

Scripts

Local Image Features **Extraction** — **LIFEx** —

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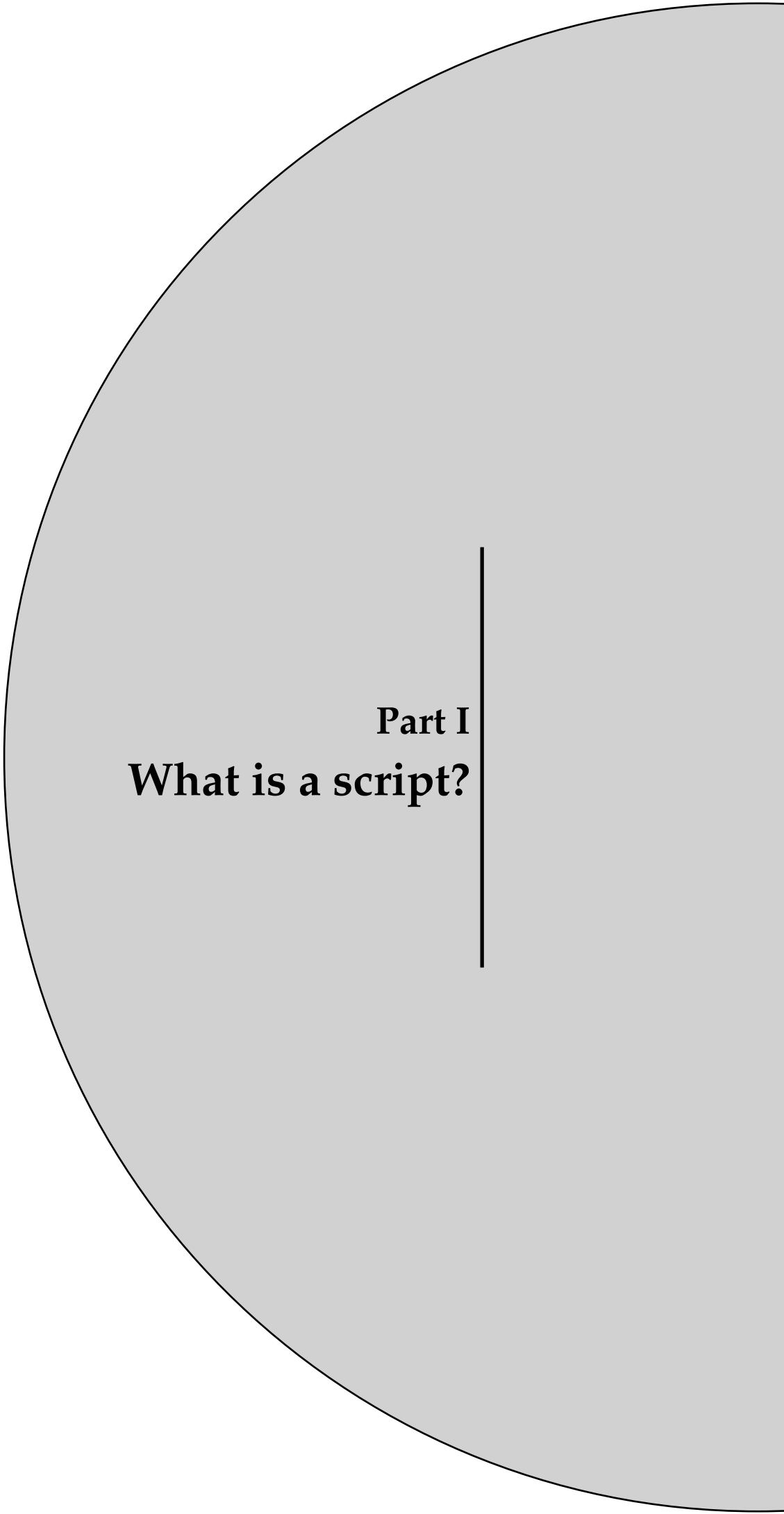
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Part I
What is a script?

Chapter 1

Introduction

1.1 Rationale and what is a script?

Scripting consists of developing instructions that automate processes in the LIFEx application. It is commonly employed to streamline repetitive tasks, perform complex computations, or adapt the user experience for research.

In practice, a script enables execution of all operations and calculations without human intervention. Multiple scripts may be prepared, arbitrarily named, stored, and modified as required, as they are plain text files.

The LIFEx scripting procedure generally consists of the following steps:

- **Script writing:** involves employing the syntax and commands of the scripting language to define executable instructions for the application.
- **Script debugging and testing** consist in executing the script, validating the output, and correcting any errors to ensure conformity with expected behavior.

1.2 General information

- Integration and execution: once functional, the script must be executed within the application, with explicit specifications for timing and mode of execution.
- Script maintenance: ongoing adaptation is required to ensure functionality with evolving application versions, including bug corrections and potential feature extensions.

1.2 General information about writing a script text file

Introduction: A LIFEx script consists of a plain text file structured as a sequence of properties, each defined on a separate line.

Properties: Properties are configuration values expressed as key=value pairs. The key serves as an identifier analogous to a variable, used to access the associated value.

Property example: Example: an application may use the property "*LIFEx.Output.Directory*" to specify the directory in which files are stored.

LIFEx.Output.Directory = /home/user/newDirectory

- key property is "*LIFEx.Output.Directory*" Upper and lower case are not important in the key. This key can also be written : "*LIFEx.output.directory*"
- value property is "*/home/user/newDirectory*"

Construction guidelines of property:

- warning: Keys must be unique identifiers, with a single definition per line.
- Leading spaces preceding a key are disregarded.
- The order of property lines is functionally irrelevant; however, categorization enhances readability for users.
- File paths must employ the / separator consistently, independent of the operating system.
- Comment lines, indicated by the character #, are excluded from application parsing.

1.3 Script execution sequence

The execution of scripts follows a mandatory sequence, as outlined below:

1. Series loading [mandatory]
2. — Series operations [optional]
3. — Series saving [optional]
4. ROI loading [optional]
5. — ROI operations [mandatory when ROI is loaded, otherwise optional]

6. — ROI saving [mandatory when ROI is loaded, otherwise optional]
7. Protocol loading [optional]
8. — MTV protocol operations [optional]
9. — Texture protocol operations [optional]
10. — ROI closing [optional and automatic]
11. ROI closing [optional and automatic]
12. Series closing [automatic]

1.4 How to run a script file

To execute a script, the file must be dragged into the patient image loading panel. Recognition as a valid script requires its size not to exceed 2 MB.

