# Material & Structure Analysis Suite



User commands Version 8.5

Z-set is distributed by

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# Chapter 1

# Introduction

# Z-set user commands

#### **Description**:

This Z-set user-commands handbook covers the command syntax and some of the details for the different files involved with basic use of the FEA related functionality.

#### Handbook Summary:

The following list summarizes the documentation for all of Z-set. As part of our 8.2/8.3 developments, greatly expanding the software documentation is one of our primary goals.

The list below is sorted in what we feel would be an appropriate sequence for the normal user, starting with installation and reviewing capabilities, to creating input files and eventually scripting and developing add-ons to the software.

- **Release Notes/Zmaster** The basic overview of the software, installation instructions, and documentation for the graphical user interface Zmaster on all platforms. The Zmaster manual also now covers all the base reference chapters such as environment variables, user parameters, function reference, command line programs, etc.
- **Examples/Training** This book is essentially the "getting started" documentation for the software. The book describes Z-mat, simulation, optimization, material models, and the FEA code use. The examples cover setup of complete models, and are meant to demonstrate the capabilities with relatively simple examples.
- **Z-mat User commands** This summarizes the command file formats and capabilities of the Z-mat interface, simulation, optimization and material files for all of Z-mat and Z-set.
- **Z-set User commands** This summarizes the command file formats for the FEA related capabilities including meshing, FEA solution, post processing, etc.
- **Developer** A guide to the user-extensible features of the software, including scripting, ZebFront material model development, making plugins, and the C++ programming API.
- Plugins A guide to add-on features available.
- **Theory** Theory manual covering formulations (under development).

# Conventions

This page summarizes the conventions used for the Z-set input files. An overview of the general command syntax (command hierarchies) is given in the beginning of the *Examples/Training* manual.

- Running of Z-set modules generally requires that a "problem name" be given. Most input and output data files are based on this name with a variety of suffixes attached. Henceforth, *problem* will often be used to indicate the problem name given while running the commands.
- The characters % and # indicate that the rest of the line is a comment. For example: \*\*\*load % external problem loading
- There are no abbreviations allowed in the use of keywords. All keywords and command names must be written entirely.
- The admissible characters for the names of user variables are: a-z, A-Z, ", +, -, \*, ., =, /, \_, ), (, \, ~
- The text entry is *always* case-sensitive.
- The use of braces [ ] in the syntax descriptions indicates an option with a default definition.
- All parameter values used in the input files and described as "real" in the syntax descriptions must have a decimal point. All standard specifications of floating point values are accepted. Two examples of real values are: 3.0 4.2e-5
- The use of parenthesis () indicates data input of real values in vector form. An example is: (0.1 0.2 1.0). In most cases the size of such vectors must be compatible with the overall problem dimension.

The symbol  $\stackrel{\textcircled{}}{\widehat{}}$  indicates a section where the calculation is sensitive to input data or format.

# Chapter 2

# Mesher

# Introduction

#### **Description**:

The Mesher module is provided in order to perform batch mesh manipulations and combinations to pre-existing meshes, and import and export Z-set format files<sup>1</sup>. These operations supersede the meshing utility programs previously supplied with Zebulon. The normal mode of operation is as a batch process launched with the command Zrun -m prob. The utilities are however also supported in some other applications, such as the batch mesher commands of the Zmaster geom and mesh menus.

An important thing to remember is that the ASCII file represents the objects and configuration options for a particular calculation. The computational modules are then run sequentially in order to modify or generate the mesh. This is therefore the configuration of an object system, and not the activation of repeated "database" manipulation commands.

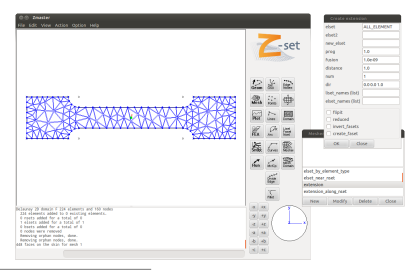
#### Batch mode:

For batch operation, an ASCII input file is submitted to the Zrun program to manipulate and combine any number of meshes. The operation will generate more files depending on the options chosen. The most common use is to generate a new ASCII .geof or binary .geo file for use in a FEA calculation.

See the syntax description for **\*\*\*\*mesher** below 2.4 for a detailed overview of the mesher syntax, with some examples.

#### Running in Zmaster:

The batch mesh operations in Zmaster operate on the current mesh (accessed under the Mesh command, or act on a newly generated mesh after all the Zmaster domains are generated (in the Geom command). This later operation happens after regenerating the mesh with the Mesh Domain button. The figure below shows the dialogs which get opened after selecting the Batch Mesher Create button in the geometry toolbar.



<sup>1</sup>example problems using the mesher can be found in \$Z7PATH/test/Mesher\_test/INP and in \$Z7PATH/test/Zmaster/MESHER

The operation of the graphical interface for running batch meshers is described more completely in the Zmaster manual. Here we want to remark on the connection between the command file syntax described in this chapter and the dialog creation.

The available meshers are listed in the bottom list box of the Mesher Create dialog with the exact names of the **\*\***-level command option in this chapter. The **\*\*\*** level commands are not available in Zmaster, so the operation always applies to the currently loaded mesh. In the figure above, the mesher option **\*\*extension** (see page 2.53) is selected. Selecting either New or Modify will open up an edit dialog with the options equivalent to the **\***-level options of the selected mesher. The option data fields can then be typed directly into the dialog box entries.

#### <u>User extensions</u>:

Because there are in fact many many things one may wish to do to a mesh in the course of building a model, the interface for adding new mesh transformation commands is especially flexible and easy to program. The details of how to create a mesher are described in the separate Z-set Developer handbook, with example code in the distribution \$Z7PATH/User-project directory.

# \*\*\*\*mesher

#### **Description**:

The basic syntax for meshing operations follows the same logic as other parts of the program. Meshing blocks are started with the four asterisk command **\*\*\*\*mesher**. The mesher can load different portions of the final mesh as independent meshes. The successful mesher will therefore learn to create a series of meshing operations, modifications, and unions for a final mesh.

#### Note:

There is a modified syntax for the mesher which can be used by Zmaster in the .mast file. The \*\*\*\*master command accepts the sub-command \*\*\*mesher which indicates that transformation commands from this chapter are to be applied every time the mesh button is pressed. One can move those commands outside the Zmaster program by changing the command to \*\*\*\*mesher and including a \*\*\*mesh/\*\*open group.

#### Syntax:

The major meshing commands are as follows:

```
****mesher
[ ***delete_file fname ]
[ ***global_parameter param-definitions ]
[ ***shell shell-command ]
[ ***function_declarations functions ]
[ ***mesh name ]
....
```

- **\*\*\*delete\_file** This allows entry of file names to delete before running the meshers. Usually this will be the new mesh file name to ensure a clean state before running.
- **\*\*\*function\_declarations** This command allows entry of function declarations ahead of the mesher applications, so that complex pre-defined functions can be used where function selection is allowed (e.g. nsets and so on).
- **\*\*\*global\_parameter** This base-level command allows setting of global parameter definitions in the mesher input to customize behavior. See the reference section in the Release/Zmaster manual.
- **\*\*\*mesh** This command opens a new mesh object which will be the subject of meshing operations. Use the **\*\*open** command to load an existing mesh in for operations. Any number of **\*\***-level transformation commands can then be applied to the mesh before the next **\*\*\***-level command. At the occurrence of the next **\*\*\***-level command, the mesh will be saved to disk with the problem name given as a parameter to **\*\*\*mesh**. Following **\*\*\*mesh/\*\*open** operations can re-open the mesh for further processing.
- **\*\*\*shell** This command can be used to launch external programs to generate sub-meshes or other operations. Probably the most useful purpose for this command is to run a Zmaster batch job.

#### Example:

An example of the mesher commands as output by Zmaster is as follows:

```
% in prob.mast
***mesher
 **extension
 *elset face
 *distance 3.000000e+00
 *num 1
****return
```

These lines will make the element set face (which was created using one of the domain tools) to be extended into a 3d mesh every time the mesh-domains button is pressed. The mesher commands could be moved into the .inp file to separate the process:

```
%
% in prob.inp.. importing prob_2d.geof a,d applying an
% extension into 3d.
%
****mesher
 ***mesher
 ***mesh prob
 **open prob_2d.geof
 **extension
 *elset face
 *distance 3.000000e+00
 *num 1
****return
```

#### Example:

One feature of the batch mesher is the ability to step through multiple meshing steps in the same file. For example a multi-part mesh can be generated using a combination of Zmaster batch jobs, and mesher transformers and unions:

```
****mesher
***shell Zrun -B part1.mast
                               % generate a new, 2d part1.geof
***mesh part1
                               % start creating the 3d, also part1.geof
  **open part1.geof
                               \% open the 2d created from Zmaster batch
                               % now start extensions
  **extension
    *elset ALL_ELEMENT
    *new_elset solid
    *prog 1
    *distance 0.5
    *num 2
    *dir (0 0 1 )
  **nset z=0
    *plane 0. 0. 1. 0.
    *limit 0.001
***shell Zrun -B part2.mast
                               % a ***-level command closes the prev. mesh
***mesh part2
                               % start the part2 mesh independently
  **open part2.geof
                               % and in the same way as above. This lets
                               % us generate complex meshes in 1 run
  %
  %
     etc
   %
****return
```

#### \*\*\*mesh

#### **Description**:

As introduced before, this command indicates the start of a new mesh object. The meshing operations which follow will act on the mesh object, until the next **\*\*\***-level command is reached. At that point the mesh will be saved with the given name. Use of this command invariably involves use of the **\*\*open** or **\*\*import** commands.

# Syntax:

The basic **\*\*\*mesh** section syntax is summarized below:

\*\*\*mesh [output-name]
\*\*export format output-name
\*\*open input-name
\*\*open\_mast mast-file.mast
\*\*import format input-name
\*\*inp\_file
\*\*output output-name
\*\*dont\_save\_final\_mesh
\*\*transform
...

The mesh output name *output-name* is the problem name, and therefore does not have the .geof suffix. It is optionally included on the same line as **\*\*\*mesh** for convenience.

**\*\*export** Write an input file in a format other than native Z-set.

- **\*\*import** Open a mesh file in a format other than the native Z-set .geof format. The format type is a keyword for one of the external mesh formats allowed.
- **\*\*inp\_file** specify that the input file (FEA loads, BCs element formulation, time sequences, etc) are to be read and output with the mesh. This is probably desired in order to get as *full* an import or export as possible. Note: this command is provided as a convenience feature, and does not imply full compatibility between formats. It is likely the different codes options just do not have an equivalent for translation, and will be passed silently. Be careful as well not to overwrite existing hand-edited input files! there are no questions before output is written.
- **\*\*open** Open a file defining the mesh which will be operated on. There must be only one instance of open in the **\*\*\*mesh** block. The file name is the problem name without the **.geof** suffix. Additional mesh components can be loaded using the **\*\*union** command described on page 2.119.
- **\*\*open\_mast** Open a Zmaster .mast file, running Zmaster batch on it beforehand. This open mode actually does a union, but normally it would be used in place of an **\*\*open** or **\*\*import** command.
- **\*\*output** specify the output file.
- **\*\*dont\_save\_final\_mesh** disables writing the mesh, as it normally is the final stage of the meshing process.

 $\dots$  continued

Aside from the basic control options, any number of mesh transformation operations can be added in this section as indicated by **\*\****transform*. The syntax and operation of these different mesher transformations are the subject of the remainder of this chapter.

#### **<u>Translations</u>**:

Version 8.4 is shipped with the following import/export translators:

CODE	DESCRIPTION
abaqus	Import/export an ABAQUS mesh and input file
ansys	Exports mesh and input file commands to ANSYS input file.
gfm	Import/export the GFM format from COSMOS/M with input file commands
gmsh	Import/export GMSH format. Only GMSH format's version 2 is supported for export.
ideas	Import I-deas unv format.
k	Imports/exports to LS-Dyna input file.
neu	Import FEMAP neutral format.

These import functions currently only import geometrical information. Boundary conditions and real constants are mapped into mesh sets (nset, elset, etc) and given names based on their values.

\*\*\*\*mesher \*\*\*mesh

# Example:

A small example input follows. See the following sections for more examples demonstrating the individual operations. This example opens up a sub-mesh named BASE1, rotates and flips it, and creates some node/boundary sets for it.

```
****mesher
***mesh BASE
**open BASE1
**rotate
    x1 1. 0. 0.
    x3 0. 1. 0.
**switch *axis -z z
**rotate x3 0. 0. 1. x1 0.984808 -0.173648 0.0
**translate 0. -5.5 -1.0
**nset base_fix *plane 0. 0. 1. 0.0
**bset base_top *plane 0. 0. 1. 1.0
****return
```

# Example:

Two examples follow, the first of mesh import and the second of problem export (mesh and loads).

```
****mesher
```

```
***mesh linear_trial
                                  % start making file linear_trial.geof
                                  % import a Cosmos/M model
  **import gfm big_mesh.gfm
  **quad_to_lin
                                  % linearize the mesh
  **elset front *func x>0.0;
                                  % generate new sets
  **elset bottom *func y<0.0;</pre>
  **renumbering frontal_only
                                  % frontal renumbering
****return
****mesher
***mesh
                                  % no name because we're giving output
  **export abaqus abaqus_lin.inp
                                  % via the export command.. abaqus format
                                  % include info from the .inp file (BCs)
  **inp_file
  **open zebu_lin.geof
                                  % This specifies the input name for both
                                  % the geof and .inp files
***return
```

# $**add_element$

#### **Description**:

This command is used to enter elements into a mesh "by hand." It is useful normally to utilize element numbers which are "far away" from those generated by other meshing commands. Normally Zmaster and batch mesher commands all presume a sequentially numbered mesh, starting with one. So if you know there will be less than 10000 elements, you could start your hand-created element numbers there.

It is normally very useful to write shell scripts (or command batch files) to generate the records for this type of command.

#### Syntax:

The **\*\*add\_element** command takes a list of element definitions, much as they are given in the **.geof** file. There are no checks if an equivalent element already exists. The element id must be unique, and the nodes with the given ids must exist before this command runs.

```
**add_element
ele-id ele-type n_1 n_2 \dots n_N r_1 \dots r_m
```

Any number of elements can be added. The entries must however start with an acceptable element number and element type. In the above,  $n_1$  means node number 1 *id*, and  $r_1$  means real constant 1 for the element (e.g. thickness). The number and type of real constant possible depends on the element type, as does the appropriate number of nodes.

A duplication of this command exists which automatically creates an element set made up of the new elements. That command has the syntax:

```
**add_element_elset eset-name
ele-id ele-type n_1 n_2 \dots n_N r_1 \dots r_m
```

# Example:

Some example uses follow.

. . .

```
**add_element
6000 c2d4 1 2 3 4
6001 c2d4 2 5 6 7
**elset new_elem
*elements 6000 6001
```

A shorter version would use the **\*\*add\_element\_elset** variant:

```
**add_element_elset new_elem
6000 c2d4 1 2 3 4
6001 c2d4 2 5 6 7
```

# \*\*add\_info

# **Description**:

This command is used to assign "real-constant" information to a set of elements if that has not yet been set in other preprocessing. One command per elset/real constant value must be used. The real constant will also be added on the the end of the real constants for the elements as they exist in the given elset.

Typical applications of this command are to assign thicknesses to plane stress elements.

