EDF R&D



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Code_Saturne documentation

Code_Saturne version 4.0.5: autovnv tool

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1 Introduction

AUTOVNV is a small framework to automate the launch of *Code_Saturne* computations and do some operations on new results.

The script needs a directory of previous *Code_Saturne* cases which are candidates to be duplicated. This directory is called **repository**. The duplication is done in a new directory which is called the **destination**.

For each duplicated case, AUTOVNV is able to compile the user files, to run the case, to compare the obtained checkpoint file with the previous one from the **repository**, and to plot curves in order to illustrate the computations.

For all these steps, AUTOVNV generate two reports, a global report which summarizes the status of each case, and a detailed report which gives the differences between the new results and the previous ones in the **repository**, and display the defined plots.

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In the **repository**, previous results of computations are required only for checkpoint files comparison purpose. They can be also usfull, if the user needs to run specific scripts.

2 Installation and prerequisites

AUTOVNV does not need a specific installation: the related files are installed with the other Python scripts of *Code_Saturne*. Nevertheless, additional prerequisites required are:

- numpy,
- matplotlib,
- python-vtk.

3 Command line options

The command line options can be found with the command: code_saturne autovnv -h.

- -f FILE, --file=FILE: gives the file of parameters for AUTOVNV. This file is mandatory, and therefore this option must be completed;
- -q, --quiet: does not print status messages to stdout;
- -u, --update: update installation pathes in scripts (i.e. SaturneGUI and runcase) only in the repository, reinitialize xml files of parameters and compile;
- -r, --run: runs all cases;
- -c, --compare: compares chekpoint files between repository and destination;
- -p, --post: postprocess results of computations;
- -m ADDRESS1 ADDRESS2 ..., --mail=ADDRESS1 ADDRESS2 ...: addresses for sending the reports.

Examples:

- code_saturne autovnv -f sample.xml: duplicates all cases from the repository in the destination, compile all user files and exits;
- code_saturne autovnv -f sample.xml -r: as above, and run all cases if defined in sample.xml;
- code_saturne autovnv -f sample.xml -r -c: as above, and compares all new checkpoint files with those from the repository if defined in sample.xml;
- code_saturne autovnv -f sample.xml -rcp: as above, and plots results if defined in sample.xml;
- code_saturne autovnv -f sample.xml -r -c -p -m "dt@moulinsart.be dd@moulinsart.be": as above, and send the two reports.
- code_saturne autovnv -f sample.xml -c -p: compares and plots results in the destination already computed.

Note:

The detailed report is generated only if the options -c, --compare or -p, --post is present in the command line.

4 File of parameters

The file of parameters is a XML formatted ascii file.

4.1 Begin and end of the file of parameters

This example shows the four mandatory first lines of the file of parameters.

```
<?xml version="1.0"?>
<autovnv>
     <repository>/home/dupond/codesaturne/MyRepository</repository>
     <destination>/home/dupond/codesaturne/MyDestination</destination>
```

The third and fourth lines correspond to the definition of the **repository** and **destination** directories. Inside the markups **repository** and **destination** the user must inform the related directories. If the **destination** does not exit, the directory is created.

The last line of the file of parameters must be:

</autovnv>

4.2 Case creation and compilation fo the user files

When AUTOVNV is launched, the file of parameters is parsed in order to known which studies and cases from the **repository** should be duplicated in the **destination**. The selection is done with the markups <study> and <case> as the following example:

The attributes are:

- label: the name of the file of the script;
- status: must be equal to on or off, activate or desactivate the markup;
- compute: must be equal to on or off, activate or desactivate the computation of the case;
- post: must be equal to on or off, activate or desactivate the post-processing of the case;
- run_id: label of the directory in which the result is stored. If this attribut is missing or set to run_id="", an automatic value will be proposed by the code.

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Only the attributes label, status, compute and post are mandatory.

If the directory specified by the attribute run_id already exists, the comptutation is not performed again. For the post-processing step, the existing results are taking into account only if no error file is detected in the directory.

With the attribute status, a single case or a complete study can be switched off. In the above example, only the case Grid1 of the study MyStudy1 is going to be created.