1.5 Can I run several script files at the same time?

Multiple script files may be executed sequentially and automatically by selecting them simultaneously and dragging them into the patient image loading panel.

1.6 Can I automatically build a script from an existing directory structure of patient/study/series with ROI ?

Two script writing approaches are available in LIFEx:

- Direct script writing (for expert users or for complex cases where the directory structure cannot be easily recognized) : This is described throughout this document.
- Automatic detection and generation of a "direct" script from a complete directory structure (This is described in a dedicated section: 2.3).



Part II
General concepts

Chapter 1

How to create a script?

1.1 Write a script

Scripts allow chaining image and ROI loading to produce a consolidated results file in .csv format.

1.2 Script for getting started

Example: the script loads two series (PT0, PT1), each with two associated ROIs (Patient 0 and Patient 1).

The script may be copied into a text file (e.g., script.txt). File paths must be modified to reflect the actual directories: ex. `directory/subDirectory`

1.3 Main remarks

script.txt

```
# Common
# result directory -> mandatory
LIFEx.Output.Directory={directory/subDirectory}

# Patient 0 / Series 0 / ROI 0 / ROI 1
# loading series -> mandatory
LIFEx.Patient0.Series0={directory/subDirectory}/PT0
LIFEx.Patient0.Roi0={directory/subDirectory}/RoiVolume/S0R0.uint16.nii.gz
LIFEx.Patient0.Roi1={directory/subDirectory}/RoiVolume/S0R1.uint16.nii.gz

# Patient 1 / Series 0 / ROI 0 / ROI1
# loading series -> mandatory
LIFEx.Patient1.Series0={directory/subDirectory}/PT1
LIFEx.Patient1.Roi0={directory/subDirectory}/RoiVolume/S1R0.uint16.nii.gz
LIFEx.Patient1.Roi1={directory/subDirectory}/RoiVolume/S1R1.uint16.nii.gz
```

How to create?

1.3 Main remarks relevant to the script writing

- What are the image formats that can be managed using LIFEx scripts?
 - Image files may be provided in NIfTI-1 format (.nii or .nii.gz), requiring full path specification in the script;
 - For DICOM images, the root directory (excluding filenames) must be specified. All files within the directory are then loaded.
- What are the ROI formats that can be managed using LIFEx scripts?
 - ROI formats supported include NIfTI-1 and RTStruct. In both, the full file path must be provided; for RTStruct, all ROIs are imported without exception.
- Syntax of all pathways of files in LIFEx scripts:
 - File paths must exclude accents and spaces.
 - Path syntax is not strictly dependent on the operating system.
 - * Windows paths follow Unit:/Directory/File.extension or Unit:/Your Directory/ for series of DICOM images example C:/Home/Users1/File1
 - * Linux paths: /Your Directory/Your File.extension
 - * macOS paths: /Your Directory/Your File.extension

1.4 Perform operations

Intermediate ROI operations may be executed after Series and ROI loading but prior to feature extraction and results recording:

Main syntax of one operation: The operation syntax is defined as:

```
{key}.Operation0=nameOp0,arg1,arg2,...,argn
```

with "nameOp0" is the title of the button of the ROI action to be performed, arg1 the first argument, arg2 the second argument, ...

Main syntax of many operations to the same Series or ROI: so each uses the result of the previous.

```
{key}.Operation0=nameOp0,arg1,arg2,...,argn|nameOp1,arg1,arg2,...,argn
```

with "nameOp0" is the first operation, "nameOp1" is the second operation.

Operations are executed in the order in which they are written (left to right).

Here's an example:

```
{key}.Operation0=SpatialResampling,2,2,2|Texture,false,false,false,1,3D,Absolute,0.314,192,0,60
```

Explanation:

- series 0 is selected to start this operation 0;
- resample rescaling is realized on series 0
- texture protocol is realized on resample rescaling series result.

Order operations between lines: If you have more than one operation to perform you can describe the actions with an order given by the key_operationN:

```
{key}.Operation0=...
```

```
{key}.Operation1=...
```

```
{key}.Operation2=...
```

How to create?

Chapter 2

Script properties

2.1 Common Properties

Optionally open the results directory at script completion (default)

```
LIFEx.Output.Directory.OpenAtTheEnd=true
```

Define a common output directory for the saving of all series or ROI:

```
LIFEx.Output.Directory={directory/subDirectory/}
```

LIFEx.Password: enables access to experimental or unreleased functions, facilitating controlled feature release.

LIFEx.Password:

```
LIFEx.Password=****
```

2.2 US Properties

LIFEx.GuiUpdate: controls whether the GUI is displayed. Disabling it (false) can accelerate script execution, particularly in server environments (LIFEx>=7.4.1) [true (default) || false]

```
LIFEx.GuiUpdate:
```

```
LIFEx.GuiUpdate=false||true
```

LIFEx.SeriesRoiGuiUpdate: defines whether series and ROI images are rendered during script execution. Disabling (false) accelerates computation (LIFEx>=7.5.6) [true (default) || false]

```
LIFEx.SeriesRoiGuiUpdate:
```

```
LIFEx.SeriesRoiGuiUpdate=false||true
```

LIFEx.MessagesGuiUpdate: determines whether textual messages are displayed. Setting to false suppresses output and increases performance (LIFEx>=7.5.6) [true (default) || false]

```
LIFEx.MessagesGuiUpdate:
```

```
LIFEx.MessagesGuiUpdate=false||true
```

2.2 US Properties

For ultrasound DICOM data, disabling the red, green, and blue channels enables acquisition of the averaged signal alone (all channels are active by default).

The following properties may be added to deactivate individual RGB channels:

```
script.txt
```

```
LIFEx.PropertiesSeriesUS.haveRedChannel=false  
LIFEx.PropertiesSeriesUS.haveGreenChannel=false  
LIFEx.PropertiesSeriesUS.haveBlueChannel=false
```

2.3 Auto-completion with existing structure

It is possible to let LIFEx automatically complete the file tree (Series and ROIs), as well as the operations (on Series and ROIs). This generation is available starting from version 26.3.8.

The sections below describe possible use cases for such generation.

2.3.1 Auto-completion with one series per patient

Unlike the traditional (direct) script construction, when using the BuilderScript functionality, the series and ROI are identified in the keys by letters of the alphabet (A, B, C, ...). If you do not have enough letters, you can combine several: AA, AB, AC, ...

2.3 Auto-completion with existing structure

The numbering of operations remains numeric.