# Syntax:

The **\*\*add\_info** command takes the following syntax:

```
**add_info
*elset eset-name
*info real-value
```

# Example:

An example use follows:

# \*\*add\_node

#### **Description**:

This command is used to enter nodes into a mesh "by hand," and parallels the **\*\*add\_element** command. It is useful normally to utilize node numbers which are "far away" from those generated by other meshing commands.

# Syntax:

The **\*\*add\_node** command takes a list of node definitions as they appear in the .geof file. The node dimension will be sized according to the number of coordinates entered in after the node id. Note that zero coordinates must therefore be entered.

There is checking only if the node id already exists. No check is performed for the node coordinate.

```
**add_node

id-num x<sub>1</sub> x<sub>2</sub> [ x<sub>3</sub>]

...
```

A duplication of this command exists which automatically creates a node set made up of the new nodes. That command has the syntax:

```
**add_node_nset nset-name
id-num x<sub>1</sub> x<sub>2</sub> [ x<sub>3</sub>]
...
```

Any number of nodes can be added. The entry fields are read until another **\*\*-level** command is reached. The dimension of the node is assigned according to the number of entered coordinates  $(x_1, \text{ etc above})$ .

#### Example:

An example use follows.

\*\*add\_node\_nset some\_nodes
200 1. 0. 0.
201 2. 0. 0.
202 2. 1. 0.
203 1. 1. 0.

# \*\*boolean\_operation

#### **Description**:

This command is used to perform boolean operation between meshes (union, intersection or difference). It uses the GNU Triangular Surface library (GTS) plugin for operations on mesh skins. For volume meshes, INRIA remeshing plugins are required (see **\*\*yams\_ghs3d**).

#### Syntax:

The command has the following syntax:

```
**boolean_operation
 *operation
              operation-type
*file1
          geof-filename
*file2
          geof-filename
*output_file gts-output-filename
                    ٦
 [ *keep_1st
               nset
 [ *keep_2nd
               nset
                     ٦
 [ *dist_crit distance
                         ٦
 [ *surface_crit value ]
 [ *refinement func(x,y,z) ]
 [ *absolu ]
 [ *tolerance tol
                   ٦
 [ *min_size min
                   1
 [ *max_size max ]
 [ *gradation grad ]
 [ *optim_style opt ]
 [ *options yams-command-line-options
```

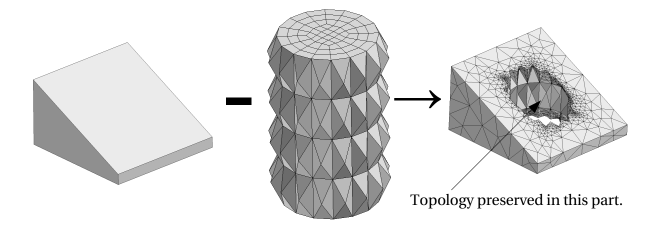
Many keywords correspond to the yams syntax and one should refer to the manual. The operation order is file1 OPERATION file2. For a complete description of the syntax see **\*\*yams\_ghs3d** documentation.

٦

- \*operation operation-type must be intersection, union or difference.
- **\*output\_file** file name containing GTS output after the boolean operation.
- \*keep\_1st this will preserve the topology of specified *nset* the in first mesh.
- \*keep\_2nd this will preserve the topology of specified *nset* the in second mesh.
- \*dist\_crit merge vertices closer than the specified distance.
- **\*surface\_crit** eliminate degenerate triangles with relative surface areas smaller than the specified *value*.

\*\*\*mesher
\*\*\*mesh
\*\*boolean\_operation

```
****mesher
***mesh difference
**boolean_operation_ghs3d
*operation difference
*file1 cube.geof
*file2 volume.geof
*output_file difference.gts
*options -m 100 -FEM
*optim_style 1
*keep_2nd skin
****return
```



#### <u>Note</u>:

The "difference" operation requires the substracted mesh to be slightly bigger than the other one. Therefore, it may be necessary to "thicken" it slightly, using e.g. **\*\*thicken\_bset** (page 2.112) or **\*\*porcupine** (page 2.86).

# \*\*bounding\_box

# **Description**:

This mesher is used as a utility to print the bounding box which the mesh fits in

# Syntax:

There are no options with this command. The bounding box will be calculated using the current mesh configuration. That is after all preceeding meshers are applied. Any number of calls to this can of course be applied.

\*\*bounding\_box

# Example:

An example output from this command follows.

Bounding box :

xmin=0 xmax=3 ymin=0 ymax=3 zmin=0 zmax=1

Note this is printed among the general screen messages, and stored no where else. The responses round to integer values if there is no decimal needed.

#### \*\*bset

#### **Description**:

This command creates a boundary set according to various input data. The syntax is the same as for the **\*\*nset** command, except an ordered boundary is produced. A typical use is to generate first an **nset** using the **\*\*nset** command, and then simple make a copy of it in the bset format using the **\*use\_nset** option. The command can also be used to find the outer boundary of an elset.

#### Syntax:

The bset mesher takes the following command syntax:

```
**bset name ... options
```

with the following options available:

- \*axes switches the axes defining a cylindrical coordinate system when using the \*plane
  selection. an example is \*axes 1 3 2 where the cylinder is rotated about the 2 axis
  instead of the default 3-rd axis.
- **\*elset** *elset1 ... elsetN* use only the nodes which are used by the given element sets. This can be used to make a "wrap" of separate elsets, or to easily limit the selection criterion.
- **\*function** Create a bset using the nodes which satisfy the given function. Remember to include a semicolon at the end of the function definition.
- \*limit Limit for nodes to qualify to be accepted by a function or plane function (default
   is 1.e-3).
- \*nodes node1...nodeN uses the given nodes to make up the set.
- \*plane makes a boundary given a plane equation (4 real values). The 1st 3 values are the components of the plane normal, and the fourth is the intercept (or equiv. for cyl. coordinates).
- \*req\_number allows the user to set the number of nodes of the \*use\_nset option which must match the element faces in order to create the bsets. For example if the nset given includes just the edge nodes of an element around a corner, the "wrapping" face around the corner can be added.
- **\*surface** Indicates that the set should be a bounding surface (outer) of the acceptable nodes (see also unshared\_faces page 2.121).
- \*type cartesian | cylindrical set the type of coordinate system to be used with the \*plane option.
- \*use\_dimension dim sets the dimension of bset to create. Normally the bset will be a
  faset for 3D meshes (the max space dimension found in the mesh), and will be a liset
  for 2D meshes.

 $\dots$  continued

\*\*\*\*mesher \*\*\*mesh \*\*bset

- \*use\_bset bset-name use the nodes in the given bset to apply the criteria given. Several entries may be given.
- \*use\_nset nset-name make the nset named into a bset. Several entries may be given. This replaces the use of all nodes in the mesh to apply the selection criteria.

#### Example:

To create the outer "wrapping" surface (unshared) for the element set PIPE2:

```
**bset wrap1 *surface *elset PIPE2
```

Making an outer face at a radius of 20.

```
**bset face_r=20
*limit 1.e-3
*type cylindrical
*plane 1. 0. 0. 20.
```

More examples are given for the \*nset command on page 2.80.

# \*\*bset\_align

#### **Description**:

This command aligns lisets (in 2D) and fasets (in 3D) such that their "normals" are aligned in the same direction sign-wise. The normals are still normal to the face.

The best normal alignment is very important for certain boundary conditions (e.g. pressure) and contact surfaces to be correctly defined.

# Syntax:

\*\*bset\_align
 \*bsets bset1 ... bsetN
[ \*normal (direction) ]
[ \*towards (point) ]
[ \*away\_from (point) ]
[ \*inwards ]

By default (i.e. if neither normal, towards nor away\_from are specified), the bset are oriented outwards (or inwards, if it is specified).

- **\*bsets** is the list of bsets on which the reorientation is applied. The shorthand ALL may be used to select all bsets.
- **\*normal** is a global direction that all normals will be aligned with (positive scalar product)

\*towards all normals will be directed towards this point

\*away\_from all normals will be directed away from this point

\*inwards all normals will be directed inwards (instead of the default ourtwards)

These last 4 options are of course mutually exclusive. Note that the vectors must be entered using vector notation (c.f. page 1.3).

#### <u>Note</u>:

This mesher replaces **\*\*faset\_align** (page 2.59), which was only available in 3D.

```
**bset_align
*bsets top bottom
*normal (0. -1. 0.) % all normals will point down
**bset_align
*bsets exterior % all normals will point outwards
**bset_align
*bsets left right
*towards (0. 0. 0.) % all normals will point to the origin
```

# \*\*bset\_to\_mast

#### **Description**:

This command generates a .mast file fragment from an existing liset. This fragment can be added to an existing mast file, to generate a new geometry.

#### Syntax:

The command has the following syntax:

\*\*bset\_to\_mast
\*liset liset-name
\*output mast-file

\*liset is the name of the liset to consider

**\*output** specifies the name of the resulting mast file

```
****mesher
***mesh
**dont_save_final_mesh
**open section.geof
**unshared_edges border
**bset_to_mast
*liset border
*output starter.mast
****return
```

# $**bset\_to\_mesh$

#### **Description**:

This command creates a 2D mesh from an existing boundary set (bset). Currently, it only handles best on the x - y plane, and linear elements.

In a certain way, it does the opposite of an **\*\*extension**.

# Syntax:

The command has the following syntax:

**bset_to_mesh	
*bset_name	bset-name
<pre>*new_mesh_name</pre>	resulting- $geof$ - $name$
[ *keep_bset ]	liset-name(s)

**\*bset\_name** is the name of the bset to consider.

**\*new\_mesh\_name** is the name of the resulting geof file.

\*keep\_bset is a list of existing lisets to keep in the 2D mesh.

```
***mesher
***mesh tmp.geof
**open tube3D.geof
**bset_to_mesh
*bset_name face.1
*new_mesh_name tube_section.geof
****return
```

# \*\*build\_fronts

#### **Description**:

This command fastly creates many continuous lisets related to a given nset.

Resulting output as many as required lisets named liset-name0, liset-name1, etc.

# Syntax:

\*\*build\_fronts
[ \*elset elset-name ]
[ \*nset nset-name ]
[ \*bset liset-name ]

**\*elset** restrict operation to a given elset (default is **ALL\_ELEMENT**).

**\*nset** nodes that must be used to creat lisets.

\*bset base name of created lisets.

# \*\*build\_parallel\_boundary\_files

#### **Description**:

This mesh operation is used to make data files for use with boundary conditions allowing boundary set binary files to be used to apply variable conditions in a parallel computation. Normally such files are supplied at as constant values for element faces, and the binary files must be split in order to represent faces existing on parallel computation sub-domains.

Please see the **\*\*\*pressure** command on page 3.68 for further details for the file loading.

# Syntax:

The command has the following syntax:

**build	d_parallel_boundary_files
*file	file-name
*bset	bset-name

- \*file specifies the input file (*problem*) name. The output file names will look like: *file-name*.005 for the output map 5.
- \*bset the name of the boundary set where these BC values apply. This set **must** be ordered exactly the same as the stored file data.

# Example:

An example use follows. The first snippit is the mesher applied to a set of binary files in order to split them up correctly for a parallel computation:

```
****mesher

***mesh press_calc

**open meshfile.geof

*build_parallel_param_files

*file thermal_calc.ctnod

*card 200 % total number to process

**build_parallel_boundary_files

*file internal_pressure.bin external_pressure.bin shell_pressure.bin
*bset internal_pressure external_pressure shell_pressure
****return
```

# \*\*build\_parallel\_param\_files

# **Description**:

This mesh operation is used to make data files for use with the **\*\*\*parameter** command in a parallel finite element calculation (see page 3.155). It takes a parameter binary file which would normally be used directly with the **\*\*\*parameter \*\*file** command described on page 3.157 and splits it appropriately for the different sub-domains.

# Syntax:

The command has the following syntax:

\*\*build\_parallel\_param\_files
\*file list-of-files
\*card nb-outputs-for-each-file
[ \*ip ]

- \*file specifies either one or a list of binary input file names for reading the parameter values. These files will be read using the Ni format which defaults to big-endian. The output file name will be formatted for example *file-name*.005 (for the output map 5).
- \*card the number of output maps for each file specified in the \*file parameters. This command assumes that the file records 1-1 the current mesh node numbering and size.
- \*ip indicates that the parameter values are at integration points, not nodes. All the element integration locations are assumed to map exactly the same as the calculation generating the parameter file.

# \*\*phi\_psi\_no\_refine

#### **Description**:

This command is used to compute the two levelsets describing a planar semi-circular crack. The first levelset  $\Phi$  implicitly describes the crack plane as the surface where  $\Phi = 0$  whereas the second levelset  $\Psi$  implicitly describes a cylinder as the surface where  $\Psi = 0$  whose intersection with the plane forms the crack.

So the crack location is defined by:

 $\begin{cases} \Phi = 0\\ \Psi < 0 \end{cases}$ 

This mesher command computes the nodal values of these two levelsets, and stores the result in two files named phi.dat and psi.dat. These two files can then be used in an XFEM analysis.

# Syntax:

The command has the following syntax:

```
**phi_psi_no_refine
 *center (1. 0.5 0.5)
 *normal (0. .2 .7)
 *rho .3
 *vtk_output levelset
```

 $\texttt{*center} \quad \mathrm{defines \ the \ crack \ center}$ 

\*normal is the plane normal vector

**\*rho** is the crack radius

**\*vtk\_output** if present, allows to export the computed levelsets into two vtk legacy files for debugging purposes using Paraview

# $**cfv_build$

#### **Description**:

This command is used to mesh a regular shape around a crack front (the command name stands for "Crack Front Volume build"). This volume can be used to perform accurate integrals for energy release rate computations with De Lorenzi or G-theta formulations. In order to be connected to a tetrahedral mesh, "c3d5" pyramid elements are located on every face which must be connected to the rest of the mesh.

The crack front be can specified by a liset (a Catmull-Rom spline interpolation is used), or volume must be completly discribed by a crack front method component (i.e. gtheta). Warning: this method is only available for 3D meshes, and it generates linear meshes only.

# Syntax:

The command has the following syntax:

**cfv_build	
[ *liset liset-name propagation-direction-vector	]
[ *ask_crack_to crack-method-name ]	
<b>*rc</b> internal-square-length	
*ri internal-radius	
<b>*re</b> external-radius	
<b>*nbc</b> number-of-square	
*nbri number-internal-cuts	
<b>*nbre</b> number-external-cuts	
[ *delta square-deformation ]	
[ *delta_p pyramid-height ]	
[ *connect connection-radius ]	
[ *opening opening-angle ]	

Note that at least on of the command **\*liset** or **\*ask\_crack\_to** must be specified in order to select how to build the volume.

\*liset specifies whether one liset describes the crack front. A vector is required to define the orientation of the crack.

\*ask\_crack\_to specifies which crack-method describes the volume.

\*rc defines the semi-edge length of the interior square of the section.

\*ri defines the radius of the interior circle of the section.

**\*re** defines the radius of the exterior circle of the section.

\*nbc is used to define the number of sub-squares dividing interior square part of the section.

\*nbri defines the number of radial cuts between the square part and the interior circle.

\*nbre defines the number of radial cuts between the interior and the exterior circles.

\*delta is used to define the square deformation.

\*delta\_p used to specify the height pyramid elements.

- **\*connect** used to specify volume should be connected to a hexahedral mesh: a layer of pyramidal elements is inserted.
- \*opening used to create a volume with a meshed crack open in direction propagationdirection-vector. Initialy this crack is filled by elset "compl".

