 244
mean 142 204 245
99.5% 143 205 246

```

The user can set the maximum number of breakpoints to be identified, the confidence interval at which the calculation must be performed and an option or a plot. The output dataframe contains the day of the year for each of the breakpoints and their respective confidence intervals.

9 Pushing forward the analysis: pixel - based phenology

In order to thoroughly exploit the capabilities of an imagery archive, spatial analysis represents the most promising feature. Hence, specific functions are built to fit curves and extract thresholds on each pixel included in a region of interest instead of averaging the greenness index over the entire ROI. The computation for this analysis may be quite intense, therefore it is suggested to either conduct it in small ROIs or on images with reduced size. For this approach a specific function `updateROI()` was designed. The default of the function was illustrated before. To illustrate the group of new functions for the spatial analysis we present a new dataset from Torgnon Larch Site. At this site we defined 4 ROIs corresponding to individual or small groups of trees which showed a somewhat different phenology by simply looking at the pictures, as shown in the plot below. ROIs distinguish plants where the autumn phenology occurs slightly earlier, or later and an evergreen spruce tree (fig.11).

```
> PrintROI('larch/2012/REF/20121004T1400.jpg', 'larch/2012/ROI/')
```

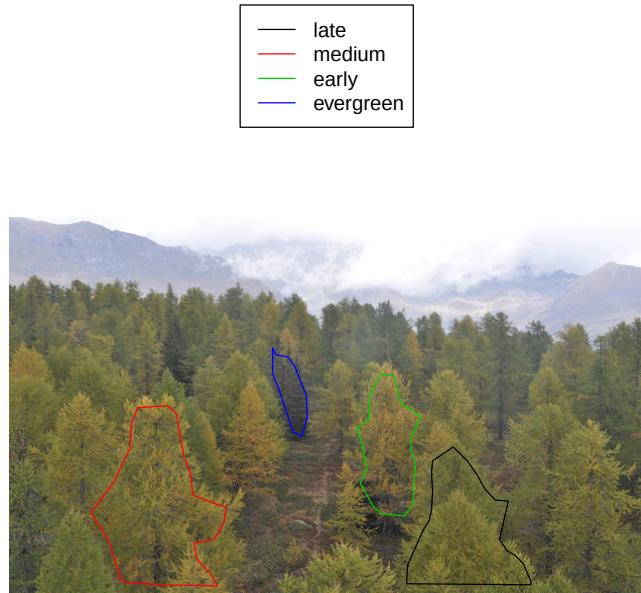


Figure 11: ROIs extracted at the Torgnon Larch site

The function that allows to extract VIs in the pixel-based analysis is the same as in the ROI-averaged approach, except that `spatial` is set to TRUE.

```
> extractVIs(images.2012, folders640.2012$roi,
+             folders640.2012$VI, spatial=TRUE)
```

The structure of the object in output is quite different from the one in the ROI-average approach, and is as follows:

- A list with 4 elements, the 4 ROIs
- Within each element of this list is another list with one element for each time step processed
 - Within each element of this list is a data.frame with three columns corresponding to the digital number of red green and blue, respectively, and there is a row for each pixel in the ROI.

```
> class(VI.data.spatial)
```

```
[1] "list"
```

```

> names(VI.data.spatial)
[1] "late"      "medium"    "early"     "evergreen"

> str(VI.data.spatial$late[[1]])
'data.frame':      11371 obs. of  3 variables:
 $ red   : num  171 196 207 197 214 213 182 190 157 183 ...
 $ green: num  178 200 211 200 217 216 185 193 156 187 ...
 $ blue  : num  186 209 220 205 222 223 192 202 172 198 ...

```

Hence, in the ROI named `late` there are 11371 pixels. The function `spatialGreen` which default is shown below, performs the whole computation of relative indexes, filtering and fitting for each pixel.