LIFEx will generate an auto-completed script based on a source (builder) script containing the following keys:

```
minimal builder script.txt
```

```
LIFEx.Input.Directory={directory/InputDirectory/}  
LIFEx.Output.Directory={directory/OutputDirectory/}  
LIFEx.SeriesA.OperationA={Series operation}
```

Note that the "PatientN" level has disappeared compared to the direct construction of scripts, since it will be recreated during this procedure.

Neither file names nor directory names matter. The application automatically identifies each file as either an image (Series) or a segmentation (ROI) from the input directory given.

The directory structure does not need to follow a strict format. Automatic completion will work as long as the application can detect images and ROIs in the sub-tree. Please avoid placing multiple image series in the same patient folder, and ensure that each ROI is located close to its corresponding image.

In this case, LIFEx will start by scanning the "Input Directory" and all its subfolders in order to build a detailed script, including the construction of patient0, patient1, ..., patientN. This new script file will be saved with an additional extension, so as not to overwrite the original script file.

Script properties

File tree example (/home/example/input):

```
/
├── PAT0/
│   ├── SERIES CT
│   │   └── series_file0.nii.gz
│   └── ROI
│       ├── roi0_file.nii.gz
│       └── roi1_file.nii.gz
├── PAT1/
│   ├── SERIES CT
│   │   ├── series_file0.dcm
│   │   ├── series_file1.dcm
│   │   └── series_file2.dcm
│   └── ROI
│       ├── roi0_file.nii.gz
│       └── roi1_file.nii.gz
├── PAT2/
│   ├── series_file0.nii.gz
│   ├── roi0_file.nii.gz
│   └── roi1_file.nii.gz
└── PAT3/
    └── TMP0/
        ├── series_file0.nii.gz
        ├── roi0_file.nii.gz
        └── roi1_file.nii.gz
```

2.3 Auto-completion with existing structure

Example of basic script:

builder script.txt

```
LIFEx.Input.Directory=/home/example/input
LIFEx.Output.Directory=/home/example/ouput
LIFEx.SeriesA.MarkerDirectory=SERIES CT
LIFEx.SeriesA.OperationA=spatialresampling,2,2,2|save nii uint16
```

Auto-completion will write a new content script as follows:

generated script.txt

```
lifex.output.directory=/home/example/output

lifex.patient0.roi0=/home/example/input/PAT0/ROI/
lifex.patient0.series0=/home/example/input/PAT0/SERIES CT/series_file0.nii.gz
lifex.patient0.series0.operation0=spatialresampling,2,2,2|save nii uint16

lifex.patient1.roi0=/home/example/input/PAT1/ROI/
lifex.patient1.series0=/home/example/input/PAT1/SERIES CT/
lifex.patient1.series0.operation0=spatialresampling,2,2,2|save nii uint16

lifex.patient2.roi0=/home/example/input/PAT2/roi0_file.nii.gz
lifex.patient2.roi1=/home/example/input/PAT2/roi1_file.nii.gz
lifex.patient2.series0=/home/example/input/PAT2/series_file0.nii.gz
lifex.patient2.series0.operation0=spatialresampling,2,2,2|save nii uint16

lifex.patient3.roi0=/home/example/input/PAT3/TMP0/roi0_file.nii.gz
lifex.patient3.roi1=/home/example/input/PAT3/TMP0/roi1_file.nii.gz
lifex.patient3.series0=/home/example/input/PAT3/TMP0/series_file0.nii.gz
lifex.patient3.series0.operation0=spatialresampling,2,2,2|save nii uint16
```

Windows: On Windows, the ':' character may appear escaped with a backslash (\:). This does not affect the script when it is read back. Otherwise, all \ characters are properly converted to /

2.3.2 Auto-completion with several series per patient

When working with multiple series, the script can specify which reference series should be used, determining the loading order of the series. It is also possible to deselect one series and select another to run the same script on different datasets. Here are some examples to explain this functionality.

• Example 1 : Selection

Selection: Two patients, each with four series and two ROIs. Note that in this case, one of the four series will not be loaded - this introduces the concept of a series selection. One of the two ROI will not be loaded too with the same principle.

2.3 Auto-completion with existing structure

Note: If the concept of "selection" not used is applied, then all series (or ROIs) are selected by default.

File tree example (/home/example/input):

```
/
├── PAT0/
│   ├── CT AC
│   │   └── seriesCT0_file.nii.gz
│   ├── PT0
│   │   └── seriesPT0_file.nii.gz
│   ├── PT1
│   │   └── seriesPT1_file.nii.gz
│   ├── PT2
│   │   └── seriesPT2_file.nii.gz
│   └── ROI
│       ├── roi0_file.nii.gz
│       └── roi1_file.nii.gz
├── PAT1/
│   ├── CT AC
│   │   ├── seriesCT0_file0.dcm
│   │   ├── seriesCT0_file1.dcm
│   │   ├── seriesCT0_file2.dcm
│   │   └── ...
│   ├── PT0
│   │   ├── seriesPT0_file0.dcm
│   │   ├── seriesPT0_file1.dcm
│   │   └── seriesPT0_file2.dcm
│   ├── PT1
│   │   ├── seriesPT1_file0.dcm
│   │   ├── seriesPT1_file1.dcm
│   │   └── seriesPT1_file2.dcm
│   ├── PT2
│   │   ├── seriesPT2_file0.dcm
│   │   ├── seriesPT2_file1.dcm
│   │   └── seriesPT2_file2.dcm
│   └── ROI
│       ├── roi0_file.nii.gz
│       └── roi1_file.nii.gz
```

Script properties

Original script as follows:

2.3 Auto-completion with existing structure

builder script.txt

```
LIFEx.Input.Directory=/home/example/input  
LIFEx.Output.Directory=/home/example/output
```

```
LIFEx.SeriesA.OperationA=P  
LIFEx.SeriesB.OperationA=Q  
LIFEx.SeriesC.OperationA=R  
LIFEx.SeriesD.OperationA=S  
LIFEx.SeriesA.MarkerDirectory=CT|true  
LIFEx.SeriesB.MarkerDirectory=PT0|true  
LIFEx.SeriesC.MarkerDirectory=PT1|false  
LIFEx.SeriesD.MarkerDirectory=PT2|true
```

```
LIFEx.RoiA.OperationA=T  
LIFEx.RoiB.OperationA=U  
LIFEx.RoiA.MarkerDirectory=roi0|true  
LIFEx.RoiB.MarkerDirectory=roi1|false
```

Script properties

Replace the operations P, Q, R, S, T, U with the ones you want.