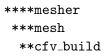
Various elsets are created:

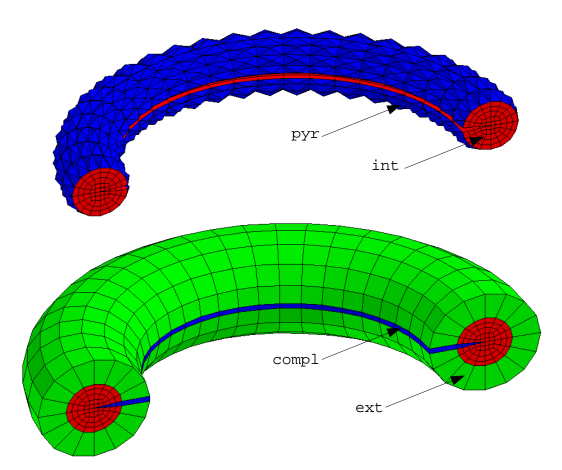
- "int" contains the interior part of the volume with only hexahedral elements (on which integrations should be computed).
- "pyr" contains the pyramidal elements used to connect the volume to tetrahedral meshes.
- "ext" contains external pyramidel elements if the connect keyword is specified.
- "compl" contains elements inserted inside the crack if the opening keyword is specified.

Various nsets are created:

- "front" contains the nodes located at the crack front.
- "skin" contains the nodes located on the skin made of pyramidal elements used to connect the volume to tetrahedral meshes.

```
****mesher
***mesh volume.geof
  **open crack.geof
  **cfv_build
   *liset LISET_FRONT (-1. 0. -1.)
   *rc .3
   *ri .6
   *re 1.
   *nbc 2
   *nbri 1
   *nbre 2
   *opening .1
   *delta .07
   *delta_p .2
   *connect 2.
****return
```





# $** check\_orientation$

#### **Description**:

This command verifies that all elements are correctly mapped such as to create a valid geometry (i.e. face normals all pointing out and properly connected).

#### Syntax:

This command takes an optional command to select the elset upon which the orientation check should be made.

\*\*check\_orientation
\*elset eset-name

**\*elset** specify an element set to check and adjust so as to be correctly defined. By default the element set is ALL\_ELEMENT.

# \*\*classical\_to\_zstrat

#### **Description**:

This command is used to transform 3D volumic elements into "Z-strat" solid elements. Z-strat elements are quadratic in the plane and linear in their thickness. They are typically used for composite applications.

# Syntax:

The command has the following syntax:

\*\*classical\_to\_zstrat
 \*axis axis-number
[ \*layered bool ]
[ \*elset elset-name ]

**\*axis** is a number defining the linear axis.

- \*layered is a boolean indicating that a layered geometry should be used (e.g. c3d12l or c3d16l instead of c3d12 and c3d16). The default value is not layered (FALSE).
- **\*elset\_name** indicates that the mesher should apply only to the elements in the given elset. The default is to apply on ALL\_ELEMENT.

```
**classical_to_zstrat
*axis 3 % elements become linear in the z axis
*layered TRUE % make layered elements
*elset E1 % apply on E1 elset
```

# $** condense\_out\_elset\_domain$

#### **Description**:

This mesher is for parallel problems, where a specific feature across the mesh is desired to be wholely contained in one domain. The mesh should already be split into domains, from which the elements in the given elset will be "condensed out" into a new domain (i.e. the original split should be n - 1 domains, with n the total desired number). Note that doing this does not ensure that the domains will be well sized for balanced load, or that poorly conditioned partial domains with isolated small groups of elements could be created.

#### Syntax:

The command has the following syntax:

**condens	se_out_elset_domain	
*elset $e$	elset-name	

\*elset specifies a pre-existing elset name which will make up the new domain.

# \*\*compute\_predefined\_levelset

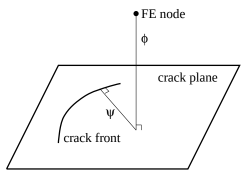
#### **Description**:

This command is used to generate levelsets (scalar field given at nodes of the FE mesh) defining cracks with classical geometries in an xfem analysis (page 3.205).

In this case the discontinuity is described by 2 levelsets  $\phi$  and  $\psi$ , where (see figure):

- $\phi$  is the signed distance of a node to the crack plane,
- $\psi$  is signed the distance to the crack front of the orthogonal projection of a node on the crack plane.

such that points for which  $\phi = \psi = 0$  are located on the crack front, and positive  $\psi$  values correspond to nodes ahead of the front.

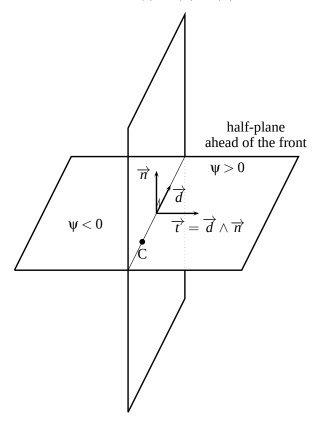


# Syntax:

```
**compute_predefined_levelset
    *type (circular|elliptic|line)
[ *rho radius ]
    *center c
    *normal (n)
[ *rho2 minor_radius ]
[ *direction (d) ]
[ *do_fem_output ]
[ *elset_name name ]
[ *elset_distance dist ]
[ *phi_name phi_file ]
[ *psi_name psi_file ]
```

- command \*type is used to specify the crack geometry (predefined types available are: circular, elliptic, line),
- *radius* is the radius (circular cracks) or the major radius value (elliptic cracks). It has no effect for line cracks.
- (c) is a vector defining the position of the centre for circular or elliptic cracks. In the case of line cracks, the coordinates of a point on the crack front is expected.
- (n) is a vector defining the normal to the crack plane (see figure above),

- for elliptic cracks, command rho2 is needed to specify the minor radius value.
- command \*direction defines a vector (d) whose meaning is the following, depending on the particular geometry:
  - the direction of the minor axis for an elliptic crack. Note that in this case (d) is expected to be orthogonal to (n) (otherwise an error occurs).
  - a vector parallel to the crack front for line cracks. The sign of this vector also defines the crack propagation direction in this case, as described on the following figure:  $\psi > 0$  is obtained in direction  $(t) = (d) \land (n)$



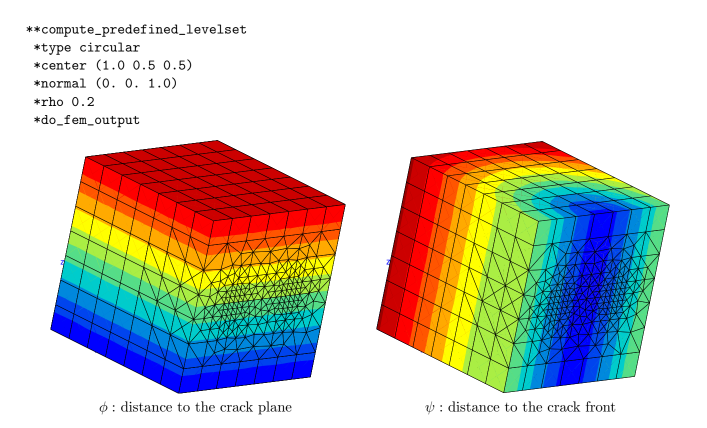
- optional command  $*do_fem_ouput$  allows the generation of FE results with  $(\phi, \psi)$  values calculated on nodes of the input mesh (see the example for the iso-contours generated).
- command **\*elset\_name** creates an elset (with name *ename*) containing elements situated within a distance of less than *dist* from the crack front.
- commands **\*phi\_name** et **\*psi\_name** allows to redefine the names of the output ASCII files used to store the levelset values on the mesh nodes (default names are "**phi.dat**" and "**psi.dat**"). Those files are used to define the discontinuity by the **\*\*\*xfem\_crack\_mode** command.

\*\*\*\*mesher

#### \*\*\*mesh

 $** \texttt{compute\_predefined\_levelset}$ 

# Example:



# \*\*continuous\_liset

## **Description**:

This command is used to reorder a scrambled line set. It first looks for a potential line set beginning, and then puts all other segments in the right order. The result is a well ordered line set, even is the initial segment ordering is completely scrambled.

# $\underline{Syntax}$ :

The command has the following syntax:

\*\*continuous\_liset
 \*liset\_name liset-name
[ \*start\_at\_node node-id ]
[ \*start\_near (x,y,z) ]

The **\*liset\_name** is used to indicate the name of the source line set to be re-ordered. If this line set is a loop (hence has no "natural" beginning), the optional **start\_at\_node** parameter specifies where to begin the liset. Alternatively, the **start\_near** parameter specifies the localization of this first node its (possibly approximate) coordinates.

# \*\*crack\_2d

## **Description**:

This command is used to insert a crack in a 2D mesh and apply 1/4 node displacements at the tip.

A duplicate of the command exists which automatically modifies the nodal positions on edges surrounding the crack tip to 1/4 node positions to simulate a singular field better. To get this option replace crack\_2d with crack\_2d\_quarter\_nodes

# Syntax:

The command has the following syntax:

```
**crack_2d_quarter_nodes
*liset liset-name
[ *node node-id ]
[ *crack_nset nset-name ]
[ *half ]
```

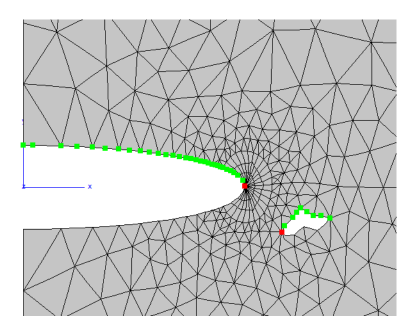
The options for this command are described below:

- **\*liset** give the liset name which describes the crack plane. This liset will be kept after the meshing operation.
- **\*node** Give node ids or nset names indicating nodes which are crack tips. These nodes terminate the crack surface creation.
- \*crack\_nset enter a name for a new nset to be created, which makes up the opposing surface from the crack definition liset. This lets one do contact, or other manipulations on the newly created surface.
- \*half Only make a closed crack tip at the first end of the given liset. That is for an edge-crack, make the line set originating where the crack tip is to be and leading out to the free surface.

# Example:

An example use is shown below taken from the test case Jint\_test/INP/ccp\_2.inp which is shown in the figure below.

\*\*\*\*mesher \*\*\*mesh \*\*crack\_2d



This case includes 2 cracks, one of which is open ended (the first) and one which automatically detects the crack tip nodes.

```
****mesher
```

```
***shell
  rm -f ccp_2.geof
  Zrun -B ccp_2.mast
***mesh ccp_2
 **open ccp_2.geof
 **crack_2d_quarter_nodes
 *liset crack
 *half
 **crack_2d_quarter_nodes
 *liset crk
****return
```

# $** crack_3 d_q uarter_nodes$

#### **Description**:

This command is used to set 1/4 node positions for adjacent faces of a 3d crack, as defined by a line set. It does not yet provide crack insertion via splitting face sets.

## Syntax:

The command has the following syntax:

\*\*crack\_3d\_quarter\_nodes
\*liset liset-name

where the *liset-name* is the name of a valid line set defining the crack tip line.

# \*\*create\_interface\_elements

#### **Description**:

This mesher takes a pre-existing boundary set and creates interface elements (e.g. debonding) at that location. The mesher can be used either between a boundary set and an existing node set, which assumes that best and next are not actually connected, or by using a boundary set only, where additional nodes will be inserted to make an interface condition.

## Syntax:

The command has the following syntax:

>	<pre>**create_interface_elements</pre>
	*boundary bset-name
Γ	<pre>*elset elset-name ]</pre>
Γ	<pre>*nset nset-name ]</pre>
Γ	*axi ]
[	<pre>*reduced ]</pre>

flag that the interface elements are axisymmetric. Default is not axisymmetric. \*axi

- **\*boundary** indicates the boundary set from which to create interface elements. There will be 1 interface element per boundary segment in the set.
- \*elset gives the name of the elset to create with the new interface elements. If this command is not included the name interface will be used for a new elset.
- **\*nset** indicates the opposing node set name. This option indicates that the mesh is already disconnected at the interface.
- **\*reduced** flag indicating reduced integration. Default is full integration.

# \*\*create\_interface\_elements\_between\_elsets

#### **Description**:

This mesher takes pre-existing element sets and creates interface elements (e.g. debonding/cohesive elements) at their common boundaries.

## Syntax:

The command has the following syntax:

;	**create_	interface_elements_between_elsets
	*elsets	elset-name1 elset-name2 [ elset-name(n) ]
[	*axi	]
[	*reduced	]
[	*remove_	sets ]

- \*elsets gives the names of the elsets between which the boundary will be created (note: elsets and not elset). The keyword ALL\_ELSET may be used to designate all elsets (except the standard elset ALL\_ELEMENT) present in the mesh. Names for the new element sets will be automatically generated, and a new elset EI\_ALL will be generated containing all new interface elements, as well as an elset VOL\_ALL containing all volume elements (so that the union of EI\_ALL with VOL\_ALL should yield ALL\_ELEMENT).
- \*axi flag that the interface elements are axisymmetric. Default is not axisymmetric.
- **\*reduced** flag indicating reduced integration. Default is full integration.
- **\*remove\_sets** indicates that only the interface elements should be kept, and that the associated elsets generated during the meshing are thrown away. The default is to keep all elsets.

# \*\*create\_interface\_elset

#### **Description**:

This mesher applies interface (debonding) elements between all elements of a given elset.

## <u>Note</u>:

Currently (Z-set 8.3 and newer), this command is implemented for linear 2d elements only (i.e. it will create i2d4 elements).

#### Syntax:

The command has the following syntax:

\*\*create\_interface\_elset
\*apply\_to elset-name
\*elset\_name new-elset-name

- \*apply\_to indicates the element set which is to be broken up and have interface elements inserted at each element edge. The default element set is ALL\_ELEMENT.
- **\*elset\_name** The new element set containing the interface elements. The default name is **debond**.

# \*\*cut\_surface

#### **Description**:

This command performs a robust surface intersection operation. It requires a 3D volume mesh that better be refined (using Distene remeshing tools) closely to a given meshed surface.

Resulting output is made of a conform surface mesh (elset Nsurface) decomposed in various elsets: parts of original elsets boundaries (Nskin) and their newly cut parts (Nskinc), created cut surface (Nlip). Some output nset are also produced: Nlip containing created cut surface nodes, Nskin skin nodes, Nfront front nodes (for surface containing a front liset i.e. for 3D crack insertion), Nseg\_pb containing original volume mesh edges nodes for edges cut an even times by the cut surface (smaller original volume elements are required in this area, as cut surface curve radius must be too small there).

This fast and robust algorithm is based on the given volume elements intersection, thus a remeshing process is required before FE computations (for better surface meshing, and volume filling), on the output Nsurface.

Syntax:

\*\*cut\_surface
[ \*elset elset-name-list ]
[ \*elset\_to\_cut elset ]
[ \*nset\_to\_cut nset ]
[ \*nset\_not\_to\_cut nset ]
[ \*surface elset-name ]
[ \*tolerance tol ]
[ \*front liset-name-list ]
[ \*front\_ini nset ]
[ \*filter tol ]
[ \*allow\_quad ]
[ \*inside ]

- **\*elset** list of elsets that should be cut by surface. If more than one is given, boundaries between the elsets will be kept during the process (to be able to keep various elsets in the output).
- **\*surface** is the given surface elset mesh to intersect with.
- **\*elset\_to\_cut** only this elset may be cut (default is to cut ALL\_ELEMENT).
- \*nset\_to\_cut only edges contained in this nset may be cut.
- \*nset\_not\_to\_cut edges contained in this nset are not allowed to be cut.
- \*tolerance if an edge intersection node is closer from an edge node than this distance value assume the edge node is on the cut surface. In most cases (in fact, in all currently tested cases), the zero default value of this parameter has proved to be robust.
- **\*front** list of lisets representing the extremity of the cut surface that must be accurately inserted (i.e. for 3D crack insertion).