After the creation of the directories in the **destination**, for each case, all user files are compiled. The AUTOVNV stops if a compilation error occurs: neither computation nor comparison nor plot will be performed, even if they are switched on.

Notes:

- During the duplication, every files are copied, except mesh files, for which a symbolic link is used.
- During the duplication, if a file already exists in the **destination**, this file is not overwritten by AUTOVNV.

4.3 Run cases

The computations are activated if the option -r, --run is present in the command line.

All cases described in the file of parameters with the attribute compute="on" are taken into account.

After the computation, if no error occurs, the attribute compute is set to "off" in the copy of the file of parameters in the **destination**. It is allow to restart AUTOVNV without re-run successfull previous computations.

Note that it is allowed to run several times the same case in a given study. The case has to be repeated in the file of parameters:

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```
</case>
</study>
</autovnv>
```

If nothing is done, the case is repeated without modifications. In order to modify the setup between two runs of the same case, an external script has to be used to change the related setup (see sections 4.5 and 6).

4.4 Compare checkpoint files

The comparison is activated if the option -c, --compare is present in the command line.

In order to compare two checkpoint files, markups <compare> have to be added as a child of the considered case. In the following exemple, a checkpoint file comparison is switched on for the case Grid1 (for all variables, with the default threshold), whereas no comparison is planed for the case Grid2. The comparison is done by the external script cs_io_dump with the option --diff.

The attributes are:

- repo: id of the results directory in the **repository** for example repo="20110704-1116", if there is a single results directory in the RESU directory of the case, the id can be ommitted: repo="";
- dest: id of the results directory in the destination:
 - → if the id is not known already because the case has not yet run, just let the attribute empty dest="", the value will be updated after the run step in the destination directory (see section 5);
 - → if AUTOVNV is restarted without the run step (with the command line code_saturne autovnv -f sample.xml -c for example), the id of the results directory in the destination must be given (for example dest="20110706-1523"), but if there is a single results directory in the RESU directory of the case, the id can be ommitted: dest="", the id will be completed automatically;
- args: additional options for the script cs_io_dump
 - ⋄ --section: name of a particular variable;
 - \diamond --threshold: real value above which a difference is considered significant (default: 1e-30 for all variables);
- status: must be equal to on or off: activate or desactivate the markup.

Only the attributes repo, dest and status are mandatory.

Several comparisons with different options are permitted:

Comparisons results will be sumarized in a table in the file report_detailed.pdf (see 5):

Variable Name	Diff. Max	Diff. Mean	Threshold
VelocityX	0.102701	0.00307058	1.0e-5
VelocityY	0.364351	0.00764912	1.0e-3

4.5 Run external additional preprocessing scripts with options

The markup prepro> has to be added as a child of the condidered case.

The attributes are:

- label: the name of the file of the considered script;
- status: must be equal to on or off: activate or desactivate the markup;
- args: additional options to pass to the script.

Only the attributes label and status are mandatory.

An addionnal option "-c" (or "--case") is given by default with the path of the current case as argument (see exemple in section 6 for decoding options).

Note that all options must be processed by the script itself.

Several calls of the same script or to different scripts are permitted:

All preprocessing scripts are first searched in the MESH directory from the current study in the **repository**. If a script is not found, it is searched in the directories of te current case. The main objectif of running such external scripts is to create or modify meshes or to modify the current setup of the related case (see section 6).

4.6 Run external additional postprocessing scripts with options for a case

The launch of external scripts is activated if the option -p, --post is present in the command line.

The markup <script> has to be added as a child of the condidered case.

The attributes are:

- label: the name of the file of the considered script;
- status: must be equal to on or off: activate or desactivate the markup;
- args: the arguments to pass to the script;
- repo and dest: id of the results directory in the repository or in the destination;
 - → if the id is not known already because the case has not yet run, just let the attribute empty dest="", the value will be updated after the run step in the destination directory (see section 5);
 - → if there is a single results directory in the RESU directory (either in the repository or in the destination) of the case, the id can be ommitted: repo="" or dest="", the id will be completed automatically.

If attributes repo and dest exist, their associated value will be passed to the script as arguments, with options "-r" and "-d" respectively.

Only the attributes label and status are mandatory.

Several calls of the same script or to different scripts are permitted:

All postprocessing scripts must be in the POST directory from the current study in the **repository**. The main objectif of running external scripts is to create or modify results in order to plot them.

Example of script, which searches printed informations in the listing, note the function to process the passed command line arguments:

```
parser.add_option("-d", "--dest", dest="dest", type="string",
                     help="Directory of the result in the destination")
    (options, args) = parser.parse_args(argv)
   return options
def main(options):
   m = os.path.join(options.dest, "listing")
    f = open(m)
    lines = f.readlines()
    f.close()
   g = open(os.path.join(options.dest, "water_level.dat"), "w")
    g.write("# time, h_sim, h_th\n")
   for 1 in lines:
      if l.rfind("time, h_sim, h_th") == 0:
          d = 1.split()
          g.write("%s %s %s\n" % (d[3], d[4], d[5]))
    g.close()
if __name__ == '__main__':
    options = process_cmd_line(sys.argv[1:])
    main(options)
```

4.7 Run external additional postprocessing scripts with options for a study

The launch of external scripts is activated if the option -p, --post is present in the command line.

The purpose of this functionality is to create new data based on several runs of cases, and to plot them (see section 4.8) or to insert them in the final detailed report (see section 4.10).

The markup <postpro> has to be added as a child of the considered study.

The attributes are:

- label: the name of the file of the considered script;
- status: must be equal to on or off: activate or desactivate the markup;
- args: the additional options to pass to the script;

Only the attributes label and status are mandatory.

The options given to the script in the command line are:

• -s or --study: label of the current study;

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- -c or --cases: string which contains the list of the cases
- -d or --directories: string which contains the list of the directories of results.

Additional options can be pass to the script throught the attributes args.

Note that all options must be processed by the script itself.

Several calls of the same script or to different scripts are permitted.

4.8 Post-processing: curves

The post-processing is activated if the option -p, --post is present in the command line.

The following example shows the drawing of four curves (or plots, or 2D lines) from two files of data (which have the same name profile.dat). There are two subsets of curves (i.e. frames with axis and 2D lines), in a single figure. The figure will be saved on the disk in a pdf (default) or png format, in the POST directory of the related study in the destination. Each drawing of a single curve is defined as a markup child of a file of data inside a case. Subsets and figures are defined as markup childs of <study>.