```

> spatialGreen(spatial.list, fit, threshold, filters='default',
+ parallel=TRUE, save=FALSE, path=NULL, assign=TRUE)

```

Arguments of `spatialGreen()` are similar to those of `greenProcess()` and `autoFilter()`, in that a `fit` argument allows to choose a fitting method, a `threshold` argument allows to define a threshold method, the argument `filters` controls number and sequence of filtering steps. `spatialGreen` is an extremely computationally intense function, therefore it required particular care to memory saturation. The argument `parallel` allows to parallelize processes into all but one of your processors. Argument `save` provides the option to save each pixel's fit into a small RData object (saved in `path`). This will prevent your memory from being filled up by the creation of a very large object. If you decide to save your temporary fits than you don't need to assign the product of `spatialGreen`. Instead, if `save` is FALSE `assign` must be TRUE otherwise the result of your long analysis will be only printed on screen. The main argument `spatial.list` for the function `spatialGreen()` must contain data from one single ROI. Here is how we used `spatialGreen()` to process the ROI named `late` in our Larch site.

```

> spatialGreen(spatial.list = VI.data.spatial$late, fit = 'gu',
+ threshold = 'gu', filters = 'default', parallel = TRUE,
+ save = TRUE, path = late.path, assign = FALSE)

```

In the code above we decided to save temporary fits from each pixel without constructing a huge object (with `save = TRUE` and `assign = FALSE`). Each pixel of the ROI was therefore processed to extract VIs, filter them, fit the curve from Gu et al (2009), extract thresholds with the same approach. After

have run the function, the folder `late.path` will contain 11371 .Rdata files numbered progressively that represent the fitting results for each pixel. After the analysis, the function `extractParameters()` allows to extract from the fitting all relevant parameters that can be analyzed. Here is an example of extraction of parameters from the ROI named `late` in the larch images shown above:

```
> late.spatial <- extractParameters(path = late.path)
```

Arguments of the function `extractParameters()` include `path`, i.e. the path where temporary files are saved, `list`, alternatively to argument `path` if you have assigned the results from `spatialGreen()` in an object in the workspace, and `update` similar to the function `update` shown before, if you want to apply another threshold method to an already existing fit. Let's look at the structure of the object `late.spatial` to illustrate the parameters extracted from each fitted pixel:

```
> str(late.spatial)

'data.frame':      11371 obs. of  19 variables:
 $ UD        : num  NA 142 142 143 142 ...
 $ SD        : num  NA 154 151 152 156 ...
 $ DD        : num  NA 285 284 282 279 ...
 $ RD        : num  NA 303 307 303 305 ...
 $ maxline   : num  NA 0.46 0.471 0.471 0.474 ...
 $ baseline  : num  NA 0.335 0.335 0.334 0.334 ...
 $ prr       : num  NA 0.01055 0.01512 0.01596 0.00998 ...
 $ psr       : num  NA -0.00627 -0.00498 -0.00588 -0.0044 ...
 $ plateau.slope: num  NA -0.000124 -0.000194 -0.000158 -0.000241 ...
 $ y0        : num  0.335 0.335 0.335 0.335 0.334 ...
 $ a1        : num  0.121 0.125 0.136 0.136 0.141 ...
 $ a2        : num  0.121 0.125 0.136 0.137 0.141 ...
 $ t01       : num  139 134 137 138 134 ...
 $ t02       : num  301 302 306 302 304 ...
 $ b1        : num  2.25 4.25 3.18 3.03 5 ...
 $ b2        : num  1.131 0.986 1.058 1.338 1.1 ...
 $ c1        : num  25.2 19.1 14.5 18.4 13.1 ...
 $ c2        : num  0.0915 0.0631 0.0469 0.0757 0.041 ...
 $ RMSE      : num  0.00886 0.00993 0.01472 0.01016 0.00779 ...
```

It is actually a `data.frame` containing one row for each pixel and 19 columns with the extracted thresholds and the parameters of the curve fitting. Additionally, the root mean square error from the fitting is computed and extracted. Note that missing values in this `data.frame` are not surprising because some fitting may fail to converge, and hence return NA for all parameters and thresholds. Another cause of NA might be that the thresholds do not respect the expected chronology.

An ad hoc function was designed to display the results from the spatial analysis, `plotSpatial()`. Here is an example:

```
> ## roi.data.path <- '.../ROI/roi.data.Rdata'
> ## image.path <- '.../REF/20121004T1400.jpg'
> plotSpatial(data = late.spatial, param = 'UD',
+ roi.data.path = roi.data.path,
+ roi.name = NULL, image.path = image.path,
+ probs=c(0.01, 0.99),
+ plot.density = TRUE, digits = 0, upadj=4)
```

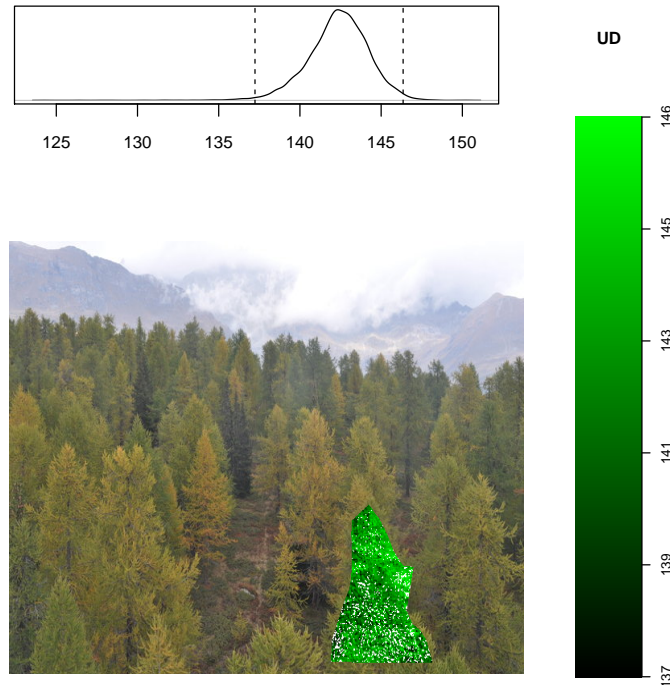


Figure 12: Spatial distribution of Upturn Date in the ROI named late, Torgnon Larch Stand 2012

In the plot shown in fig.12, the spatial distribution of `param` is displayed in a black to green scale superimposed on an image retrieved in `image.path` using the coordinates in `roi.data.path`. The argument `probs` allows to set limits to the range of `param` for plotting based on quantiles. The default is `c(0.01, 0.99)`, meaning that values lower than 1st percentile and higher than 99th percentile are removed before plotting, and will result in white pixels in the plot. The argument `plot.density` allows to plot in the upper area of the plot a density distribution of the plotted parameter. The vertical dashed lines in the density plot represent the quantiles as chosen in `probs` to cut the green color scale. Upturn date in the ROI called late, which includes almost only one individual tree can vary by as much as 10 days (fig.12). Now let's look together at the three ROIs where apparent differences in autumn phenology were observed (i.e. late, medium and early). To show results from more than one ROI with `plotSpatial` one has to first put all results in a list:

```
> all.spatial <- list(early=early.spatial, late=late.spatial,
+                      medium=medium.spatial)
```

Note that it is important that the list be named, with names corresponding to ROI names. Now we are ready to run:

```

> plotSpatial(all.spatial, 'UD', roi.data.path, NULL,
+             image.path, plot.density=TRUE, digits=0,
+             upadj=4)

```

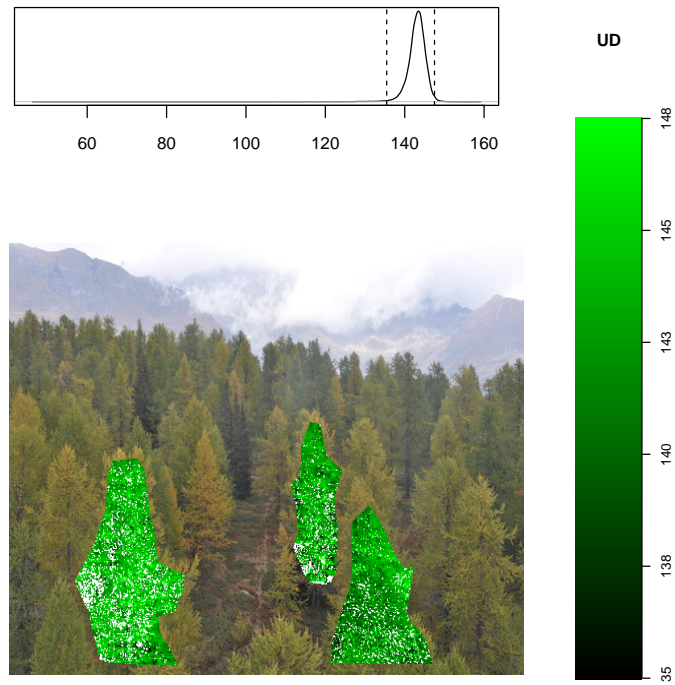


Figure 13: Spatial distribution of Upturn Date in all ROIs, Torgnon Larch Stand 2012

The plot in fig.13 shows no such a big difference in the spring phenology of the selected ROIs, the range of UD being slightly more than 10 days. A different pattern shows up when looking at autumn phenology (fig.13):