- **\*front\_ini** initial crack front, for newly cracked surface insertion (this nset represents the previous crack front), the newly surface will contain this previous front.
- \*filter tolerance distance used to fuse pairs of same created surface elements (i.e. front 3D crack advance, to keep only one side of previously built lip). Default value is zero (no surface will be fused), if required a usual value is 1.e-5 times the mesh characteristic length.
- **\*allow\_quad** quad elements can be created during the algorithm. Default is not allowed as non-planar elements can be created an make a surface remeshing process fail.
- \*inside crack surface advance may not pass through the boundary of the mesh (e.g. when 2 cracks are coalescing).

# $^{**}dg_{-}transform$

#### **Description**:

This command is used to transform a mesh for continuous Galerkin formulation to a mesh suitable for discontinuous Galerkin formulation.

## Syntax:

The command has the following syntax:

**dg_transform		
[ *elset	]	elset-name
[ *bset	]	bset-name
[ *name	]	name-of-bset-to-create

**\*elset** specifies elset where operations should be made.

- **\*bset** specifies beet where discontinuity should be inserted (should not work well for now, see VC if needed).
- \*name specifies name that will be given to pairs of bset where DG interface formulation will be imposed (format will be namea0/nameb0, namea1/nameb1, ... ,nameaN/namebN).

# $**deform\_mesh$

#### **Description**:

This mesher deforms the active mesh via results taken from a previous analysis. It is a simple implementation assuming 1-1 correlation between the meshes. It can be used to load any types of displacement results, but is most commonly used to create bucking analysis imperfections via the modes of a previous eigen solution.

## Syntax:

The command has the following syntax:

\*\*deform\_mesh
\*map solution-no
\*input\_problem pb\_name
\*magnitude disp
\*format fmt

- \*format gives the solution format to use for deforming the mesh. These can be any of the ones with a valid results database implementation (see e.g. page 2.117).
- **\*input\_problem** The prefix name of the problem results which will be used for the deformation.
- \*map specifies the solution or map number from the results file which will be used to get the deforming displacements. It must be one listed in the .ut file. The default is to take the last output available.
- **\*magnitude** the scale factor for the displacements to be applied to the mesh. The default is 1.

# Example:

An example for deforming a mesh with a previous Eigen solution follows:

```
****mesher
 ***mesh buckle.geof
 **open T_54_9.geof
 **deform_mesh
 *map 1
 *input_problem lanczos
 *magnitude 0.05
```

# \*\*delete\_elset

#### **Description**:

This command is used to remove elements from the mesh by a given element set name. Elements are also removed from any other elset of which they are part. Nodes (in nsets for instance) and integration points (in ipsets) which are "orphaned" by this command will be removed as well. If elsets or ipsets become empty as a result of this, they will also be removed. The command can be used to create cavities in the mesh, which are otherwise difficult to obtain.

#### Syntax:

The delete elset command simply takes a list of valid elset names. There are no options with this command.

\*\*delete\_elset elset-name

#### Note:

If you wish to simply remove the *element set* and not the elements themselves, use the **remove\_set** command (see page 2.95).

# \*\*div\_quad

#### **Description**:

This mesher is used to perform a quadrilateral to triangular transformation.

## Syntax:

The command has the following syntax:

\*\*div\_quad
[ \*elset elset-name ]

**\*elset** gives an elset for which the division will be applied. the default elset is **ALL\_ELEMENT**.

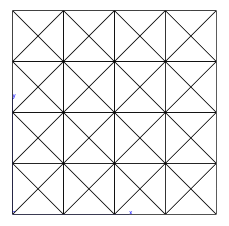
# Note:

This mesher does not preserve any sets related to the elsets which are sub-divided.

#### Example:

The following shows the division take for a ruled mesh.

7		



\*\*elset

## **Description**:

This command creates an element set according to various input data. If the element set already exists, new elements will simply be added. Elements will be included in the set only once.

## Syntax:

The elset command takes a variety of options to specify the elements to be added (and each of them may be specified more than once):

```
**elset name
[ *add_elset eset1 ... esetN ]
[ *allow_partial ]
[ *attached_to_nodes id_1 ... id_N ]
[ *attached_to_nset nset1 ... nsetN ]
[ *elements id_1 ... id_N ]
[ *func function ; ]
[ *not_in_elsets eset1 ... esetN ]
[ *remove_elset eset1 ... esetN ]
[ *sequence start_id end_id [ increment ] ]
[ *use_elsets eset1 ... esetN ]
```

where the options can be taken from the following:

\*add\_elset Adds all the elements in the named (pre-existing) element sets.

- \*allow\_partial Indicates that a function describing element node positions for the elset is true if one or more of the nodes meets the given function criteria. By default, all nodes must fit the equation.
- \*attached\_to\_nodes indicates that all elements attached to nodes with the given ids are to be added to the elset.
- \*elements Enter element ids in directly. Non-existing element ids will be silently ignored.
- \*func Enter a function which is used to select the elements. The function should be a chain of multiplied boolean expressions in the coordinate space (see example below).
- \*not\_in\_elsets will add the complement to the given element sets as a new elset. Not compatible with other options such as \*func and so on.
- - $\dots$  continued

\*\*\*\*mesher \*\*\*mesh \*\*elset

- \*sequence will add a sequence of elements from *start\_id* to *end\_id* with increment *increment* (Z-set 8.4 and newer). The latter can be negative, but in that case, *start\_id* should be greater than *end\_id*. If this is not the case, *-increment* will be used instead. If *increment* is omitted, a default value of 1 is used. Non-existing element ids generated with this command will be silently ignored.
- **\*use\_elsets** use only the elements in the named element sets to check the given function against. The default is to use all elements in the mesh.

#### Example:

Some example uses follow.

```
% Shows the use of functions
  **elset front
   *func x>0.0:
  **elset bottom
   *func y<0.0;
  **elset look
   *func (y<0.0)*(z>20.0);
   *allow_partial
\% Creates/adds to the elset "first_10" with elements 1 through 10
% (better use the *sequence command)
  **elset first_10
   *elements 1 2 3 4 5 6 7 8 9 10
% Creates/adds to the elset "sequence_example" with elements
% 1 4 29 30 32 34 ... 48 50 53
  **elset sequence_example
   *elements 1 4 29
   *sequence 30 50 2
   *elements 53
% Creates/adds to the elset "inverse_sequence" with elements
% 50 48 46 ... 32 30
  **elset inverse_sequence
   *sequence 50 30 -2
% Creates/adds to the elset "twice" with elements 1 through 15
  **elset twice
   *sequence 1 10
  **elset twice
   *sequence 5 15
```

\*\*\*mesher
\*\*\*mesh
\*\*elset\_by\_element\_type

# $**elset\_by\_element\_type$

# **Description**:

This mesher creates one elset per element type.

#### Syntax:

The command has the following syntax:

\*\*elset\_by\_element\_type

and takes no option.

# \*\*elset\_explode

#### **Description**:

This mesher is used to provide a per-elset exploded view of a mesh, which can be useful for presentation purposes. Because this mesher adds nodes between elset interfaces, the display of ctnod results files will be invalidated (including the displacements), and there is no checking to that  $regard^2$ 

# Note:

By default, this mesher does not actually explode the elsets! If the shrink factor is not specified, it merely creates the nodal interfaces in order to do so later with a **\*\*function** (page 2.60) or **\*\*translate** (page 2.117) mesher afterwards.

#### Syntax:

\*\*elset\_explode
[ \*elsets elset-list ]
[ \*shrink factor ]

- **\*elsets** list of elsets used as domains to separate. By default, all elsets are used, which can lead to bad result.
- \*shrink specify the factor used separate the different parts of the mesh. The new nodes position is computed as  $\underline{\mathbf{v}}' = \underline{\mathbf{v}} + f \cdot (\underline{\mathbf{e}} \underline{\mathbf{c}})$  with  $\underline{\mathbf{v}}$  and  $\underline{\mathbf{v}}'$  respectively the initial and final node position, f the shrinking factor,  $\underline{\mathbf{c}}$  the center of the mesh  $\underline{\mathbf{e}}$  the center of the elset.

# Example:

One can use this mesher after a calculation has been performed to separate the mesh, then reassign the visualization mesh in the *problem.ut* file. The following splits the 2 elset problem zsteel\_rubber found in the static test directory:

```
***mesher
***mesh toto
 **open zsteel_rubber.geof
 **elset_explode
 *elsets acier rubber
 **translate
 *elset_name acier
 *x -20.0
 *y -20.0
```

The ut file can be now modified to use the toto.geof file just created.

```
**meshfile toto.geof
**node U1 U2 Pn RU1 RU2 RPn
**integ sig11 sig22 sig33 sig12 dv F11 F22 F33 F12 F21
```

 $<sup>^{2}</sup>$  actually the record sizes will be interpreted incorrectly, and the results loaded are therefore a randomized mix of values. No crash occurs however.

\*\*element

- 1 1 1 1 4.0000000000000000e+00
- 2 1 1 2 8.000000000000000e+00
- 3 1 1 3 1.20000000000000e+01
- 4 1 1 4 1.60000000000000000e+01
- 5 1 1 5 2.0000000000000000e+01

And the results can be then viewed on the exploded mesh via Zmaster. When doing this make sure and turn off deformed mesh and to include either (or both) **\*\*contour\_by\_element** and **\*\*value\_at\_integration** output options in the calculation.

# \*\*elset\_split

#### **Description**:

The **\*\*elset\_split** command is used to create parallel problem domains and write a *problem*.cut file based on different named elsets. Normally one would use an automated mesher such as **\*\*metis\_split** to do this, but the elset split ensures repeatability and fine control over the process.

# <u>Note</u>:

Any elements in the mesh which are not assigned to a domain using the **\*domain** sub-option will be assigned to the 1st domain.

#### Syntax:

The command has the following syntax:

```
**elset_split
*no_binary
*domain elset-name
...
*domain_startswith stem
```

- **\*no\_binary** indicates that no binary cut file should be written (default is to write the binary cut).
- \*domain enter new domain made up of the elements in the given elset name. Repeat the command for all domains.
- **\*domain\_startswith** is a shorthand to automatically enroll all domains whose name starts with the given *stem*.

# Example:

There are some examples of this command in the Parallel\_test directory.

```
****mesher
***delete_file mpc1b_small.geof
***shell Zrun -B mpc1b_small.mast
***mesh mpc1b_small
 **open mpc1b_small.geof
 **quad_to_lin
 **elset right *add_elset R2 R4
**elset left *add_elset R1 R3
**elset_split
 *domain right
 *domain left
 *no_binary
```

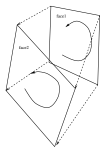
#### \*\*extension

#### **Description**:

This command extends planar (2D or 3D shell/face) geometries into 3D. One can use either an element set or a face set which is already defined.

#### Note:

For extensions which are between two element sets, care must be taken to properly define the sets. Order of domain selection is critical here. The figure here shows the proper order for ruled mesh definition in Zmaster. Select the edge segments in the sense of the arrowed paths when taking the same view to your mesh extension.



#### Syntax:

**extension		
*elset	$eset1$   All_Elset	
*elset2	eset2	
*new_elset	$new\_elset$	
*distance	dist	
*num	num	
*prog	prog	
*dir	dir-vec	
*flipit		
*reduced		
<pre>*invert_fasets</pre>		
<pre>*remove_initial_nsets</pre>		

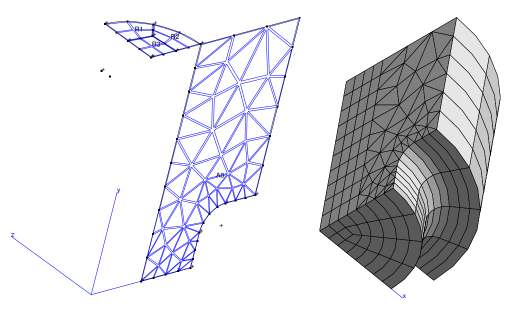
- \*elset eset1 defines the element set or faset which will be extended. There is no limitation on the type of elements or faces contained on the face. Mixed order is not allowed however. The option ALL\_ELSET extends each elset of the mesh<sup>3</sup>. If no elset is specified, the elset ALL\_ELEMENT is taken.
- **\*elset2** eset2 This optional command makes the extension between two mesh/face sets as given. Note that the mesh topology must be exactly the same. This command is most useful with the mapped mesh domains available in Zmaster. The degenerate case of 2 edges being the same (so the extended mesh forms a wedge) is handled. Do not use **\*distance** with this option. This option is disabled when the first elset name is ALL\_ELSET.
- \*new\_elset new-elset This optional command gives the elset name for the created elements
  instead of being the same as the input name. This is disabled when the input elset name
  is ALL\_ELSET.

<sup>&</sup>lt;sup>3</sup>Available in version 8.2 or newer

- \*distance This command is used to define the extension distance, *dist* (real). Do not use it when the extension is between two meshes.
- \*num This command takes an integer value for *num* in order to define the number of elements which will be created in the cross section.
- \*prog Geometric progression of the extension.
- \*dir The direction along which to extend. This direction is specified in vector form and does not need to be normalized. The default is (0. 0. 1.).
- \*flipit Used when the created elements are inverted.
- **\*reduced** Create reduced integration elements.
- \*invert\_fasets In order to properly calculate the normals (and other properties) of a face set, the nodes in each element need to be ordered in a certain manner within the faset. This is very important for applying correctly many types of boundary conditions. After extension, the nodes in the newly created faset have the same order as in the parent faset, which implies that the normal points in the same direction, which may not always be the desired direction. This option allows the order within a faset to be inverted, so that the direction of the normal of the newly created faset is reversed as well.
- **\*remove\_initial\_nsets** Initial nsets are also extended and the suffix "-ext" is added to these new nsets names. This option allows to remove initial nsets and keep extended one with the initial names.

#### Example:

The following example shows the use of extensions in different directions, and making use of the 3d plane meshing in Zmaster.



\*\*\*\*mesher \*\*\*mesh \*\*extension

\*\*extension \*elset R1 \*dir (0. 1. 0.) \*distance -2.0 \*num 10 \*\*extension \*elset R2 \*dir (0. 1. 0.) \*distance -2.0 \*num 10 \*\*extension \*elset R3 \*dir (0. 1. 0.) \*distance -2.0 \*num 10 \*\*sweep \*elset AO \*angle 90.0 \*num 4

## \*\*extension\_along\_nset

#### **Description**:

This mesher is a modification of the **\*\*extension** mesher in order to allow an extension path to be defined with a set of nodal points.

#### Syntax:

This command takes the same commands as the **\*\*extension** command (see page 2.53) and 2 additional commands relating to the path.

\*\*extension\_along\_nset
\*nset\_to\_follow nset-name
\*global\_normal normal-vector

Note that many of the commands from extension, though available, make no sense for this mesher because of the pre-defined path (such as num, direction, etc).

\*global\_normal Input a vector allowing the initial face plane to be calculated. The tangent vector of the path is calculated based on linear segments from and to adjacent nodes.