```
<study label='Study' status='on'>
    <case label='Grid1' status='on' compute="off" post="on">
        <data file="profile.dat" dest="">
            <plot fig="1" xcol="1" ycol="2" legend="Grid level 1" fmt='r-s'/>
            <plot fig="2" xcol="1" ycol="3" legend="Grid level 1" fmt='r-s'/>
    </case>
    <case label='Grid2' status='on' compute="off" post="on">
        <data file="profile.dat" dest="">
           <plot fig="1" xcol="1" ycol="2" legend="Grid level 2" fmt='b-p'/>
            <plot fig="2" xcol="1" ycol="3" legend="Grid level 2" fmt='b-p'/>
        </data>
    </case>
    <subplot id="1" legstatus='on' legpos ='0.95 0.95' ylabel="U ($m/s$)" xlabel="Time ($s$)"/>
    <subplot id="2" legstatus='on' legpos ='0.95 0.95' ylabel="U ($m/s$)" xlabel="Time ($s$)"/>
    <figure name="velocity" idlist="1 2" figsize="(4,5)" format="png"/>
</study>
```

4.8.1 Define curves

The curves of computational data are build from data files. These data must be ordered as column and the files should be in results directory in the RESU directory (either in the **repository** or in the **destination**). Commentaries are allowed in the file, the head of every commentary line must start with character #.

In the file of parameters, curves are defined with two markups: <data> and <plot>:

- <data>: child of markup <case>, defines a file of data;
 - \rightarrow file: name of the file of data
 - \rightarrow repo or dest: id of the results directory either in the **repository** or in the **destination**;
 - ⇒ if the id is not known already because the case has not yet run, just let the attribute empty dest="", the value will be updated after the run step in the destination directory (see section 5);
 - ⇒ if there is a single results directory in the RESU directory (either in the **repository** or in the **destination**) of the case, the id can be ommitted: **repo="""** or **dest="""**, the id will be completed automatically.

The attribute file is mandatory, and either repo or dest must be present (but not the both) even if it is empty.

- <plot>: child of markup <data>, defines a single curve; the attributes are:
 - → fig: id of the subset of curves (i.e. markup <subplot>) where the current curve should be plotted;
 - \rightarrow xcol: number of the column in the file of data for the abscisse;
 - → ycol: number of the column in the file of data for the ordinate;
 - \rightarrow legend: add a label to a curve;
 - → fmt: format of the line, composed from a symbol, a color and a linestyle, for example fmt="r--" for a dashed red line;
 - \rightarrow xplus: real to add to all values of the column xcol;
 - \rightarrow yplus: real to add to all values of the column ycol;
 - \rightarrow xfois: real to multiply to all values of the column xcol;
 - → yfois: real to multiply to all values of the column ycol;
 - \rightarrow xerr: draw horizontal error bar (see section 4.8.5);
 - \rightarrow yerr: draw vertical error bar (see section 4.8.5);
 - → some standard options of 2D lines can be added, for example markevery="2" or markersize="3.5". These options are summarized in the table 2. Note that the options which are string of characters must be overquoted likes this: color="'g'."

Property	Value Type
alpha	float (0.0 transparent through 1.0 opaque)
antialiased or aa	True or False
color or c	any matplotlib color
dash_capstyle	butt; round; projecting
dash_joinstyle	miter; round; bevel
dashes	sequence of on/off ink in points ex: dashes="(5,3)"
label	any string, same as legend
linestyle or ls	-;;; :; steps;
linewidth or lw	float value in points
marker	+; ,; .; 1; 2; 3; 4;
markeredgecolor or mec	any matplotlib color
markeredgewidth or mew	float value in points
markerfacecolor or mfc	any matplotlib color
markersize or ms	float
markevery	None; integer; (startind, stride)
solid_capstyle	butt; round; projecting
solid_joinstyle	miter; round; bevel
zorder	any number

Table 2: Options authorized as attributes of the markup plot.

The attributes fig and ycol are mandatory.

Details on 2D lines properties can be found in the matplotlib documentation. For more advanced options see section 4.8.7.

4.8.2 Define subsets of curves

A subset of curves is a frame with two axis, axis labels, legend, title and drawing of curves inside. Such subset is called subplot in the nomenclature of matplotlib.

<subplot>: child of markup <study>, defines a frame with severals curves; the attributes are:

- \rightarrow id: id of the subplot, should be an integer;
- → legstatus: if "on" display the frame of the legend;
- → legpos: sequence of the relative coordinates of the center of the legend, it is possible to draw the legend outside the axis;
- → title: set title of the subplot;
- \rightarrow xlabel: set label for the x axis;
- \rightarrow ylabel: set label for the y axis;
- \rightarrow xlim: set range for the x axis;
- \rightarrow ylim: set range for the y axis.

The attributes fig and ycol are mandatory.

For more advanced options see section 4.8.7.

4.8.3 Define figures

Figure is a compound of subset of curves.

<fi>qure>: child of markup <study>, defines a pictures with a layout of frames; the attributes are:

- \rightarrow name: name of the file to be written on the disk;
- \rightarrow idlist: list of the subplot to be displayed in the figure;
- \rightarrow title: add a title on the top of the figure;
- \rightarrow nbrow: impose a number of row of the layout of the subplots;
- \rightarrow nbcol: impose a number of column of the layout of the subplots;
- → format: format of the file to be written on the disk, "pdf" (default) or "png" 1;
- \rightarrow standard options of figure can be added (table 3), for example figsize="(3,4)".

Property	Value Type
figsize	width x height in inches; defaults to (4,4)
dpi	resolution; defaults to 200

Table 3: Options authorized as attributes of the markup figure.

Details can be found in the matplotlib documentation. For more advanced options see section 4.8.7.

The attributes name and idlist are mandatory.

¹Other format could be choosen (eps, ps, svg,...), but the pdf generation with pdflatex will failed.

4.8.4 Experimental or analytical data

A particular markup is provided for curves of experimental or analytical data: <measurement>; the attributes are:

- \rightarrow file: name of the file to be read on the disk;
- → path: path of the directory where the file of data is. the path could be ommitted (path=""), and in this case, the file will be searched recursively in the directories of the considered study.

The attributes file and path are mandatory.

In order to draw curves of experimental or analytical data, the markup <measurement> should be used with the markup <plot> as illustrated below:

```
<study label='MyStudy' status='on'>
    <measurement file='exp1.dat' path=''>
            <plot fig='1' xcol='1' ycol='2' legend='U Experimental data'/>
            <plot fig='2' xcol='3' ycol='4' legend='V Experimental data'/>
    </measurement>
    <measurement file='exp2.dat' path =''>
            <plot fig='1' xcol='1' ycol='2' legend='U Experimental data'/>
            <plot fig='2' xcol='1' ycol='3' legend='V Experimental data'/>
    </measurement>
    <case label='Grid1' status='on' compute="off" post="on">
        <data file="profile.dat" dest="">
            <plot fig="1" xcol="1" ycol="2" legend="U computed" fmt='r-s'/>
            <plot fig="2" xcol="1" ycol="3" legend="V computed" fmt='b-s'/>
        </data>
    </case>
</study>
<subplot id="1" legstatus='on' ylabel="U ($m/s$)" xlabel= "$r$ ($m$)" legpos ='0.05 0.1'/>
<subplot id="2" legstatus='off' ylabel="V ($m/s$)" xlabel= "$r$ ($m$)"/>
<figure name="MyFigure" idlist="1 2" figsize="(4,4)" />
```

4.8.5 Curves with error bar

In order to draw horizontal and vertical error bar, it is possible to specify to the markup <plot> the attributes xerr and yerr respectively. The value of theses attributes could be:

• a single column number where the average uncertainty is in the file of data:

```
<measurement file='axis.dat' path =''>
     <plot fig='1' xcol='1' ycol='3' legend='Experimental data' xerr='2' />
</measurement>
```

• a sequence of two column numbers where the minimum and the maximum of the uncertainty are in the file of data:

Notes:

The attributes xerr and yerr could not be both at the same time in the markup plot. If horizontal and vertical bars has be drawn together, repeat the markup plot like this:

```
<measurement file='axis.dat' path =''>
     <plot fig='1' xcol='1' ycol='3' legend='Experimental data' xerr='2' />
     <plot fig='1' xcol='1' ycol='3' yerr='4' />
</measurement>
```

4.8.6 Monitoring points or probes

- file: name of the file to be read on the disk;
- fig: id of the subset of curves (i.e. markup <subplot>) where the current curve should be plotted;
- dest: id of the results directory in the destination:
 - → if the id is not known already because the case has not yet run, just let the attribute empty dest="", the value will be updated after the run step in the destination directory (see section 5);
 - → if AUTOVNV is restarted without the run step (with the command line code_saturne autovnv -f sample.xml -c for example), the id of the results directory in the destination must be given (for example dest="20110706-1523"), but if there is a single results directory in the RESU directory of the case, the id can be ommitted: dest="", the id will be completed automatically;

The attributes file, fig and dest are mandatory.

In order to draw curves of probes data, the markup crobes> should be used as a child of a markup <case> as illustrated below:

4.8.7 Matplotlib raw commands

The file of parameters allows to execute additional matplotlib commands (i.e Python commands), for curves (2D lines), or subplot, or figure. For every object drawn, Autovnv associate a name to this object that can be reused in standard matplotlib commands. Therefore, childs markup <plt_command> could be added to <plot>, <subplot> or <figure>.

It is possible to add commands with **Matlab style** or **Python style**. For the Matlab style, commands are called as methods of the module plt, and for Python style commands or called as methods of the instance of the graphical object.

Matlab style and Python style commands can be mixed.

• curves or 2D lines: when a curve is drawn, the associated name are line and lines (with line = lines[0]).

• subset of curves (subplot): for each subset, the associated name is ax:

4.9 Post-processing: scalar map

The post-processing is activated if the option -p, --post is present in the command line.

AUTOVNV is able to draw colored maps, with optional contours, from results on cut planes of the computational domain. The results must be a set of ensight files. The following example shows the drawing of a scalar map using the master file of result RESULTS.case. The figure will be saved on the disk in a **png** format, in the POST directory of the related study in the **destination**. Each drawing of a single scalar map is defined as a markup child of a file of data inside a case.

In the file of parameters, scalar map are defined with two markups: <resu> and <scalar>. Four other markups can be used: <scale>, <contours>, <title> and <axes> as childs of the markups <scalar>.

- <resu>: child of markup <case>, defines the master file of set of ensight files;
 - \rightarrow file: name of the master ensight file
 - \rightarrow repo or dest: id of the results directory either in the **repository** or in the **destination**;
 - ⇒ if the id is not known already because the case has not yet run, just let the attribute empty dest="", the value will be updated after the run step in the destination directory (see section 5);
 - ⇒ if there is a single results directory in the RESU directory (either in the **repository** or in the **destination**) of the case, the id can be ommitted: **repo="""** or **dest="""**, the id will be completed automatically.

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The attribute file is mandatory, and either repo or dest must be present (but not the both) even if it is empty.

- <scalar>: child of markup <resu>, defines a scalar map;
 - → name: name of the image file on the disk
 - \rightarrow variable: name of the variable to draw (must be present in the master ensight file)
 - → normal: tuple to set the normal of the cut plane of the computational domain
 - → center: tuple to set the center of the cut plane of the computational domain
 - \rightarrow stretch: tuple to stretch the view of the cut plane
 - → time-step: select the physical time of the value to display
 - \rightarrow size: size of the png image
 - \rightarrow zoom: adjust the zoom for the point of view of the cut plane
 - \rightarrow wireframe: switch on or off the mapping on the grid²

See default values for these attributes in the table 4.

Attributes	Default values
normal	(0.,0.,1.)
center	(0.,0.,0.)
stretch	(1.,1.,1.)
time-step	-1 i.e. the last record
size	(500,400)
zoom	1.0
wireframe	off

Table 4: Default attributes values of the markup scalar.

The attributes name and variable are mandatory.

- <scale>: child of markup <scalar>, defines a color bar;
 - → color: set a predefined palette in "hsv", "gray", "hot", "flag", "jet", "blue_to_yellow", "spring", "summer", "winter", "autumn"
 - → range: tuple to set the range of values to display
 - \rightarrow coord: tuple to set the normalized coordinates of the color bar
 - \rightarrow levels: number of levels for the color bar
 - → height: set the height of the color bar
 - → width: set the width of the color bar
 - → position: simple predefined position for the color bar in "North", "South", "West", or "East"
 - \rightarrow legend: legend of the color bar
 - \rightarrow fontsize: font size for the legend
 - → format: float format to display the value of the levels of the color bar

See default values for these attributes in the table 5.

No attribute is mandatory.

• <contours>: child of markup <scalar>, draw iso-contours of the selected variable on the cut plane;

²But the view might not be the true computational grid.

Attributes	Default values
color	default vtk palette
range	all values
coord	(0.9,0.09)
levels	10
height	0.8
width	0.1
position	East
legend	no legend
fontsize	20
format	%4.4f

Table 5: Default attributes values of the markup scale.

→ status: must be equal to on or off, activate or desactivate the markup

 \rightarrow nval: number of isovalues

 \rightarrow range: tuple to set the range of iso-values to display

 \rightarrow color: any matplotlib color.

See default values for these attributes in the table 6.

Attributes	Default values	
status	off	
nval	10	
range	all variable values	
color	k	

Table 6: Default attributes values of the markup contours.

No attribute is mandatory.

• <title>: child of markup <scalar>, defines a title for the scalarmap;

 \rightarrow label: text for the title

 \rightarrow fontsize: font size for the title

 \rightarrow coord: tuple to set the normalized coordinates of the title

See default values for these attributes in the table 7.

Attributes	Default values
label	no title
fontsize	20
coord	(0.5, 0.99)

Table 7: Default attributes values of the markup title.

No attribute is mandatory.

• <axes>: child of markup <scalar>, defines axes bars around the scalarmap;

→ fontsize: font size for the ticks of the axes

 \rightarrow format: float format to display the ticks of the axes

 \rightarrow levels: number of ticks of all axes

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```
\rightarrow xlabel: text of the X axe \rightarrow ylabel: text of the Y axe \rightarrow zlabel: text of the Z axe
```

See default values for these attributes in the table 8.