2.3 Auto-completion with existing structure

Auto-completion will write a new content script as follows:

```
generated script.txt

LIFEx.Output.Directory=/home/example/output

LIFEx.Patient0.Roi0.Operation0=...
LIFEx.Patient0.Roi0=PAT0/ROI/roi0_file.nii.gz
LIFEx.Patient0.Series0.Operation0=P
LIFEx.Patient0.Series1.Operation0=Q
LIFEx.Patient0.Series2.Operation0=S
LIFEx.Patient0.Series0=PAT0/CT AC/seriesCT0_file.nii.gz
LIFEx.Patient0.Series1=PAT0/PT0/seriesPT0_file.nii.gz
LIFEx.Patient0.Series2=PAT0/PT2/seriesPT2_file.nii.gz

LIFEx.Patient1.Roi0.Operation0=...
LIFEx.Patient1.Roi0=PAT1/ROI/roi0_file.nii.gz
LIFEx.Patient1.Series0.Operation0=P
LIFEx.Patient1.Series1.Operation0=Q
LIFEx.Patient1.Series2.Operation0=S
LIFEx.Patient1.Series0=PAT1/CT AC
LIFEx.Patient1.Series1=PAT1/PT0
LIFEx.Patient1.Series2=PAT1/PT2
```

Script properties

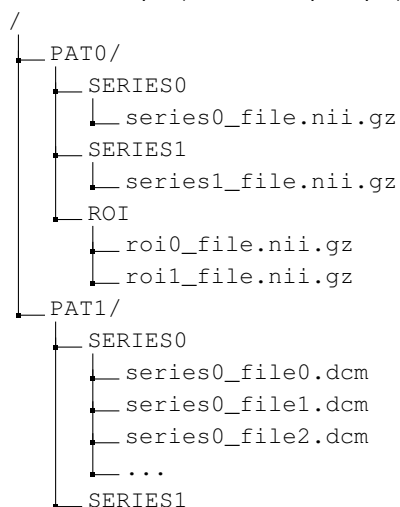
Note 1: Note that any reference to ROI1 has disappeared because the MarkerDirectory key "roi1|false" is set to false.

Note 2: Note that any reference to Series PT1 has disappeared because the MarkerDirectory key "PT1|false" is set to false.

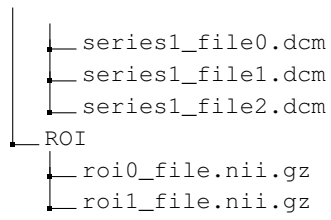
• Example 2: reference

Two patients, each with two series and two ROIs. In this case, both series are loaded but only one of them as reference series.

File tree example (/home/example/input):



2.3 Auto-completion with existing structure



Original script as follows:

```
builder script.txt
```

```
LIFEx.Input.Directory=/home/example/input/
LIFEx.Output.Directory=/home/example/output
```

```
LIFEx.SeriesA.MarkerDirectory=SERIES0|true
LIFEx.SeriesA.OperationA=...
```

```
LIFEx.SeriesB.MarkerDirectory=SERIES1|true|reference
LIFEx.SeriesB.OperationA=...
```

```
LIFEx.RoiA.OperationA=...
LIFEx.RoiB.OperationA=...
```

Script properties

2.3 Auto-completion with existing structure

Auto-completion will write a new content script as follows:

```
generated script.txt

LIFEx.Output.Directory=/home/example/output

LIFEx.Patient0.Series0.Operation0=...
LIFEx.Patient0.Series1.Operation0=...
LIFEx.Patient0.Roi0.Operation0=...
LIFEx.Patient0.Roi0=PAT0/ROI
LIFEx.Patient0.Series0=PAT0/SERIES1/series1_file.nii.gz
LIFEx.Patient0.Series1=PAT0/SERIES0/series0_file.nii.gz

LIFEx.Patient1.Series0.Operation0=...
LIFEx.Patient1.Series1.Operation0=...
LIFEx.Patient1.Roi0.Operation0=...
LIFEx.Patient1.Roi0=PAT1/ROI
LIFEx.Patient1.Series0=PAT1/SERIES1/
LIFEx.Patient1.Series1=PAT1/SERIES0/
```

Script properties

Note: The series "SERIES1" becomes the reference series (as specified by the key/value LIFEx.SeriesB.MarkerDirectory=...|reference and receives index 0 (and loaded in first).

2.3.3 Execution after auto-completion (v>=26.3.8)

It is possible to not request the execution of the generated script immediately after its construction. To do this, add the following key/value to the initial script:

```
script.txt

LIFEx.ExecutionAfterBuilderConstruction=false
```

In this case, the execution is triggered immediately. This is equivalent to dragging and dropping the generated script file into the application.

This feature is optional and is not enabled by default. If this line does not exist with the value "true", the script is generated but not executed.

Chapter 3

Series operations

3.1 Syntax examples

Store series in anonymized DICOM format:

```
LIFEx.Patient0.Series0.Operation0=Save anonymous dcm
```

Note: When writing DICOM files via a script, the default format is DcmEnhanced. Generating DicomLegacy format is not supported at this time.

Standardize series units to SUVbw:

```
LIFEx.Patient0.Series0.Operation0=SUVbw  
available units: suvbw || suvlbm_james120 || suvlbm_james128 || suvlbm_janma || kbqml ||  
cpxvx
```

3.2 Series common operations

Standardize series units to SUVbw prior to SpatialResampling:

```
LIFEx.Patient0.Series0.Operation0=SUVbw|SpatialResampling,2,2,2
```

Apply 2x2x2 resampling prior to saving in NIfTI format:

```
LIFEx.Patient0.Series0.Operation0=SpatialResampling,2,2,2|save nii uint16
```

Export results in anonymized NIfTI format:

```
LIFEx.Patient0.Series0.Operation0=Save Anonymous nii uint16
```

Define the output directory for series export:

```
LIFEx.Output.Directory={directory/subDirectory/}
```

3.2 Series common operations

Define a global operation list to be applied across all loaded series:

```
LIFEx.Operation0=list of operations to be performed on all series
```

3.3 Series operations

3.3.1 Save DICOM Series

Note: DICOM export requires the input series to be in DICOM format.