**\*nset\_to\_follow** Enter an existing nset path for the mesh extension to take place.

#### Example:

The following example shows extension along an nset, using a curve generated in Zmaster and meshed using line 13d2 line elements (under Ruled mesh). A boundary set called lines was also created in Zmaster which creates the nset along which the extension will take place.

```
****mesher

***mesh extension_nset_linear
**open curve_linear.geof
**union
*add zzz.geof
*elset zzz
*tolerance 0.
**extension_along_nset
*nset_to_follow lines
*global_normal (1. 0. 1.)
*elset zzz
**delete_elset lines
****return
```

## $** extract\_surface$

#### **Description**:

This command builds beets corresponding to each surface of a given elset.

#### Syntax:

The command has the following syntax:

\*\*extract\_surface
[ \*elset ] elset-name
[ \*criterion ] angle

\*elset restricts the operation to the named elset (defaults to ALL\_ELEMENT),

\*criterion is the angle (in degrees) from which two adjacent elements are considered part of different surfaces (defaults to  $45^{\circ}$ ).

# \*\*extrude\_shell

#### **Description**:

The command **\*\*extrude\_shell** is used to extrude 3d shell elements into "Z-Strat" solid elements.

## Syntax:

The command has the following syntax:

\*\*extrude\_shell
[ \*elset\_name elset-name ]
[ \*layered ]
[ \*thickness thickness ]

- **\*elset\_name** indicates that the mesher should apply only to the elements in the given elset. The default behavior is to use ALL\_ELEMENT for the elset.
- \*layered is a flag setting that a layered geometry should be used (e.g. c3d121 or c3d161 instead of c3d12 and c3d16). The default is not layered.
- **\*thickness** gives an uniform thickness thickness to the elements. Default value is taken arbitrarily as 0.1.

# \*\*faset\_align

#### **Description**:

This command aligns fasets such that their "normals" are aligned in the same direction signwise. The normals are still normal to the face.

The faset normal alignment is very important in order that boundary conditions (e.g. pressure) and contact surfaces are correctly defined.

## Syntax:

\*\*faset\_align
\*faset faset1 ... fasetN
\*normal direction
\*towards point
\*away\_from point

The normal must be entered using vector notation (c.f. page 1.3). If it is not entered at all, either the option **\*towards** or **\*away\_from** must be entered with a "target" point entered in vector form.

This command is now replaced by the more general **\*\*bset\_align** (page 2.18).

#### Example:

The following is from the example problem Straw/LIN\_PLASTIC/straw.inp

```
***mesh straw
 **open straw_base.geof
 **nset fix
  *point 0. -1. 0.
 **nset RO-surf
  *func (y>=0.0)*(y<=1.0);
 **nset R1-surf
  *func (y<=0.0)*(y>=-1.0);
 **bset target
  *use_nset nodes-ext
  *elset A2
 **faset_align *faset R6.1
                             *normal
                                      (0. -1.0 0.)
 **faset_align *faset target *normal
                                      (0. -1.0 0.)
```

## \*\*function

#### **Description**:

This command modifies nodal positions according to the given function. These functions are as described in the chapter on functions and scripting (page 5.2). Function transforms can be applied to the whole structure (default), to nset(s), and to the nodes of an elset or elsets.

#### Syntax:

**function		
*func	func(x,y,z)	
*xtrans	func(x,y,z)	
*ytrans	func(x,y,z)	
*ztrans	func(x,y,z)	
*nset	$nset1 \dots nsetN$	
*elset	$elset1 \dots elsetN$	

- \*func used for function definitions which are henceforth available. One can use a "learn" command line option as well to specify an external file with function definitions.
- \*xtrans \*ytrans \*ztrans functions which specify a coordinate reassignment in x, y, or z directions. Note that these are absolute values (trans is a bit of a misnaming), so an actual translation of 0.40 in the z direction would use \*ztrans z-0.40;

**\*nset** specify nset(s) for the function to be applied to.

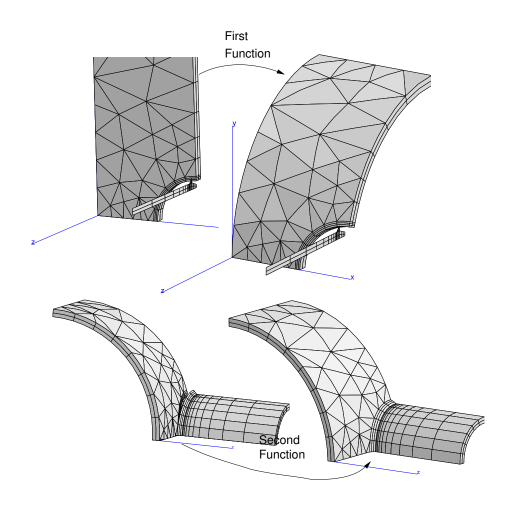
**\*elset** specify elset(s) for the function to be applied to.

#### Example:

The following are the function statements from the example \$Z7PATH/test/Mesher\_test/INP/func-xform.inp

```
**function
 *elset pipe1
 *function_def R:= 6.366197722 + z;
 *function_def theta:=y/6.366197722;
 *function_def Z:=R(z)*cos(theta(y))-6.366197722;
 *function_def Y:=R(z)*sin(theta(y));
 *ztrans Z(y,z);
 *ytrans Y(y,z);
...
**function
 *elset weld
 *elset pipe2
 *function_def f1:= (z>=1.0)*1.0 + (z<1.0)*(z>0.0)*z + (z<=0.0)*0.0;
 *function_def f1:= (z>=0.2)*1.0 + (z<0.2)*(z>0.0)*(z*z/0.04);
 *ztrans f1(z)*z + (1.-f1(z))*Z(y,z);
 *ytrans f2(z)*y + (1.-f2(z))*Y(y,z);
```

\*\*\*\*mesher
\*\*\*mesh
\*\*function



#### \*\*fuse\_nset

#### **Description**:

This command joins nodes forming a "mini-union" algorithm, or can also be used to "clean up" a mesh with possible gaps in nodes. This is very useful when batch mesher transforms are applied to an initially exploded arrangement of sub-meshes, which are then to be transformed together and glued. It is also an effective method to make cracks (in the absence of using **\*\*crack\_2d**).

#### Syntax:

\*\*fuse\_nset
[ \*nset1 nset-name ]
[ \*nset2 nset-name ]
[ \*tolerance tol ]
[ \*cleanup\_mesh ]
[ \*cleanup\_nset nset ]
[ \*average\_locations ]
[ \*allow\_mixed\_fuse ]

\*allow\_mixed\_fuse fuse nodes of different dimension.

- \*average\_locations this indicates that the new node position for joined nodes is in the middle of their original positions. By default the node position in node set one is taken.
- \*cleanup\_mesh this changes the mode of operation to a "clean up" mode, where all nodes closer than the tolerance are joined.
- \*cleanup\_nset cleanup only the nodes of a given nset.
- \*nset1 when not running in \*cleanup\_mesh mode, this specifies the name of one of the two node sets which will be compared. It is required for standard use.
- **\*nset2** enter the node set name to be compared with *nset1*.
- \*tolerance enter a real value for the magnitude of distance between nodes at or below which they will be considered the same (its default value is that of the global parameter Mesher.MeshFusion).

# \*\*geof\_format

#### **Description**:

This mesher is used to convert the mesh format from ASCII based, human readable form to a binary file, and back again. Please note that the binary file will be read by Zebulon and Zmaster much faster than the formatted mesh, and will also take up less disk space.

# Syntax:

The command has the following syntax:

\*\*geof\_format
\*format file-name
\*unformat file-name

The options **\*format** and **\*unformat** set the operation to respectively convert the current mesh file to a formatted .geof file, or convert the formatted file to a binary .geo file. The default operation is to format the mesh. The mesh will be immediately saved using the current mesh name (set after the **\*\*\*mesh** command). Thus mesher commands after the **\*\*geof\_format** will not be included in that mesh file.

#### $\underline{Note}$ :

Alternatively, the mesh is automatically saved in binary mode if its filename ends with .geo, as in \*\*\*mesh mymesh.geo.

# \*\*hexa\_to\_tetra

#### **Description**:

This command is used to tetrahedrize a mesh (i.e. transform all its c3d8 or c3d8r elements to c3d4).

#### Syntax:

The command has the following syntax:

\*\*hexa\_to\_tetra
[ \*elset ] elset-name

**\*elset** restricts the operation to the named elset

#### Note:

There is currently another mesher that has the same objective: hexa\_only\_to\_tetra. The latter's main advantage is that it tries to keep a conforming mesh (i.e. cuts the faces coherently between two adjacent elements). This cutting operation is not trivial, nor always possible. So you should always check the result to see if the resulting mesh is conforming. Currently it only knows how to handle c3d8, doesn't have the **\*elset** option, and doesn't update the bsets and elsets accordingly (nsets are unchanged by the cutting operation).

#### Syntax:

The command has the following syntax:

\*\*hexa\_only\_to\_tetra
[ \*seed ] element-rank
[ \*talkative ]

**\*seed** selects which element should be started with (default is the first element).

\*talkative (integer) produces more output, useful if something goes wrong. 1 adds a little more output, 2 is very chatty.

#### Example:

```
***mesher
***mesh plaque_tetra.geof
**open plaque.geof
**hexa_only_to_tetra
*seed 4561
*talkative 1
****return
```

# \*\*import\_abaqus\_pressure

#### **Description**:

This command imports ABAQUS input data for pressures on element faces into Zebulon. This allows more robust pre-processors with ABAQUS export (e.g. Patran, Femap, etc), or existing user programs, to define variable pressures over a surface.

# Syntax:

The command has the following syntax:

\*\*import\_abaqus\_pressure
\*bset bset-name
\*ptable\_file out-file-name
\*use\_dimension dim

**\*bset** assign the name of the new bset for the pressures to be applied to.

\*ptable\_file

 $*use_dimension$ 

# \*\*insert\_discontinuity

### **Description**:

This command fastly duplicates nodes of a bset to create a discontinuity. If an nset is given those nodes will not be duplicates in order to represent a 3D crack front for instance.

Resulting output contains an odd number of nset named side0, side1, side2, side3, etc. linked to each pair of sides of the discontinuity (a pair is built for each discontinous zone of the given bset).

Only for 3D meshes.

# Syntax:

\*\*insert\_discontinuity
[ \*elset elset-name ]
[ \*bset faset-name ]
[ \*nset nset-name ]

\*elset restrict operation to a given elset (default is ALL\_ELEMENT).

\*bset continuous or discontinuous faset where discontinuity must be built.

**\*nset** nodes that must not be duplicated (to create a crack front for instance).

### $**inverse\_bset$

### **Description**:

This command is used to inverse the geometric direction of line or face sets. The order of all the segments of the set is reversed, as is the nodal ordering of each segment, so the face or edge normals are inverted.

Any mix of lisets or fasets can be given to this command.

# Syntax:

\*\*inverse\_bset
 \*names 11 12 13
 ...

### $\underline{Note}$ :

This command supersedes the older **\*\*inverse\_liset**.

# $**inverse\_liset$

## **Description**:

This is used to revert the way a liset is described. It can be useful for instance for contact computations in which the target liset must be followed in an order so that impacting nodes are located on the right.

## Syntax:

\*\*inverse\_liset \*names 11 12 13 ...

Any number of liset can be inverted in the same pass. Liset names have to be declared after **\*names** option.

This command is deprecated as of 8.3 in favor of **\*\*inverse\_bset**.

# \*\*join\_bsets

### **Description**:

This command is provided in order to join bsets together<sup>4</sup> It is useful to use this command after aligning bsets seperately using the **\*\*faset\_align** command.

### Syntax:

The **\*\*join\_bsets** command requires a new bset name. and needs one or more existing bsets to be entered which will be joined in the new set.

\*\*join\_bsets new-bset
 \*bsets bset1 ... bsetN
 ...

where *bset1* indicates a character name of a pre-exisiting boundary set (faset or liset).

<sup>&</sup>lt;sup>4</sup>There are no checks to eliminate duplicated segments, and lisets are not sorted.

# \*\*join\_nsets

### **Description**:

This command joins node sets. The original node sets are conserved. In the new node set the order of the nodes is the same as in the constituting node sets, and the order of the node sets within the new node set is as entered by the user. By default a check is carried out in order to prevent nodes from being added twice. This can be switched off with the **\*keep\_duplicates** option.

# Syntax:

The **\*\*join\_nsets** command requires a new node set name and one or more existing nsets to be joined into the new node set.

```
**join_nsets new-nset
*nsets nset1 [ nset2 ... nsetN ]
[ *keep_duplicates ]
```

where nset1 ... nsetN indicate the names of pre-exisiting nsets.

# \*\*lin\_to\_quad

### **Description**:

Use this command to increase the integration order of elements from linear to quadratic. The command always assumes that the midside nodes are linearly placed between the corner nodes, so if the linear mesh approximates a curve, the quadratic mesh will in fact model the chord faceted approximation.

### Syntax:

The command has the following syntax:

\*\*lin\_to\_quad
[ \*elset ] elset-name
[ \*no\_nset ]

- **\*elset** if specified, only the given elset is set to quadratic. The default is to run on the whole mesh.
- \*no\_nset if specified, nsets are not modified, and thus only contain nodes from the original
  linear mesh.

# <u>Note</u>:

In version 8.3 there is no corresponding alteration of the node sets and boundary sets which might be affected by this increase in order. The user should be careful to remove such unwanted (and in the bset case, invalid) sets by hand using the **\*\*remove\_set** command (see page 2.95).

# Note:

The inverse operation quad\_to\_lin, i.e. going from quadratic to linear elements, is described on page 2.89.

Ş

# \*\*make\_springs

### **Description**:

This command is supplied as a convenient means of adding spring elements to a mesh<sup>5</sup>. In particular, one noded springs which resist motion from the initial position can be added to nsets which is often a useful means of achieving stability in structures which are initially unstable for a quasi-static solution (e.g. soft springs added to a contact problem to allow a solution before contact begins).

# Syntax:

```
**make_springs spring-type new-elset
[ *connect_points vector1 vector2 ]
[ *connect_nodes node1 node2 ]
[ *nset nset_name ]
[ *nset_pair nset1 nset2 ]
[ *load_pmpc_eqn file-name ]
[ *proximity prox-val ]
[ *start_ele_id id-val ]
```

The algorithm checks the spring geometry whether is is a one or two noded spring, and uses the given options valid for that spring geometry.

- \*connect\_points used for 2 node elements, this command will link the two nodes closest to the given coordinates with a spring. The command may be given as many times as required.
- \*connect\_nodes used for 2 node springs, connecting by node id number rather than position. The command may be repeated.
- **\*nset** used for 1 node springs. Springs of the new set will be attached to each node contained in the node set.
- \*nset\_pair Join nodes between the two sets with a 2 node spring.
- \*proximity sets the algorithm to do a "proximity matching" between the two given nsets (thus \*nset\_pair must be used). For each node of the 1st nset the closest node at most prox-val away is linked up. So not all nodes in the 1st nset are required to be attached to springs, and each will have only one spring attached. The 2nd nset may however have multiple springs attached to one node.
- $*start_ele_id$  uses the given id number as the starting number for the new spring elements.