```

> plotSpatial(all.spatial, 'RD', roi.data.path, NULL,
+             image.path, plot.density=TRUE, digits=0,
+             upadj=4)

```

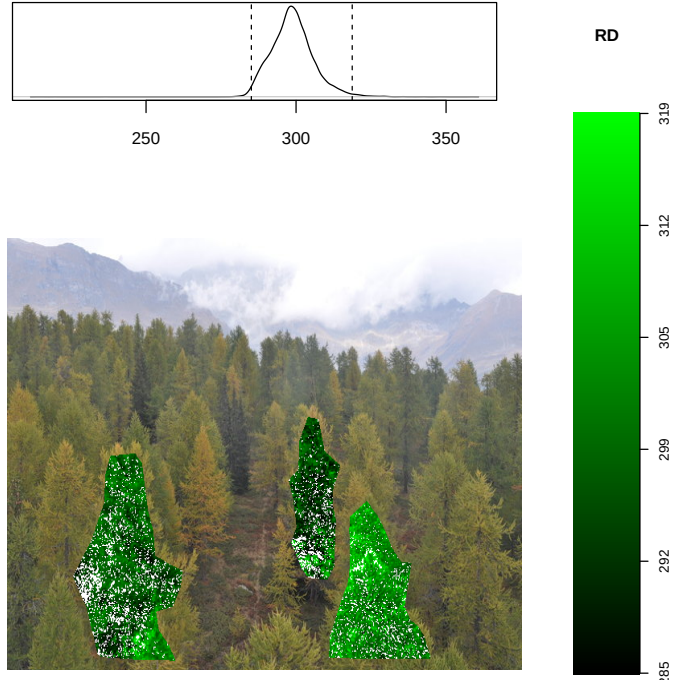


Figure 14: Spatial distribution of Recession Date in all ROIs, Torgnon Larch Stand 2012

Recession date ranges between doy 285 and 320, more than one month difference, clearly showing the expected pattern, with later autumn phenology in the ROI named late (on the right of the plot), earlier in center of the image, and in-between for the ROI on the left (fig.14). The white areas in the figure represent pixels where the fitting failed.

A more heterogeneous autumn than spring phenology in this larch stand was indeed a good tool to show the potential of pixel - based analysis of phenology and its functions. Image resolution is an important issue when deciding to run a pixel-based analysis. To reduce computer time it might be worth decrease image quality. However when the target vegetation is quite far from the camera (as in the larch site) such decrease may dramatically reduce the ability to detect phenological spatial patterns.

To further illustrate the pixel-based analysis we also show results from the Torgnon grassland site. We will use the foreground ROI shown in fig.2. At

this site, the spatial distribution of plant species responds to a peculiar micro-morphology that leads to a different species composition in convex and concave topographies. These different species show also a quite different phenology. In fig.15 we show how upturn date (UD) varies across a relatively small area within the scene of the camera.