There are two main types of DICOM objects used to represent medical images: so-called Legacy objects and Enhanced objects:

- Legacy DICOM objects correspond to the historical format, in which each image is typically stored as an independent instance containing its own metadata. This model is simple and widely compatible with existing systems, but it can lead to redundancy and limitations when handling volumes or complex image series.

script.txt

```
# loading series -> mandatory  
LIFEx.Patient0.Series0={directory/subDirectory}/PT0  
LIFEx.Patient0.Series0.Operation0=Save DCM Legacy
```

- Enhanced DICOM objects, introduced more recently in the standard, use a multi-frame structure that allows multiple images to be grouped within a single instance while sharing common information. This approach improves metadata consistency, reduces information duplication, and facilitates the management of advanced data such as dynamic, multi-parametric, or volumetric acquisitions.

```
script.txt
```

```
# loading series -> mandatory
LIFEx.Patient0.Series0={directory/subDirectory}/PT0
LIFEx.Patient0.Series0.Operation0=Save DCM Enhanced
```

3.3.2 Save anonymous DICOM Series

Export anonymized series in DICOM-Enhanced format - provided original input is DICOM:

```
LIFEx.Patient0.Series0.Operation0=Save DCM Enhanced,PatientID->xxx,PatientName->xxx
```

- You can update the value of a DICOM field (new value xxx) using the following format:
DicomFieldName->xxx

- A field that you want anonymized must be replaced using the following syntax:
DicomFieldName->REPLACED

Below is a non-exhaustive list that serves as a baseline for anonymization:

```
PerformingPhysicianName->REPLACED,
ReferringPhysicianName->REPLACED,
NameOfPhysiciansReadingStudy->REPLACED,
InstitutionName->REPLACED,
InstitutionAddress->REPLACED,
InstitutionalDepartmentName->REPLACED,
OperatorsName->REPLACED,
PersonAddress->REPLACED,
PersonTelephoneNumbers->REPLACED,
PersonTelecomInformation->REPLACED,
PatientName->REPLACED,
PatientID->REPLACED,
PatientBirthDate->REPLACED,
PatientState->REPLACED,
IssuerOfPatientID->REPLACED,
AdmittingDiagnosesDescription->REPLACED,
MedicalAlerts->REPLACED,
Allergies->REPLACED,
AdditionalPatientHistory->REPLACED,
ReasonforVisit->REPLACED,
ServiceEpisodeDescription->REPLACED,
ProtocolName->REPLACED
```

Syntax and case sensitivity are important when defining DICOM keys.

You are encouraged (and expected) to review and adapt it to meet your own requirements. Field names ("DicomFieldName") are available on the website:

<https://dicom.innolitics.com/>

Series operations

3.3 Series operations

3.3.3 Save series in others formats

Export series in NRRD format:

```
LIFEx.Patient0.Series0.Operation0=Save nrrd
```

Export series in NIfTI format (float 32 bits):

```
LIFEx.Patient0.Series0.Operation0=Save nii float32
```

Export series in NIfTI format (uint 16 bits):

```
LIFEx.Patient0.Series0.Operation0=Save nii
```

Export series in ECAT format: (.v):

```
LIFEx.Patient0.Series0.Operation0=Save ecats
```

Export MIP 3D Plans: Coronal - Sagittal - Axial in nifti format (float32):

```
LIFEx.Patient0.Series0.Operation0=Save MIP 3D Plans
```

Export dynamic representation as video: in video format (mp4):

```
LIFEx.Patient0.Series0.Operation0=Save mp4
```

Change output directory of Series saving:

```
LIFEx.Patient0.Series0.Operation0.Output.Directory={directory/subDirectory/}
```

Change Unit of Y axis of series:

Unit of Y axis of series, choose between:
kBq/mL || SUVbw || SUVlbm_morgan
|| SUVlbm_james || SUVlbm_janma || SUVibw || SUVbsa || Cpx/vx || Gy/vx || %/vx
|| .# class || .# k Pa || HU || T || mL/100g || mL/100g/min || sec || RC (.###) || min-1
|| Proba || # || #.# || #.## || #.### || #.####

```
LIFEx.Patient0.Series0.Operation0=SUVbw
```

Crop3D series:

Coordinates are expressed in voxels (on referenced series). You can obtain these two crop coordinates in the interface via the Series/Crop menu and the following fields: "start corner" and "end corner".

```
LIFEx.Patient0.Series0.Operation0=Crop3D,coorX1,coorY1,coorZ1,coorX2,coorY2,coorZ2
```

Crop2D series:

Coordinates are expressed in voxels (on referenced series). You can obtain these two crop coordinates in the interface via the Series/Crop menu and the following fields: "start corner" and "end corner".

LIFEx.Patient0.Series0.Operation0=**Crop2D,coorX1,coorY1,coorX2,coorY2**

Spatial resampling with specified voxel dimensions:

LIFEx.Patient0.Series0.Operation0=**SpatialResampling,xSpacing,ySpacing,zSpacing**

Example with spacing (x=3 mm, y=3mm, z=3mm):

LIFEx.Patient0.Series0.Operation0=SpatialResampling,3.0,3.0,3.0

Series operations

Perform linear combination of two series with user-defined coefficients:

LIFEx.Patient0.Series0.Operation0=**SeriesOperator, CoefA, SeriesA, OperatorName, CoefB, SeriesB**

OperatorName between: ADDITION || SUBTRACT || AVERAGE || DIVIDE || MULTIPLY || MAXIMUM || MINIMUM || AFFINE_TRANSFORM

Example of $S3 = 0.6 * S1 + 0.4 * S2$ then save S3 series result:

LIFEx.Patient0.Series0=S0file.nii.gz

LIFEx.Patient0.Series1=S1file.nii.gz

LIFEx.Patient0.Series0.Operation0=**SeriesOperator,0.6,S1,ADDITION,0.4,S2|SaveNii**

Apply LaplacienOfGaussian spatial filtering with defined kernel size and dimensionality:

LIFEx.Patient0.Series0.Operation0=**LaplacienOfGaussianFilter,SigmaX,SigmaY,SigmaZ, CalculationDimension,WholeBody,PaddingMethod**

Example with Sigma (x=2 mm, y=2mm, z=2mm), 3d calculation dimension, wholebody enable and reflect padding method:

LIFEx.Patient0.Series0.Operation0=LaplacienOfGaussianFilter,2,2,2,3d,true,reflect

- float value Sigma is the FWHM/2 of kernels size in millimeter.