 $<sup>^{5}</sup>$ in Zebulon, 2 noded springs are truss elements. 1 node elements are not drawn in Zmaster, and 2 node elements appear as lines

\*\*\*\*mesher \*\*\*mesh \*\*make\_springs

### Example:

A simple example adding one node springs follows.

```
**nset spring_set
*nodes
    % Bolts, nuts (horiz)
    136 164 163    43 70 38
    633 609 610    637 638 661
    % bolts (vert)
    1622 1623 1635 1620
    1686 1688 1699 1700
**make_springs l3d1 SPR
*nset spring_set
```

### Example:

Here specific locations are connected with 2 node springs.

```
**make_springs l2d2 springs
*connect_points (0. 0. 0.) (1. 1. 1.)
*connect_nodes 25 341
```

### Example:

The last example uses proximity matching for frontal renumbering across a discontinuous mesh. The spring connectors are deleted right after doing the renumbering.

```
****mesher
***shell Zrun -B make_springs_renum
***mesh make_springs_renum
**open make_springs_renum.geof
**make_springs l2d2 springs
*nset_pair face_outside face_inside
*proximity 8.5
**renumbering frontal_only
**delete_elset springs
****return
```

# $**mesh\_quad\_cube$

### **Description**:

This command is used to quickly generate a cubic mesh. You can also refer to mesh\_quad\_parallelepiped 2.75.

# Syntax:

\*\*mesh\_quad\_cube
\*ncut n
\*size length

**\*ncut** Number of segments along the cube sides

\*size Length of the cube sides

# Example:

\*\*\*mesher
 \*\*mesh cube
 \*mesh\_quad\_cube
 \*ncut 10
 \*size 1.
\*\*\*\*return

# $**mesh_quad_parallelepiped$

### **Description**:

This command is used to quickly generate a cubic mesh. You can also refer to mesh\_quad\_cube 2.74.

### Syntax:

**mogh	and only
**IIesII_	quad_cube
*ncutx	nx
<pre>*ncuty</pre>	ny
*ncutz	nz
*sizex	length_x
*sizey	$length_y$
*sizez	length_z

\*ncutx Number of segments along the parallelepiped sides parallel to the x axis

 $\texttt{*ncuty} \quad \text{Number of segments along the parallelepiped sides parallel to the y axis}$ 

\*ncutz Number of segments along the parallelepiped sides parallel to the z axis

\*sizex Length of the cube sides parallel to the x axis

**\*sizey** Length of the cube sides parallel to the y axis

\*sizez Length of the cube sides parallel to the z axis

### Example:

```
***mesher
***mesh parallelepiped
**mesh_quad_parallelepiped
*ncutx 10
*ncuty 20
*ncutz 5
*sizex 10.
*sizey 20.
*sizez 5.
****return
```

# \*\*metis\_renumbering

### **Description**:

This is a re-numbering mesher which will reduce the fill-in generated during factorization of the global matrix with sparse direct solvers (sparse\_direct or sparse\_dscpack). The software is based on the Metis package developed by the Computer Science Department at the University of Minnesota, and used by permission. Information on the Metis package is available on the Web at the link:

http://www-users.cs.umn.edu/~karypis/metis/

# <u>Note</u>:

Since version 8.3 the nodal renumbering for sparse matrices is done automatically, so this command is no longer generally used.

### Syntax:

The command has the following syntax:

```
**metis_renumbering
[ *param_files file1 ... fileN ]
```

**\*param\_files** enables to renumber in the same time external parameter files.

# \*\*metis\_split

### **Description**:

This is a mesher routine to build a sub-domain *problem*.cut file to be used with the parallel solver. The software is based on the Metis package developed by the Computer Science Department at the University of Minnesota, and used by permission. Information on the Metis package is available on the Web at the link:

http://www-users.cs.umn.edu/~karypis/metis/

# <u>Note</u>:

One can also use the ONERA splitmesh program (see p.2.107).

### Syntax:

The command has the following syntax:

```
**metis_split
[ *check_domains ]
[ *dont_check_domains ]
[ *check_domains_iter iter ]
[ *domains num ]
[ *no_binary ]
[ *no_elset ]
```

- \*check\_domains indicates that the domains should be checked to see if there are elements attached by less than an edge in 2d, or a face in 3d. This is important to avoid rigid body and conditioning problems, especially with triangular or tetrahedral meshes. This option is on by default; use the \*dont\_check\_domains option to disable it.
- \*check\_domains\_iter modifies the maximum of iterations allowed in this domain verification procedure (default value: 10).
- \*domains specifies the number of sub-domains to be used. In the absence of this subcommand, the default value of 10 will be taken.
- \*no\_binary suppress the binary file, which is normally used if there are both .cu (binary)
  and .cut (ascii) files. Useful for hand-modifying the cut files, or for verification.
- $*no\_elset$  suppresses creation of additional element sets showing the domains which were created.

### Example:

An example use follows. This case has a problem with some domains only tied by MPCs, so springs are created before to establish total connectivity. The springs are removed after the split is done, and then the cut file is re-written using the **\*\*write\_cut\_from\_elsets** command to take into account the deleted elements.

```
***mesher
***mesh calcul_parallel
**open big_problem.geof
**make_springs 13d2 pmpc1 *load_pmpc_eqn mpc_set_1.equ
```

```
**make_springs 13d2 pmpc2 *load_pmpc_eqn mpc_set_1.equ
**metis_split
    *domains 8
    *no_binary
    **delete_elset pmpc1
    **delete_elset pmpc2
    **write_cut_from_elsets
****return
```

\*\*\*mesher
\*\*\*mesh
\*\*modify\_mesh\_and\_cut

# \*\*modify\_mesh\_and\_cut

### **Description**:

This command, used for parallel computations, allows to modify the *problem.cut* and *problem.cu* files.

### Syntax:

\*\*modify\_mesh\_and\_cut
\*enable\_node\_renumbering
\*store\_node\_renumbering
\*store\_nodes

- \*enable\_node\_renumbering writes the \*\*\*renumbering option in the *problem*.cut and *problem*.cu files to enable subdomain Metis renumbering (see page 2.76) at runtime. This option should be used only with the sparse direct linear solvers.
- \*store\_node\_renumbering renumbers all subdomains using Metis renumbering and writes the new subdomain nodes order in the *problem.cut* and *problem.cu* files.
- \*store\_nodes writes for each subdomain the list of nodes which belong to the subdomain (without any renumbering operation). This list is always written in the *problem.cut* and *problem.cu* files using the ONERA splitmesh program (see page 2.107).

### Note:

Since version 8.3 the sparse solvers renumber internally so the node renumbering parts of this command are no longer of importance. The **\*store\_nodes** command will however prevent an "on the fly" calculation of the interface nodes, perhaps saving some calculation time.

\*\*\*\*mesher \*\*\*mesh \*\*nset

\*\*nset

### **Description**:

This command creates a node set according to various input data. If the node set already exists, new nodes will simply be added. Nodes will be included in the set only once.

### Syntax:

```
**nset name
[ *axes axis1 axis2 axis3 ]
[ *function function ; ]
[ *limit epsilon ]
[ *nodes node1 ... nodeN ]
[ *nodes node1 ... nodeN ]
[ *plane n<sub>1</sub> n<sub>2</sub> n<sub>3</sub> intercept ]
[ *point [nearest] p<sub>1</sub> p<sub>2</sub> p<sub>3</sub> ]
[ *sequence start_id end_id [ increment ] ]
[ *surface ]
[ *type cartesian | cylindrical ]
[ *use_bset bset1 ... bsetN ]
[ *use_elset eset1 ... esetN ]
[ *use_nset nset1 ... nsetN ]
```

- \*axes switches the axes defining a cylindrical coordinate system when using the \*plane
  selection. an example is \*axes 1 3 2 where the cylinder is rotated about the 2 axis
  instead of the default 3-rd axis.
- \*function creates a nset using the nodes which satisfy the given function.
- \*limit epsilon specifies the precision for nodes to qualify to be accepted by a function or a plane function (because the values are never exactly the same due to numerical noise). The default value is  $10^{-3}$ .
- \*nodes node1 ... nodeN adds specific (existing) node numbers from the mesh to the nset
   (probably useful with \*\*add\_node). Non-existing nodes will be silently ignored.
- \*plane makes an nset given a plane equation (4 real values). The first three values are the components of the plane normal, and the fourth is the intercept (or its equivalent for cylindrical coordinates).
- \*point  $p_1 p_2 p_3$  adds existing node having the given coordinates *and* those within a radius of *epsilon*. Add the keyword **nearest** if you only want the nearest node to be added.
- \*sequence will add a sequence of nodes with ids from *start\_id* to *end\_id* with increment *increment* (Z-set 8.4 and newer). The latter can be negative, but in that case, *start\_id* should be greater than *end\_id*. If this is not the case, *-increment* will be used instead. If *increment* is omitted, a default value of 1 is used. Non-existing node ids generated with this command will be silently ignored.

- \*surface indicates that the set should be an outer bounding surface of the acceptable nodes (see also unshared\_faces page 2.121).
- \*type cartesian | cylindrical sets the type of coordinate system to be used with the \*plane option. Default is cartesian.
- \*use\_bset bset1 ... bsetN apply the nset command to only those nodes in the given boundary sets.
- \*use\_elset eset1 ... esetN apply the nset command to only those nodes in given element sets (was \*elset).
- \*use\_nset nset1 ... nsetN apply the nset command to only those nodes in the named node sets.

### Example:

Some example uses follow.

```
**nset ring1 *elset ring1 *plane 0. 1. 0. 4.
                                               % nodes in y=4
             *point 20. 0. 0.
**nset fix
                                               % single point
**nset r=20
                                               % the outer radius
 *limit 1.e-3
                                               % of a part at R=20
                                               % with some allowance for
 *type cylindrical
 *plane 1. 0. 0. 20.
                                               % numerical errors
%
\% These switch the cylindrical axes to be axis 2
%
                 *axes 1 3 2 *type cylindrical *plane 0. 1. 0. 20.
**nset t=10
**nset t=m10
                 *axes 1 3 2 *type cylindrical *plane 0. 1. 0. 0.
%
% Interpreted functions are perhaps the most useful .. see
%
**nset fix-func *function (z==0.0)*(y>=0.0);
```

\*\*\*mesher \*\*\*mesh \*\*nset\_intersection

### $**nset\_intersection$

### **Description**:

This command is used to create the intersection of nsets.

### Syntax:

The command has the following syntax:

\*\*nset\_intersection
\*nsets nset1-name nset2-name
\*intersection\_name resulting-nset-name

**\*nsets** is followed by the name of the nsets to intersect

\*intersection\_name is the name of the resulting nset

### Example:

```
****mesher
***mesh tube.geof
**open tube_tmp.geof
**bset skin % creates the outer skin of the tube
*surface
**nset_intersection % creates the outer skin of the given domain
*nsets zone1-volume skin
*intersection_name zone1-skin
```

\*\*\*\*return

## \*\*open\_bset

### **Description**:

This command takes as input a bset (faset in3D, liset in 2D), and duplicates nodes at the corresponding interface, to insert discontinuities (contact definitions, cracks ...) in the initial mesh. In the output mesh, the following items are available:

- 2 bsets named SIDEO and SIDE1, obtained by duplicating the input bset for each side of the created discontinuity. Note that bset elements are automatically ordered such that the normal is pointing outside (a property needed if those bset are to be used for contact definitions).
- 2 elsets named SIDE0 et SIDE1 with elements connected to the previous bsets.
- set of nodes located at the tip of the discontinuity, and ordered in continuous lisets named FRONTO, FRONT1 ... FRONTn, where n is problem-dependent.
- nset FRONT with all tip nodes in the prevous lisets.

Note that this command is only allowed for linear input meshes (commands \*\*quad\_to\_lin, \*\*lin\_to\_qaud may be used to bypass this limitation).

### Syntax:

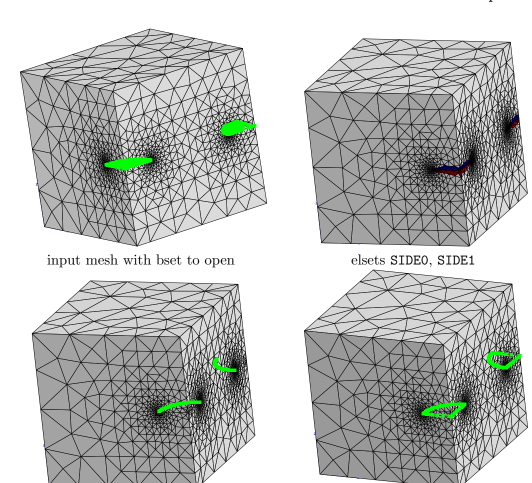
```
**open_bset
 *bset bname
[ *surface sname ]
[ *elset ename ]
[ *create_interface ]
```

- bname is the name of the bset, for which nodes need to be duplicated to create a discontinuity in the input mesh,
- the optional command **\*surface** takes as argument the name sname of an nset containing surface nodes. In this case surface nodes are removed from the front lisets (the front nodes ending points are preserved, however). See examples below.
- when the optional command \*create\_interface is specified, interface elements are automaticatly added between both sides of the discontinuity. Corresponding interface elements are included in the ouput INTERFACE elset.

# Example:

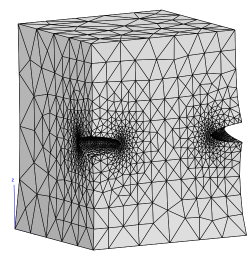
```
**open_bset
*surface SURFACE
*bset to_open
*branches
```

\*\*\*mesher
\*\*\*mesh
\*\*open\_bset



2 lisets created: FRONTO, FRONT1

FRONTS without the **\*surface** option



example FE results

# \*\*perturb\_inside

### **Description**:

This command is used to perturb a mesh geometry with a given uniform random law and a deformation factor (works only for 3D mesh if mesh surface must be preserved, e.g. without command move\_all).

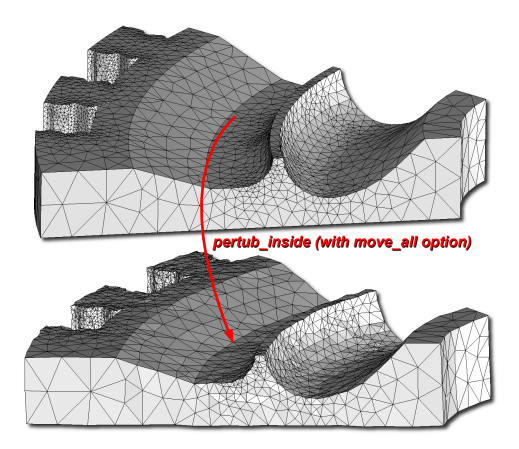
# Syntax:

The command has the following syntax:

\*\*pertub\_inside
\*factor moving-factor
[ \*move\_all ]

\*factor gives the moving factor distance.

\*move\_all impose the pertubation to be applied also on the mesh surface (works for meshes
in any dimension).



# \*\*porcupine

### **Description**:

This command builds a series of pyramids on an nset. This is generally used to in conjunction with boolean operations, in order to join quadrangular mesh faces to triangular ones. See e.g. the figure illustrating **\*\*regularize\_cfv**, on page 2.13, gives such an example (although pyramids where built differently in that particular situation).

## Syntax:

The command has the following syntax:

```
**porcupine
[ *bset bset-name ]
[ *height relative-height ]
```

- **\*bset** is the name of the bset on which pyramids are built. If it is not specified, pyramids will be built on the whole mesh skin.
- \*height is the height of the pyramid. It is a value relative to the perimeter of the base. The default value of  $\frac{\sqrt{2}}{2}$  is chosen so that a square-based pyramid has equilateral faces.