```
> plotSpatial(data = spatial.grass, param = 'UD',
+ roi.data.path = grass.roi,
+ roi.name = 'fg', image.path = grass.image.path,
+ probs=c(0.01, 0.99),
+ plot.density = TRUE, digits = 0,
+ upadj=4)
```

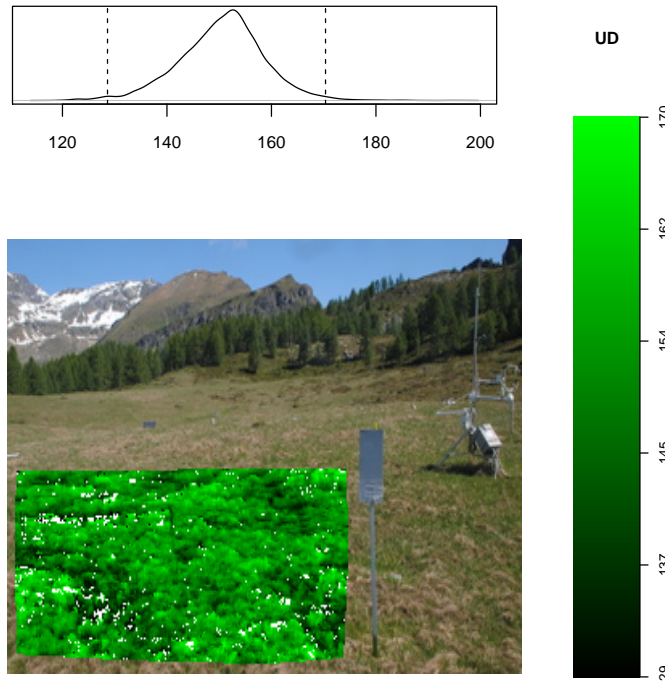


Figure 15: Spatial distribution of Upturn Date in foreground ROI, Torgnon Grassland 2012

There is a range of more than one month in the early spring phenology of plants in Torgnon grassland. We can look at the difference between the beginning (UD) and end (RD) of the growing season to compute the growing season length (gsl), by adding a column to the `spatial.grass` data.frame:

```
> spatial.grass$gsl <- spatial.grass$RD - spatial.grass$UD
```

And plot the results (fig.16)

```
> plotSpatial(data = spatial.grass, param = 'gsl',
+   roi.data.path = grass.roi,
+   roi.name = 'fg', image.path = grass.image.path,
+   probs=c(0.01, 0.99),
+   plot.density = TRUE, digits = 0,
+   upadj=4)
```

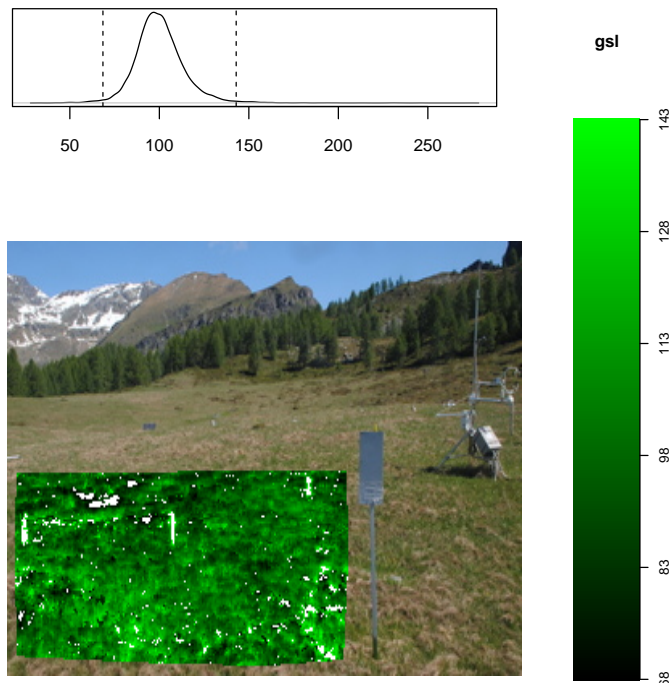


Figure 16: Spatial distribution of the growing season length (gsl) in foreground ROI, Torgnon Grassland 2012

The spatial analysis on Torgnon grassland was run on images of 320 x 214 pixels, i.e. half the resolution of the larch images. Since in the grassland the ROI falls in an area much closer to the camera than for the Larch stand, reducing the image quality did not prevent us from analyzing the spatial pattern of phenology.

10 Summary and future of the package (to be extended)

The phenopix package is currently available for download from the R-forge. The package was tested on approx 300 site-years belonging to the phenocam imagery archive, on the camera network of the project e-pheno and will soon be deployed to process images in the European Network of Flux Towers.

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