Attributes	Default values
fontsize	20
format	%6.3g
levels	3
xlabel	no text
ylabel	no text
zlabel	no text

Table 8: Default attributes values of the markup axes.

No attribute is mandatory.

4.9.1 VTK raw commands

The file of parameters allows to execute additional vtk commands (i.e Python commands). This possibility is mainly useful for interaction with actors and the camera. For each actor drawn, Autovnv associate a name to this object that can be reused in standard vtk commands. Therefore, childs markup <vtk_command> could be added to <scalar>.

The defined actors and camera are:

 \rightarrow grid: the scalar map,

</resu>

4.10 Post-processing: input files

The post-processing is activated if the option -p, --post is present in the command line.

AUTOVNV is able to include files into the final detailed report. These files must be in the directory of results either in the **destination** or in the **repository**. The following example shows the inclusion of three files: performance.log and setup.log from the **destination**, and a performance.log from the **repository**:

In the file of parameters, input files are defined with markups <input> as childs of a single markup <case>. The attributes of <input> are:

- \rightarrow file: name of the file to be included
- \rightarrow repo or dest: id of the results directory either in the **repository** or in the **destination**;
 - ⇒ if the id is not known already because the case has not yet run, just let the attribute empty dest="", the value will be updated after the run step in the destination directory (see section 5);
 - ⇒ if there is a single results directory in the RESU directory (either in the **repository** or in the **destination**) of the case, the id can be ommitted: **repo="""** or **dest="""**, the id will be completed automatically.

The attribute file is mandatory, and either repo or dest must be present (but not the both) even if it is empty.

5 Output and restart

Autovnv produces several files in the **destination** directory:

- report.txt: standard output of the script;
- auto_vnv.log: log of the code and the pdflatex compilation;
- report_global.pdf: summary of the compilation, run, comparison, and plot steps;
- report_detailed.pdf: details the comparison and display the plot;
- sample.xml: udpated file of parameters, useful for restart the script if an error occurs.

After the computation of a case, if no error occurs, the attribute compute is set to "off" in the copy of the file of parameters in the **destination**. It is allow a restart of AUTOVNV without re-run successfull previous computations. In the same manner, all empty attributes repo="" and dest="" are completed in the udpated file of parameters.

6 Tricks

• How to comment markups in the file of parameter?

The opening and closing signs for commantaries are <!-- and -->. In the following example, nothing from the study MyStudy2 will be read:

• How to add text in a figure?

It is possible to use raw commands:

• Adjust margins for layout of subplots in a figure.

You have to use the raw command subplots_adjust:

• How to find a syntax error in the XML file?

When there is a misprint in the file of parameters, AUTOVNV indicates the location of the error with the line and the column of the file:

```
my_case.xml file reading error.
This file is not in accordance with XML specifications.
The parsing syntax error is:
my_case.xml:86:12: not well-formed (invalid token)
```

• How to set a logarithmic scale?

The following raw commands have to be used:

• How to create a mesh automatically with SALOME?