# Note:

```
****mesher
```

```
***mesh trench-for-diff.geof
**open trench.geof
**porcupine
*height .2
```

% It is then used in a boolean operation

```
***mesh difference.geof
**boolean_operation_ghs3d
*operation difference
*file1 earth.geof
*file2 trench-for-diff.geof
*output_file difference.gts
*options -m 100 -FEM % Memory size (MB)
*optim_style 1
*keep_2nd peau_commune
```

#### \*\*\*return

# \*\*project\_nset

### **Description**:

This command is used to project a given nset on a specified bset. It requires bset to be made of planar faces.

### Syntax:

The command has the following syntax:

**project_nset		
*nset		nset-to-project
*bset		$bset ext{-to-project-on}$
[ *distance	]	opening-value
[ *direction	]	vector
[ *orthogonal	]	vector

- \*distance insert a distance between projected points and given bset (can be used to separate two surfaces).
- \*direction gives the direction vector followed during the projection (usually projection is done orthogonal to the bset surface).
- **\*orthogonal** is used to specify a direction on which there must be no displacement of the nset during the process.

# \*\*propag\_crack

### **Description**:

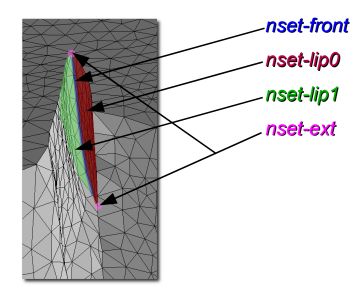
This command is used to grow 3D cracks described by an explicit mesh of the discontinuity. It verifies various conditions in order to preserve mesh's geometry.

### Syntax:

The command has the following syntax:

;	**propag_crack		
	*crack		front-nset-name lip0-nset-name lip1-nset-name ext-nset-name
Ε	<pre>*ask_speed_to</pre>	]	crack-component-name
Γ	*open	]	opening-value
Γ	*close	]	

- \*crack gives various nsets describing crack geometry: front nodes, first and second lips nodes. Last nset gives initial and final crack front nodes for opened cracks.
- **\*ask\_speed\_to** gives the name of the crack-front-component which impose the front advance.
- \*open is used to separate the lips with a given distance. This makes the remeshing process easier for the volume automatic mesher.
- **\*close** is used to collapse the lips (after remeshing if **open** command has been previously used).



# $^{**}quad\_to\_lin$

## **Description**:

This command reduces the integration order of a mesh to linear. This command can be conveniently applied using the Zquad\_to\_lin script front-end.

## Note:

There is also currently (Z-set 8.3 and newer) a mesher lin\_to\_quad (see page 2.71) to go from linear to quadratic. However, curved surfaces will be unrealistically modeled with facets. If there is the possibility of running with quadratic meshes, the basic meshing operations should be set to produce a quadratic mesh, and the mesh should be linearized before use.

### Syntax:

The command will be activated simply with the command name.

**quad_to_lin							
[ *param_file	list-of-files ]						

\*param\_file is an optional list of nodal parameter files, that will be linearized accordingly. Resulting files will be named with a .lin suffix.

## \*\*randomize

### **Description**:

This command superimposes a random displacement to nodal positions. Such a modification may be useful to model geometric imperfections.

# Syntax:

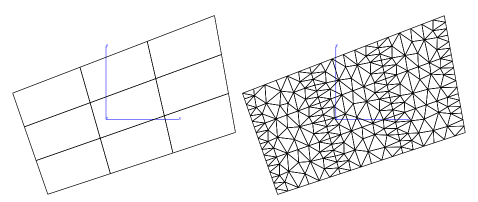
\*\*randomize
\*magnitude mag
[ \*nset nset1 ...nsetN ]
[ \*elset elset1 ...elsetN ]

- \*magnitude enter the distance (*real* value) which is the maximum magnitude of the displacement. The random displacement is applied with randomized direction in the order of space the node is given in.
- **\*nset** apply the random alteration to the given node sets.
- **\*elset** apply the random alteration to the nodes used in the elements contained in the given element sets.

# \*\*refine\_mesh\_based\_on\_element\_domains

### **Description**:

This command allows one to generate a master file (.mast) suitable to build a refined mesh. Each element of the initial mesh provides the vertices and the edges of the new mesh. A Delaunay domain is generated for each element of the initial mesh. The following figure shows an initial mesh and the new mesh that was built with the generated master file:



### Syntax:

```
**refine_mesh_based_on_element_domains
[ *edges no_edges ]
[ *master_file filename ]
```

\*edges no\_edges gives the number of nodes to use for each edge of the refined mesh (default 2).

\*master\_file filename is the filename for the master file of the refined mesh (default
refine.mast).

Note that this command currently (version 8.3) does not conserve the original elsets, and that it is implemented only for 2D linear elements.

If the **\*\*\*mesh do\_not\_save** option is omitted, a .geof file will be generated. However, this new .geof file is *identical* to the one that was loaded with the **\*\*open** command (if no other meshing operations have been carried out), and it does *not* contain the refined mesh.

### Example:

An example use follows:

```
***mesher
***mesh do_not_save
**open base.geof
**refine_mesh_based_on_element_domains
*edges 6
*master_file fine.mast
****return
```

\$

# \*\*regularize\_cfv

### **Description**:

This command is used to smooth crack front volume (see ) after remeshing process. It imposes a same distance between each node of the front and adapt the geometry of the volume meshed arround the front. It can also be used only to smooth an nset arround the front using projection on the crack front geometry. It the 3D crack propagation with remeshing process this command should be used after propag\_crack command and before any remeshing command (yams\_ghs3d).

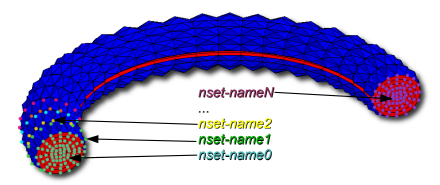
### Syntax:

The command has the following syntax:

\*\*regularize\_cfv
\*liset crack-front-liset
[ \*cut\_desc ] nset-name nset-number
[ \*nset ] nset-name

\*liset gives the crack front liset.

- \*cut\_desc describes the crack front volume, using such format: nset-name0, nset-name1
  ... nset-name(nset-number-1).
- **\*nset** gives the name on the nset on which regularization should be done (if no CFV is given).



# \*\*remove\_nodes\_from\_nset

### **Description**:

This command is used to remove a set of nodes from another given node set.

### Syntax:

The command has the following syntax:

\*\*remove\_nodes\_from\_nset
 \*nset\_name nset-name
 \*nsets\_to\_remove nset-list

The **\*nset\_name** is used to indicate the name of the source node set (the node set where the user wants to remove nodes). The **\*nsets\_to\_remove** command lists all node sets to be removed from the first node set.

# $**remove\_orphans$

# **Description**:

This command removes "orphaned" nodes from the mesh (nodes which are not attached to anything).

# Syntax:

There are no options or parameters for this command.

### $**remove\_set$

### **Description**:

This command is used to delete unwanted sets from the active mesh. In contrast to the **\*\*delete\_elset** command (page 2.45), these commands only delete the sets given, the actual mesh entities being pointed to are left as they are.

### Syntax:

The command has the following syntax:

\*\*remove\_set
\*nsets nset-list
\*bsets bset-list
\*elsets elset-list
\*ipsets ipset-list
\*nsets\_start\_with nset-prefix-list
\*bsets\_start\_with bset-prefix-list
\*ipsets\_start\_with elset-prefix-list
\*ipsets\_start\_with ipset-prefix-list
\*null\_sets

\*nsets every nset which name matches exactly with one the given names wil be removed.

- \*bsets idem for bsets.
- \*elsets idem for elsets.
- \*ipsets idem for ipsets.
- **\*nsets\_start\_with** every nset which name starts with one of the given prefixes will be removed.
- \*bsets\_start\_with idem for bsets.
- **\*elsets\_start\_with** idem for elsets.

\*ipsets\_start\_with idem for ipsets.

\*null\_sets all empty sets will be removed, i.e. sets not containing any items.

### \*\*rename\_set

### **Description**:

This command is used to rename sets from the active mesh.

(Available in Z8.4)

# Syntax:

The command has the following syntax:

```
**rename_set
[ *sets old-name1 new-name1 old-name2 new-name2 ... ]
[ *nsets old-name1 new-name1 ... ]
[ *bsets old-name1 new-name1 ... ]
[ *elsets old-name1 new-name1 ... ]
```

Any number of pairs of names can be given to each sub-command, and any combination of them is possible. As expected, the nsets command renames nsets, bsets command renames bsets and elsets command renames elsets. The sets command tries to rename all three kind of sets (and will simply ignore inexistent sets).

# Example:

```
**rename_set
*sets
  left hot_boundary % renames both the nset and the bset
  right cold_boundary
  top contact_zone
*nsets
  bottom fixed_border
  577 fixed_node % will actually create a new nset
```

# \*\*renumbering

### **Description**:

This command renumbers the mesh so that the global matrix front or bandwidth will be reduced. The algorithm is a modified Sloan renumbering scheme.

The renumbering by this command is not especially fast, and cannot directly handle disconnected mesh regions. In order to renumber across contact or MPC linked regions, use the **\*\*make\_springs** command (see page 2.72).

# <u>Note</u>:

It is advisable to do this operation as the last mesher command before saving the mesh. Alternately on unix platforms the Zrenum command can be used.

# Note:

The sparse matrix solvers should employ the metis renumbering described on page 2.76, or use built-in renumbering if available for the solver used.

# Syntax:

\*\*renumbering
[ \*frontal ]
[ \*nodal ]
[ \*w1 val ]
[ \*w2 val ]
[ \*subdomain ]

\*frontal Renumbers elements to reduce the front size in a frontal solution.

**\*nodal** Renumbers nodes to reduce the bandwidth size in a banded matrix solution (sky-line).

\*subdomain Renumbers according to subdomains for optimizing parallel computations.

# \*\*resize\_node

## **Description**:

This command is used to set the nodal position to have a desired dimension. Note that since version 8.0, the code supports mixed dimension meshes, and the dimension parameter in the .geof file is not used (left for compatibility reasons). If two coordinates are given in the node list, the node is a 2D node, and if three coordinates are given the node is a 3D node.

In Zmaster is is OK to have 3D nodes with 2D meshes, but in the FEA solver an error will be produced during the creation of DOFs and management of the shape functions.

### Syntax:

The command has the following syntax:

\*\*resize\_node
\*nsets nset1 ... nsetN
\*dim dim

The command **\*nsets** can optionally be used to specify that the dimension change only apply to those nodes. The **\*dim** command specifies the dimension to change to.

# \*\*rigid\_body

## **Description**:

Some direct solvers, like frontal and sparse\_direct, are able to automatically detect rigid body motions (i.e kernel) in the factorization step of the local matrices  $[K]_i$  (see page ??). But some others, like sparse\_dscpack, can not automatically detect these rigid body motions, so they have to be indicated to it.

Using this latter solver, the **\*\*rigid\_body** command can be used to find the rigid body motions of each sub-domain, using empirical geometrically criteria, and to write them in the *problem.cut* and *problem.cu* files. It should be noticed that this command just runs in 3D with impose\_nodal\_dof boundary conditions, in the global coordinate system.

### Syntax:

\*\*rigid\_body
\*nset nset1 ... nsetN
\*dof dof1 ... dofN

\*nset is the list of all nset names for which an impose\_nodal\_dof boundary condition
exists in the \*\*\*\*calcul module. If several dofs are imposed for an nset, the name of
this nset must appear several times.

\*dof  $% (\mathcal{A})$  is the list of the correspondent dofs.

# Note:

If the parallel computation is not a 3D one, or if other types of boundary conditions are used, the **\*\*rigid\_body** command should not be used; and the rigid body motions have to be directly specified at the end of the *problem.cut* file for each sub-domain using the following syntax. It should be noticed that in this case, the *problem.cu* has to be removed.

\*\*bc sd nrb node\_1 dof\_1 ... node\_nrb dof\_nrb

Where

sd is the number of the sub-domain for which rigid-body motions are specified.

**nbr** is the number of rigid-body motions for sub-domain *sd*.

**node\_i** is a node number.

dof\_i is the dof name for the rigid-body motion at node\_i (e.g. U1, U2 or U3).

The information concerning nodes and dofs can be obtained by observing sub-domains, or by running the same computation using a sparse\_direct linear solver and by stopping it just after sub-domain matrices factorizations.

# Note:

The sparse\_dscpack direct linear solver should not be used if the rigid body motions of a sub-domain change during computation.

\*\*\*mesher \*\*mesh \*rotate

# \*\*rotate

### **Description**:

This command is used to apply geometrical rotation of node coordinates within a mesh. The rotation will be specified using the same syntax as given for material rotations on page 3.137.

## Syntax:

\*\*rotate <ROTATION>
[ \*elset\_name name ]

### Example:

```
***mesh PIPE
 **open PIPE3
 **rotate
 *elset_name right_side
 x1 1. 0. 0.
 x3 0. 1. 0.
```

### Note: See also **\*\*symmetry**, page 2.111.

Z-set — Non-linear material & structure analysis suite

# \*\*scale

## **Description**:

This command scales the coordinate values for the mesh by a specified factor. This can be used to change units (e.g. inch-mm, meter-mm, etc).

### Syntax:

The syntax is simply the command name followed by a floating point scale factor:

### **\*\*scale** factor

An extended syntax allows different scaling factors in each direction:

**\*\*scale** xfactor yfactor zfactor

# $**set\_reduced$

# **Description**:

This command alters the degree of integration used for the element. It is a convenience feature to change quickly whether the elements are integrated fully or with reduced quadrature. With this, one does not need to alter the element type in every domain in Zmaster.

# Syntax:

\*\*set\_reduced
[ \*type reduced | normal ]
[ \*elset elset1 ... elsetN ]

**\*elset** select specific elsets to be modified. The default is to modify every element in the whole mesh.

\*type reduced make the elements reduced integration, irrespective of their current value.

\*type normal make the elements normal (full) integration, irrespective of their current value.

The default is to set to reduced integration. See also set\_normal (page 2.103).

# $**set_normal$

# **Description**:

This command is the inverse of **set\_reduced** (page 2.102): it forces normal (full) integration. It has the same syntax and sub-commands.

\*\*\*\*mesher
\*\*\*mesh
\*\*sequential\_ids

# $** sequential\_ids$

# **Description**:

This mesher renumbers a mesh sequentially from given start values. The default start node and element is  $one^{6}$ .

## Syntax:

The command has the following syntax:

\*\*sequential\_ids
\*node\_start st-node
\*element\_start st-node

 $<sup>^6\</sup>mathrm{start}$  numbers are available in 8.2+ only

# \*\*small

# **Description**:

Sets nodal coordinates with absolute values smaller than a preset value to zero<sup>7</sup>. This may be useful for instance for adding nsets with a criterion such as

\*\*nset nset\_name \*function (z==0);

which will miss points that have very small non-zero z-coordinates due to numerical noise. This command acts on all components of the position vector of the given nodes.

# Syntax:

\*\*small
[ \*limit limiting\_value ]
[ \*nset nset\_name ]

- \*limit enter the limiting value (*real* value) below which the coordinate value will be set to zero. Negative values will have their minus chopped off. The default is 1.e-6.
- **\*nset** apply to the given node set. The default is ALL\_NODE.

 $<sup>^{7}</sup>$ Available in 8.2+ only

## $** sort\_nset$

#### **Description**:

This command sorts a node set in increasing order with respect to the value of a user-specified function of the coordinates of the nodes.

#### Syntax:

The command has the following syntax:

\*\*sort\_nset
 \*nset\_name nset-to-sort
 \*criterion function ;
[ \*n2\_sort ]

\*nset\_name specifies the node set to sort. There is no default value.