- 3d calculation dimension is between [2d, 3d]

- wholebody is always true in script

- padding method is between [reflect, periodic, edge, zero]

3.3 Series operations

Apply Gaussian spatial filtering with defined kernel size and dimensionality:

LIFEx.Patient0.Series0.Operation0=**GaussianFilter,SigmaX,SigmaY,SigmaZ,CalculationDimension,WholeBody,PaddingMethod**

Example with Sigma (x=2 mm, y=2mm, z=2mm), 3d calculation dimension, wholebody enable and reflect padding method:

LIFEx.Patient0.Series0.Operation0=GaussianFilter,2,2,2,3d,true,reflect

- float value Sigma is the FWHM/2 of kernels size in millimeter.
- 3d calculation dimension is between [2d, 3d]
- wholebody is always true in script
- padding method is between [reflect, periodic, edge, zero]

Apply spatial mean filtering with defined kernel size and dimensionality:

LIFEx.Patient0.Series0.Operation0=**MeanFilter,KernelDiameterSizeVx,CalculationDimension,WholeBody,PaddingMethod**

Example with kernelDiameterSizeVx 3 vx, 3d calculation dimension, wholebody enable and reflect padding method:

LIFEx.Patient0.Series0.Operation0=MeanFilter,3,3d,true,reflect

- float value sigma is the FWHM/2 of kernels size in millimeter.
- integer value diameter kernel size between [3, 5, 7, 9, 11, 13, 15]
- 3d calculation dimension is between [2d, 3d]
- wholebody is always true in script
- padding method is between [reflect, periodic, edge, zero]

Apply wavelet-based decomposition filter with user-defined parameters:

LIFEx.Patient0.Series0.Operation0=**WaveletsFilter,TransformAlongZAxis,TransformAlongYXAxis,FamilyName,Order,Level,PaddingMethod**

Example with TransformAlongZAxis enable, TransformAlongYXAxis enable, coiflet family-Name, 1 order and 1 level:

LIFEx.Patient0.Series0.Operation0=WaveletsFilter,true,true,coiflets,1,1,reflect

- transformAlongZAxis is between [true, false]
- transformAlongYXAxis is always true
- familyName is between [coiflets, biorthogonal, daubechies, haar, reverse biorthogonal, symlets]
- daubechies order is between [1, ..., 38]
- symlets order is between [2, ..., 20]
- coiflets order is between [1, ..., 17]
- reverse/biorthogonal order is between [11, 13, 15, 22, 24, 26, 28, 31, 33, 35, 37, 39, 44, 55, 68]
- haar order is equal [1]
- Level is equal [1]
- padding method is between [reflect, periodic, edge, zero]

Apply Laws' texture energy filters with selected kernel family:

LIFEx.Patient0.Series0.Operation0=LawsFilter,KernelSize,WholeBody,PaddingMethod

Example with (l3, l3, l3 Kernel size, WholeBody enable, and reflect Padding method

LIFEx.Patient0.Series0.Operation0=LawsFilter,l3,l3,l3,true,reflect

- kernel size is between [l3, l5, e3, e5, s3, s5, w5, r5]

with l3: level (1, 2, 1)

with l5: level (1, 4, 6, 4, 1)

with e3: edge (-1, 0, 1)

with e5: edge (-1, -2, 0, 2, 1)

with s3: spot (-1, 2, -1)

with s5: spot (-1, 0, 2, 0, -1)

with w5: wave (-1, 2, 0, -2, 1)

with r5: ripple (1, -4, 6, -4, 1)

- wholebody is always true in script

- padding method is between [reflect, periodic, edge, zero]

Series operations

Chapter 4

ROI operations

4.1 Syntax examples

Example: apply an absolute threshold between 2.5 and 50:

```
LIFEx.Patient0.Roi0.Operation0=n,2.5,5
```

Example: absolute thresholding (2.5-50) followed by ROI export in NIfTI format:

```
LIFEx.Patient0.Roi0.Operation0=n,2.5,50|Save nii
```

Example: subtraction of two loaded ROIs - followed by saving of the resulting ROI:

```
LIFEx.Patient0.Roi0.Operation0=SelectedAllRoi|Subtract|SelectedAllRoi|Save nii
```

4.2 ROI operations from threshold menu

4.2 ROI operations from threshold menu

Apply absolute thresholding (min-max range):

LIFEx.Patient0.Roi0.Operation0=**n,ValueOfMinThreshold,ValueOfMaxThreshold**

Apply relative thresholding based on percentage:

LIFEx.Patient0.Roi0.Operation0=**n%,ValueOfPercentThreshold**

Fixed 40% relative threshold:

LIFEx.Patient0.Roi0.Operation0=**40%**

Fixed 70% relative threshold:

LIFEx.Patient0.Roi0.Operation0=**70%**

Apply peak-based thresholding:

LIFEx.Patient0.Roi0.Operation0=**Peak**

Apply Nestle method with specified threshold:

LIFEx.Patient0.Roi0.Operation0=**Nestle,ValueOfThreshold**

Apply PERCIST thresholding (requires preloaded Liver ROI):

LIFEx.Patient0.Roi0.Operation0=**PERCIST**

4.3 ROI operations from file/edit menu

Retain a single ROI from the loaded set

LIFEx.Patient0.Roi0.Operation0=**KeepOne**

Divide ROI into separate components

LIFEx.Patient0.Roi0.Operation0=**Split**

Union -> Union of all ROIs

LIFEx.Patient0.Roi0.Operation0=**SelectedAllRoi|Union**

Intersection -> Intersection of ROIs

LIFEx.Patient0.Roi0.Operation0=**SelectedAllRoi|Intersection**

4.3 ROI operations from file/edit menu

Subtract -> Subtraction (with only 2 ROIs loaded)

LIFEx.Patient0.Roi0.Operation0=**SelectedAllRoi|Subtract**

Morphological dilation with kernel size n voxels

LIFEx.Patient0.Roi0.Operation0=**Dilate,n**
- n is the number of voxels of length of kernel (vx)

Morphological erosion with kernel size n voxels

LIFEx.Patient0.Roi0.Operation0=**Erode,n**
- n is the number of voxels of length of kernel (vx)

Ring ROI operation

Ring ROI operation generates a hollow structure of thickness n mm (half applied as lateral thickness).

Example: *Ring,10* produces a 10 mm kernel with 5 mm lateral extent.