The following example shows how to create a mesh with a SALOME command file:

with the script salome.sh (depending of the local installation of SALOME):

#!/bin/bash export ROOT_SALOME=/home/salome/salome-640/Salome-V6_4_0-c7-v2 source /home/salome-640/Salome-V6_4_0-c7-v2/salome_prerequisites_V6_4_0_appli.sh source /home/salome-640/Salome-V6_4_0-c7-v2/salome_modules_V6_4_0.sh /home/salome/salome-640/appli_V6_4_0/bin/salome/runSalome \$* and the script of SALOME commands my_mesh.py: #!/usr/bin/env python # -*- coding: utf-8 -*import geompy import smesh # create a box box = geompy.MakeBox(0., 0., 0., 100., 200., 300.) idbox = geompy.addToStudy(box, "box") # create a mesh tetra = smesh.Mesh(box, "MeshBox") algo1D = tetra.Segment() algo1D.NumberOfSegments(7) algo2D = tetra.Triangle() algo2D.MaxElementArea(800.) algo3D = tetra.Tetrahedron(smesh.NETGEN) algo3D.MaxElementVolume(900.) # compute the mesh tetra.Compute()

• How to carry out a grid convergence study?

The following exemple shows how to carry out a grid convergence study by running the same case three times and changing the parameters between each run with the help of a prepro script.

Here the mesh, the maximum number of iterations, the reference time step and the number of processes are modified, before each run, by the script prepro.py.

The file of parameters is as follows:

export the mesh in a MED file tetra.ExportMED("./my_mesh.med")

Recall that the case attribute run_id should be given a different value for each run, while the label should stay the same and that the prepro script should be copied in the directory MESH of the study or in the directory DATA of the case.

The prepro script is given below. Note that it can be called inside the file of parameters without specifying a value for each option:

```
#!/usr/bin/env python
# -*- coding: utf-8 -*-
# Standard modules import
import os, sys
import string
from optparse import OptionParser
 _____
#-----
# Application modules import
from Pages.ScriptRunningModel import ScriptRunningModel
def process_cmd_line(argv):
   """Processes the passed command line arguments."""
   parser = OptionParser(usage="usage: %prog [options]")
   parser.add_option("-c", "--case", dest="case", type="string",
                   help="Directory of the current case")
   parser.add_option("-p", "--param", dest="param", type="string",
                   help="Name of the file of parameters")
   parser.add_option("-m", "--mesh", dest="mesh", type="string",
                   help="Name of the new mesh")
   parser.add_option("-n", "--iter-num", dest="iterationsNumber", type="int",
                   help="New iteration number")
   parser.add_option("-u", "--n-procs", dest="n_procs", type="int",
                   help="Number of processes (units)")
   parser.add_option("-t", "--time-step", dest="timeStep", type="float",
                   help="New time step")
```

if __name__ == '__main__':

```
parser.add_option("-a", "--perio-angle", dest="rotationAngle", type="float",
                      help="Periodicity angle")
    (options, args) = parser.parse_args(argv)
   return options
def main(options):
   from cs_package import package
   from Base.XMLengine import Case
   from Base.XMLinitialize import XMLinit
   from Pages.SolutionDomainModel import SolutionDomainModel
   from Pages.TimeStepModel import TimeStepModel
   from Pages.SteadyManagementModel import SteadyManagementModel
   fp = os.path.join(options.case, "DATA", options.param)
   if os.path.isfile(fp):
        try:
            case = Case(package = package(), file_name = fp)
            print("Parameters file reading error.\n")
            print("This file is not in accordance with XML specifications.")
            sys.exit(1)
        case['xmlfile'] = fp
        case.xmlCleanAllBlank(case.xmlRootNode())
        XMLinit(case).initialize()
        if options.mesh:
            s = SolutionDomainModel(case)
            1 = s.getMeshList()
            s.delMesh(1[0])
            s.addMesh((options.mesh, None))
        if options.rotationAngle:
            s.setRotationAngle(0, options.rotationAngle)
        if (options.iterationsNumber):
            s = SteadyManagementModel(case)
            t = TimeStepModel(case)
            if s.getSteadyFlowManagement() == 'on':
                s.setNbIter(options.iterationsNumber)
            else:
                t.setIterationsNumber(options.iterationsNumber)
        if (options.TimeStep):
            t = TimeStepModel(case)
            t.setTimeStep(options.TimeStep)
        if (options.n_procs):
            mdl = ScriptRunningModel(case)
            mdl.setString('n_procs', str(options.n_procs))
        case.xmlSaveDocument()
```

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options main(opt	<pre>= process_cmd_line(sys.argv[1:]) tions)</pre>	
#		