LIFEx.Patient0.Roi0.Operation0=**Ring,n**

Example with 10 millimeters (= 5 millimeters of lateral extent):

LIFEx.Patient0.Roi0.Operation0=**Ring,10**

Select all loaded ROIs

LIFEx.Patient0.Roi0.Operation0=**SelectedAllRoi**

Export ROI as NIFTI file:

LIFEx.Patient0.Roi0.Operation0=**Save nii**

Export ROI as DICOM file:

LIFEx.Patient0.Roi0.Operation0=**Save dcm**

Export ROI as CSV (comma-separated values):

LIFEx.Patient0.Roi0.Operation0=**Save csv**

Export ROI in Turku PET Centre format (.dft):

LIFEx.Patient0.Roi0.Operation0=**Save dft**

Placeholder for Recovery Coefficient export (not implemented):

LIFEx.Patient0.Roi0.Operation0=**Save dftrc**

ROI operations

4.4 ROI operations from measure menu

Export ROI as NRRD (Pmod-compatible):

```
LIFEx.Patient0.Roi0.Operation0=Save nrrd
```

Define output directory for ROI saving:

```
LIFEx.Patient0.Roi0.Operation0.Output.Directory={directory/subDirectory/}
```

Define a global output directory for saving all ROIs:

```
LIFEx.Output.Directory={directory/subDirectory/}
```

4.4 ROI operations from measure menu

Compare ROI operation

Compare ROI operation generates a comparison result file. The comparison is carried out two by two, on all the ROIs loaded in the application.

```
LIFEx.Patient0.Roi0.Operation0=Compare
```

4.5 ROI operations from sort menu

Quarter division for labeling ROI:

```
LIFEx.Patient0.Roi0.Operation0="Quarter Division"
```

Complex example:

```
LIFEx.Patient0.Roi0.Operation0=Quarter Division|Selected All Roi|Save nii
```

Chapter 5

Protocol operations

5.1 Protocol TEXTURE operations

5.1.1 Rationale

For large cohorts or multiple ROIs per patient, scripting enables automated extraction of indices (SUV, volume, histogram-based, textural) without user intervention. Prior ROI preparation is required. Scripts may be executed repeatedly with varied parameter sets to assess their influence on extracted features.

Example: For 100 patients each with 10 ROIs, the *nbGreyLevels* parameter may be varied (64, 128, 256) to evaluate its impact, requiring three separate script executions., see the "Can I run several script files at once?" p.11.

Another example: You have 10 ROI per patient and a set of 100 patients to be processed. You are interested in studying the impact of the *SpatialResampling* parameter on the 10*100 ROI, by setting this parameter to 2x2x2 mm and to 4x4x4 mm. You can write and run a first script by setting the *SpatialResampling* parameter to 2x2x2 mm. Then duplicate the script and change 2x2x2 mm by 4x4x4 mm to produce a second script to be run. Running the two scripts will calculate all indices for you automatically and store them in csv files. Setting the spatial resampling parameter to 2x2x2 mm in the script file can be done as in series operations chapter.

5.1 Protocol TEXTURE operations

A preliminary ROI verification (CheckTex) ensures suitability for texture analysis, avoiding time-consuming invalid computations. This will check whether the ROI includes a single cluster and contain a number of voxels greater than that required for consistent textural feature calculation. If one of these two conditions is not met, a warning message will be displayed.

All these lines contain the series paths, ROIs and parameters needed to redo the texture analysis you've just designed in the user interface. You can then drag this file into the application's series/images/scripts reading area to run the whole procedure. You can also use this file as a starting point for modifying certain parameters in order to re-execute a new analysis under the same conditions.

5.1.2 Explanation of field contents

Check: Setting Check=true enables a fast preliminary computation restricted to simple features (voxel count, min, mean, max).

KeepTheLargestCluster: The KeepTheLargestCluster option retains only the largest connected component within an ROI, discarding smaller clusters.

IsEnabledRoiUnion: merges all segmented regions into a single ROI prior to texture calculation.

BinSize and NbGreyLevels (how set the quantization): Grey-level discretization can be parameterized via BinSize or NbGreyLevels, with automatic computation of the unset parameter.

For instance:

if $BinSize = 0$ then $BinSize = (boundMax - boundMin) / (nbGreyLevels - 1)$
if $nbGreyLevels = 0$ then $nbGreyLevels = ((boundMax - boundMin) / binSize) + 1$
if $BinSize = 0$ and $nbGreyLevels = 0$ then $nbGreyLevels = 64$ by default
 $boundMin$ and $boundMax$ are the minimum and maximum values in the processed ROI.

Discretization has been set with:

$$discretizedValue = \text{floor}\left(\left(nbGreyLevels * \frac{originalValue - boundMin}{boundMax - boundMin}\right)\right) + 1$$

• Have the minimum and maximum bounds of a whole cohort of patients in a script (without texture calculation)

It is easy to find the BoundMax of all the ROI of all your patients in order to put this bound in the final script. This has to be done in 2 steps with exactly the same script (by changing one parameter). Here is the process.

- step 1: change this value to "true" on the Check key. It allows calculating the min, max bounds of all the ROI in a single pass of the script.
- step 2: run the script a first time to have only bound results; You then retrieve the max of the max of all the ROIs and correct the value of the MaxBound key in the script file
- step 3: change to false value of Check key in the script file
- step 4: run the script a second time to have all feature results

5.1.3 Automatic texture script creation

Since version 7.6.0, LIFEx is capable of automatically generating texture scripts based on GUI-defined protocols. To do so, follow the standard texture protocol in the interface. However, instead of executing the texture calculations by pressing the run button, you can generate the corresponding script by pressing the script button (to the left of the run button).

A text file will open with the contents of the script written for you, taking into account the interface's texture parameters. This script also includes the loading of previously opened Series and ROI.

The generated script can be reloaded into a new LIFEx session for automatic execution of texture analysis.

5.1.4 Operations

Operations between { } are optional. If they are added, do not place the braces { } in the script.

- Texture protocol execution using global ROI and absolute discretization scheme:

template:

```
template: LIFEx.Patient0.Series0.Operation0=Texture, KeepTheLargestCluster, Check,
isEnabledRoiUnion, DistanceWithNeighbours, DimensionProcessing, Absolute, Bin-
Size, NbGreyLevels, MinBound, MaxBound
{,xSpatialResampling, ySpatialResampling, zSpatialResampling}
{, filterName, ..., ...}
see: filters in "Available Series operations" section
```

example1: LIFEx.Patient0.Series0.Operation0=Texture,true,false,false,1,3D,**Absolute**,0,64,0,20
 Texture with KeepTheLargestCluster=true
 and with check=false
 and with isEnabledRoiUnion=false
 and with DistanceWithNeighbours=1 and with DimensionProcessing=3D
 and with Absolute discretization
 and with BinSize=0
 and with NbGreyLevels=64
 and with MinBound=0
 and with MaxBound=20

example2: LIFEx.Patient0.Series0.Operation0=Texture,true,false,false,1,3D,**Absolute**,0,64,0,20,2.5,2.5,2.5
 Texture with KeepTheLargestCluster=true
 and with check=false
 and with isEnabledRoiUnion=false
 and with DistanceWithNeighbours=1 and with DimensionProcessing=3D
 and with Absolute discretization
 and with BinSize=0
 and with NbGreyLevels=64
 and with MinBound=0
 and with MaxBound=20
 and with xSpatialResampling=2.5 mm
 and with ySpatialResampling=2.5 mm
 and with zSpatialResampling=2.5 mm

example3: LIFEx.Patient0.Series0.Operation0=Texture,true,false,false,1,3D,**Absolute**,0,64,0,20,2.5,2.5,2.5,
 LaplacianOfGaussianFilter,2.000,2.000,2.000,3d,WholeBody,reflect
 Texture with KeepTheLargestCluster=true
 and with check=false
 and with isEnabledRoiUnion=false
 and with DistanceWithNeighbours=1 and with DimensionProcessing=3D
 and with Absolute discretization
 and with BinSize=0
 and with NbGreyLevels=64

5.1 Protocol TEXTURE operations

and with MinBound=0
and with MaxBound=20
and with xSpatialResampling=2.5 mm
and with ySpatialResampling=2.5 mm
and with zSpatialResampling=2.5 mm
and with LaplacianOfGaussianFilter filter

- Execution with relative mean-standard deviation discretization strategy (global ROI):

template:

```
LIFEx.Patient0.Series0.Operation0=Texture, KeepTheLargestCluster, Check, isEnabledRoiUnion, DistanceWithNeighbours, DimensionProcessing, RelativeMeanSd, BinSize, NbGreyLevels, MinBound, MaxBound  
{, xSpatialResampling, ySpatialResampling, zSpatialResampling}  
{, filterName, ..., ...}  
see: filters in "Available Series operations" section
```

example: LIFEx.Patient0.Series0.Operation0=Texture,true,false,false,1,3D,**RelativeMeanSd**,0,64
Texture with KeepTheLargestCluster=true, check=false, isEnabledRoiUnion=false
and with DistanceWithNeighbours=1
and with DimensionProcessing=3D
and with RelativeMeanSd discretization
and with BinSize=0
and with NbGreyLevels=64

- Execution with relative min-max discretization strategy (global ROI):

template:

```
LIFEx.Patient0.Series0.Operation0=Texture, KeepTheLargestCluster, Check, isEnabledRoiUnion, DistanceWithNeighbours, DimensionProcessing, RelativeMinMax, BinSize, NbGreyLevels, MinBound, MaxBound  
{, xSpatialResampling, ySpatialResampling, zSpatialResampling}  
{, filterName, ..., ...}  
see: filters in "Available Series operations" section
```

example: LIFEx.Patient0.Series0.Operation0=Texture,true,false,false,1,3D,**RelativeMinMax**,0,64
Texture with KeepTheLargestCluster=true, check=false, isEnabledRoiUnion=false
and with DistanceWithNeighbours=1
and with DimensionProcessing=3D
and with RelativeMinMax discretization
and with BinSize=0
and with NbGreyLevels=64

- TextureMap protocol enables voxel-wise feature extraction with kernel-based processing and specified model (WholeBody, VoxelsOfRoi, BoundingBoxOfRoi) (local Map):

template:

```
LIFEx.Patient0.Series0.Operation0=TextureMap, KernelProcessing, MapModel,
BinSize, NbGreyLevels, MinBound, MaxBound

with KernelProcessing between 3 || 5 || 7
with MapModel between WholeBody||VoxelsOfRoi||BoundingBoxOfRoi
Don't forget the ROI loading line if you're using MapModel VoxelsOfRoi||BoundingBoxOfRoi
```

example: LIFEx.Patient0.Series0.Operation0=**TextureMap,3,WholeBody,0,64,0,20**
 TextureMap with KernelProcessing=3
 and with MapModel=WholeBody
 and with BinSize=0
 and with NbGreyLevels=64
 and with MinBound=0
 and with MaxBound=20

5.2 MTV Protocol operations

5.2.1 Introduction

MTV (Metabolic Tumor Volume) computation can be automated through scripts, chaining image and ROI loading into consolidated .csv outputs.

5.2.2 Operations

- MTV protocol execution is invoked via the operation Mtv, KeepTheLargestCluster.:

template:

```
LIFEx.Patient0.Series0.Operation0=Mtv,KeepTheLargestCluster
```

example: LIFEx.Patient0.Series0.Operation0=**Mtv,false**

- KeepTheLargestCluster on MTV: defines whether only the largest cluster should be kept when an ROI contains several clusters.
 If the value of this key is true, then only the largest cluster is kept (the others are deleted during the calculation).
 If this value is false, then the ROI remains composed of several clusters if this is the case.
 -> In the MTV protocol, we strongly recommend leaving this parameter at **false**, otherwise all clusters will not be taken into account when calculating the MTV (this depends mainly on the goal set).

5.2.3 Script example

This section shows a script example of MTV protocol.

You can copy/paste this script into a text file named script.txt (for example). Don't forget to change some information: ex. directory/subDirectory

Protocol operations

5.2 MTV Protocol operations

script.txt

```
# result file -> mandatory
LIFEx.Output.Directory={directory/subDirectory}

# Patient0 / Series / ROI
# loading series -> mandatory
LIFEx.Patient0.Series0={directory/subDirectory}/PT0
LIFEx.Patient0.Series0.Operation0=Mtv,false
# loading ROI -> mandatory
LIFEx.Patient0.Roi0={directory/subDirectory}/RoiVolume/R1.uint16.nii.gz
LIFEx.Patient0.Roi1={directory/subDirectory}/RoiVolume/R2.uint16.nii.gz

# Patient 1 / Series
# loading series -> mandatory
LIFEx.Patient1.Series0={directory/subDirectory}/PT1
LIFEx.Patient1.Series0.Operation0=Mtv,false
# loading ROI -> mandatory
LIFEx.Patient1.Roi0={directory/subDirectory}/RoiVolume/R1.uint16.nii.gz
LIFEx.Patient1.Roi1={directory/subDirectory}/RoiVolume/R2.uint16.nii.gz
```

5.2.4 Output files

The root value of the key "*LIFEx.Output.Directory*" will be used for the construction of 2 files (2 files MTV_ROI and 2 files MTV_SUMMARY) will be saved at the end of the script execution by the application:

- MTV_ROI_results.csv: will contain all the features extracted from each ROI;
- MTV_ROI_legend.csv: will contain the legend of all features extracted from each ROI
- MTV_SUMMARY_results.csv: will contain aggregated features calculated from several ROIs.
- MTV_SUMMARY_legend.csv: will contain the legend of aggregated features calculated from several ROIs.