- \*criterion enter a function of the coordinates in order to sort by. It is a function of two points  $P_1(x_1, y_1, z_1)$  and  $P_2(x_2, y_2, z_2)$  within the nset (without the  $z_1$  and  $z_2$  for two-dimensional meshes). It may also compare node ids (the corresponding variables are named id1 and id2). The function must return whether  $P_1$  is "greater" than  $P_2$ : criterion = -1 means  $P_1 < P_2$  (no action taken), criterion = 0 means  $P_1 = P_2$  (no action taken) and criterion = +1 means  $P_1 > P_2$  (the order of the points will be changed). Do not forget the semicolon at the end. There is no default value.
- \*n2\_sort uses a brute-force  $O(n^2)$  sorting method, instead of the  $O(n \log n)$  quicksort, which is used by default. This option is included because the quicksort routine is broken on some systems.

#### Example:

```
****mesher
***mesh out.geof
**open in_geof % mesh containing unsorted nsets left and radial
**sort_nset
*nset_name left
*criterion x1<x2; % sort nset according to increasing x. Note the ;
**sort_nset
*nset_name radial
*criterion (x1*x1+y1*y1+z1*z1)>(x2*x2+y2*y2+z2*z2); % sort nset
% according to decreasing distance from the
% origin [0. 0. 0.]. This generates an error
% message for two-dimensional meshes.
****return
```

# \*\*split

# **Description**:

This is an interface for the ONERA splitmesh program which can be used to build the subdomain information for a parallel computation. There is a different interface for 8.0 and 8.2. 8.0 requires that the splitmesh executable be installed on your system (not supplied by default), while it is integrated into 8.2.

# Syntax:

For 8.0 the command has the following syntax:

```
**split
*splitmesh_location path
*domains num
*mincon int-value
*no_elset
```

\*splitmesh\_location specify the path to the splitmesh executable program.

\*domains enter the number of domains (separate processes) in a parallel computation.

\*no\_elset don't generate the element sets for domain visualization.

The syntax for 8.2 is the same, except that the **\*splitmesh\_location** is not required (or available).

# Example:

An example use follows from the test Parallel\_test/INP/arm2.inp

```
****mesher
    ***split
    **splitmesh_location /home/saturne/feyel/bin/splitmesh
    **domains 4
    ***renumbering
    **subdomain
    **subdomain
    ***geof_format
    **unformat
****return
```

\*\*sweep

## **Description**:

This command extends planar (2D or 3D shell/face) geometries into 3D Through a sweep rotation. One may give an arbitrary rotation axis, point to make quite arbitrary sweeps. The method works for linear/quadratic meshes in the positive or negative rotation directions. Degenerate cases are handled (inside edge is on the axis, so the inner elements become pyramids, etc).

Extension may be given with different start/end meshes. In this case, the first mesh is swept forward, while the second swept back. The mid point nodes are the interpolation between the two.

## Note:

please be careful concerning the order of elements between the two meshes (domains) in this case. The topology must be equivalent.

If no elset is given the whole mesh will be used.

# Syntax:

**sweep							
[	*elset	eset	1	]			
Ε	*elset2	eset	2	]			
[	*new_els	set	ne	w-e	lset	]	
Γ	*angle	deg	ree	s-re	otati	on	]
[	*num		nu	m	]		
Γ	*prog		j	proį	gress	sion	]
[	*axis		(	dir-	vec	]	
[	*center		(	cent	t-vec	; ]	
[	*fusion		(	dist	]		

- \*elset eset1 defines the element set or faset which will be extended. There is no limitation on the type of elements or faces contained on the face. Mixed order is not allowed however. The default is all elements.
- \*elset2 eset2 This optional command makes the extension between two mesh/face sets as given. Note that the mesh topology must be exactly the same. This command is most useful with the mapped mesh domains available in Zmaster. The degenerate case of 2 edges being the same (so the extended mesh forms a wedge) is handled. Do not use \*distance with this option.
- **\*new\_elset** *new-elset* This optional command gives the elset name for the created elements instead of being the same as the input name.
- \*angle The angle to sweep through in degrees. May be positive or negative. The default is 360 degrees.
- \*axis Use this command to specify the rotation axis in vector form. The default is
   (0. 1. 0.).

\*\*\*\*mesher \*\*\*mesh \*\*sweep

- \*center Use this command to specify the rotation center in vector form. The default is
   (0. 0. 0.).
- **\*fusion** specifies the distance below which 2 nodes are considered identical. Default value is that of the global parameter MeshFusion.

\*\*\*\*mesher \*\*\*mesh \*\*switch\_element

# \*\*switch\_element

#### **Description**:

This command is used to inverse the element axis orientation. Note that the element orientation can be visualised in Zmaster.

#### Syntax:

The command has the following syntax:

\*\*switch\_element
\*elset elset-name
\*axis axis

The options for this command are described below:

\*elset is the elset to which the inversion is applied

\*axis Give the axis aorund which the rotation is performed

# \*\*symmetry

# **Description**:

This command is used to do a symmetry of a nset. Its modifies nodal positions by a symmetry's transformation (central, axial or planar), described with a point and a normal. The type *line* is only valid for 2D meshes (input vectors should have 0. as z-coordinate), and the type *plane* is only valid for 3D meshes.

## Syntax:

The **\*\*symmetry** command takes the following syntax:

\*\*symmetry
\*type [point | line | plane]
\*point origin-point
\*normal normal-vector
[ \*nset nset ]

\*type Defines the type of symmetry's axe : a point, a line or a plane.

\*nset The nset whose you want to transform (default value: ALL\_NODE).

\*point This defines the origin for the type of symmetry.

**\*normal** Normal's direction for the projection of the symmetry.

# Example:

The following example shows the use of symmetry of the x - z-plane for all points:

```
**symmetry
*type plane
*nset ALL_NODE
*point (0.,0.,0.)
*normal (0.,1.,0.)
```

# \*\*thicken\_bset

## **Description**:

This command is used to add a layer of elements on a specified bset. This is sometimes necessary before using a boolean operation (page 2.13).

# Syntax:

The command has the following syntax:

**thicken_bset					
*bset	bset-name				
*height	height-of-elements				
[ *direction	vector ]				
[ *towards	point ]				

**\*bset** elements are attached to the specified bset. There is currently no default.

\*height specifies the height of the newly built elements (default is 1).

 $\ast \texttt{direction}$  specifies a direction for the extension.

**\*towards** is an alternative to **\*direction**: elements are built towards the specified point (or away from, if height is negative).

# Example:

```
***mesher
***mesh trench.geof
**open trench_for_difference.geof
[...]
**thicken_bset
*bset TheSurface
*height -30.
*towards (0. 0. 0.)
****return
```

\*\*\*\*mesher \*\*\*mesh \*\*to\_2d

\*\*to\_2d

# **Description**:

This command transform 3d meshes into 2d meshes. This command only modify the dimension of node location vector.

# Syntax:

The command has the following syntax:

\*\*to\_2d

\*\*\*\*mesher \*\*\*mesh \*\*to\_3d

\*\*to\_3d

# **Description**:

This command transform 2d meshes into 3d meshes. This command only modify the dimension of node location vector.

# Syntax:

The command has the following syntax:

\*\*to\_3d

\*\*\*mesher \*\*mesh \*\*to\_cax

\*\*to\_cax

# **Description**:

This command transform 2d plane meshes into axisymmetric meshes.

## Syntax:

The command has the following syntax:

\*\*to\_cax

In pratice, this command symply changes the element types, from c2d to cax.

# \*\*transform\_fili

### **Description**:

This command is used to correct a mesh previously imported from Abaqus and generated by the SAFRAN FILI tool. It replaces degenerated elements with Z-set standard ones (prism and pyramid elements).

In some cases this mesher may fail on quadratic meshes, so it is recommended to work on linear meshes only (see the lin\_to\_quad mesher on page 2.71).

# Syntax:

The command has the following syntax:

```
**transform_fili
[ *elset elset-name ]
```

**\*elset** specifies the elset where the operation is applied (default is to correct the whole mesh).

# \*\*translate

#### **Description**:

This command is used to apply a rigid body translation of the mesh.

## Syntax:

The syntax consists of the command name followed by the 3 translation components:

\*\*translate dx dy dz

The command also has an extended syntax that provides additional functionalities:

\*\*translate
 \*x dx
 \*y dy
 \*z dz
 \*nset nset-name
 \*nodes node1 ... nodeN
 \*elset elset-name

\*x, \*y, \*z specifies the translation

\*nset restricts the translation to the given nset

 ${\tt *nodes}$  restricts the translation to the given node ids

 $\texttt{*elset} \quad \text{restricts the translation to the given elset}$ 

# Example:

\*\*translate 0. -5.5 -1.0

\*\*\*mesher
\*\*\*mesh
\*\*unconnected\_parts

# $** unconnected\_parts$

#### **Description**:

This command creates one elset per disconnected part of an input mesh: a new list of elsets is created (part1, ..., partN), with N the number of disconnected parts of the input mesh.

# Syntax:

\*\*unconnected\_parts

The command has no options.

#### Example:

An example is shown below:

```
****mesher
 ***mesh
 **open sphere_cylinder.geof
 **unconnected_parts
****return
```

\*\*\*\*mesher \*\*\*mesh \*\*union

## \*\*union

#### **Description**:

This command creates the union of meshes.

#### Syntax:

\*\*union
 \*add mesh-name
[ \*elset eset-name ]
[ \*merge\_nset merge ]
[ \*set\_change\_name chg-name ] % 8.2 (obsolete)
[ \*set\_base\_name base-name ] % 8.2 w(obsolete)
[ \*base\_name base-name ]
[ \*tolerance val ]
[ \*translate\_new\_meshes ] % 8.2 (obsolete)
[ \*tr trans-vector ] % 8.2 (obsolete)
[ \*translate trans-vector ]

\*add mesh-name adds the mesh in file mesh-name.geof to the active mesh.

- **\*elset** eset-name A new element set will be created with the name eset-name and containing all the elements added in this operation.
- \*merge\_nset limit the fusion of nodes to the given node set. The node set is on the current mesh, not the newly added mesh given by mesh-name.
- \*base\_name rename all sets from old-name to base-name.old-name.
- \*tolerance Gives the critical distance for node fusion between the two meshes. If too large, elements will collapse because node neighbors are joined, too small and there may be gaps at the interface. Use zero to join two meshes without node fusion, such as in the joining of two parts in contact. The default variable is in the global parameters, probably 1.e-3 (in the dimension of the mesh).

#### Example:

A short example follows of some union sections from the test case in Mesher\_test/INP/PIPE.inp

```
**union
 *add pipe-data/PIPE1.geof
 *elset PIPE1
 *tolerance 0.15
**union
 *add pipe-data/PIPE2.geof
 *elset PIPE2
 *tolerance 0.15
```

# \*\*unshared\_edges

## **Description**:

This command makes an edge boundary set (liset) for the outer edges of a mesh. It operates similarly to the **\*\*unshared\_faces** for 2D meshes, and produces an edge output.

# Syntax:

\*\*unshared\_edges bset-name
\*elset elset1 ... elsetN

\*elset used to specify a number of elements sets to find the unshared faces rather than the whole mesh. Note this still gives the faces which are not shared with any other element, whether in the listed elsets or not. It is just that the faces checked are a sub-set of the total mesh, and the resulting set will be on the named elsets only.

# \*\*unshared\_faces

# **Description**:

This command makes a boundary set (which can be converted to an nset using **\*\*nset**) from the outer boundary of a given element set, or of the whole mesh.

# Syntax:

\*\*unshared\_faces bset-name
[ \*elsets elset1 ... elsetN ]

\*elsets used to specify a number of elements sets to find the unshared faces rather than the whole mesh. Note this still gives the faces which are not shared with any other element, whether in the listed elsets or not. It is just that the faces checked are a sub-set of the total mesh, and the resulting set will be on the named elsets only.

## <u>Note</u>:

This command is currently faster than the **\*\*bset** bset-name **\*surface** equivalent.

## Note:

This command replaces the deprecated and ambiguous **\*\*unshared** command.

# \*\*volume\_to\_shell

#### **Description**:

Starting with a 3D volume mesh, this command extracts the given best and creates the corresponding surfacic 3D-shell mesh.

#### Syntax:

The command has the following syntax:

\*\*volume\_to\_shell
[ \*bset bset-name ]

**\*bset** specifies which bset to extract. The default is to extract the whole mesh skin.

# \*\*yams\_ghs3d

## **Description**:

This command can be used for remeshing an input Zébulon mesh by means of the DIS-TENE remeshing tools. There are basically 3 different pieces of software that may be used to build the new mesh:

- i) Yams is used for remeshing a surface mesh (that may be built from the skin of a 3D mesh or directly from a shell mesh) according to various criterions,
- ii) Ghs3d is a general purpose 3D mesher that takes as input a surface mesh and fills the volume with the trahedras,
- iii) given a metric map, Meshadapt performs either surface, volume, or surface+volume adaptation at the same time. The metric map is made of the desired sizes of vertices (edges) connected to each vertex (node) of the mesh. Since this map is given on the initial mesh, several iterations may be needed in the adaptation process.

Those tools are fully interfaced for Zébulon and the yams\_ghs3d command automatically writes out DISTENE input and reads in DISTENE output files to build the Zébulon mesh. DISTENE binaries are included in the standard Zébulon distribution (in the \$Z7PATH/PUBLIC/lib-\$Z7MACHINE/Zmesh/ folder), but an optional license key is needed to run those softwares. Yams, Ghs3d, and Meshadapt manuals are also available in the Zmesh/ folder, and the user may have a look at those if a fine tuning of the remeshing process is needed.

Note that only linear elements are handled by yams\_ghs3d. A combination of the quad\_to\_lin (before yams\_ghs3d) and lin\_to\_quad commands (after) may then be used to allow the use of quadratic meshes. Warning: if a computation result is available, the mesh will be deformed before remeshing.

#### Syntax:

The command has the following syntax:

```
**yams_ghs3d
 [ *yams_only ]
 [ *nb_iter_surf niter-surf ]
 [ *nb_iter_vol niter-vol ]
 [ *nb_iter niter ]
 [ *force_meshadapt ]
 [ *refinement_origin origin-nset ]
 [ *refinement func(x,y,z); ]
 [ *refinement_file fname ]
 [ *absolu ]
 [ *min_size min
                  ٦
 [ *max_size max ]
 [ *gradation grad ]
 [ *tolerance tol ]
 [ *optim_style opt
                     ]
 [ *preserve_faset bset-name1 bset-name2 ...
                                             ]
```

[ \*preserve\_liset bset-name1 bset-name2... ]
[ \*yams\_options yams-options ]
[ \*ghs3d\_options ghs3d-options ]
[ \*meshadapt\_options meshadapt-options ]

\*yams\_only forces surface remeshing only (ie. performing only step i)).

- \*nb\_iter\_surf if a metric map is given, this command allows to specify the number nitersurf of iterations involved in step i) to satisfy a given metric map for the surface mesh (default is niter-surf=3).
- \*nb\_iter\_vol if a metric map is given, this command allows to specify the number niter-vol of iterations involved in step iii) to satisfy a given metric for the volume mesh (default is niter-vol=2),
- \*nb\_iter this command can be used to define both *niter-surf* and *niter-vol* at the same time (*niter-surf=niter-vol=niter* in this case).
- \*force\_meshadapt when a metric map is given as input to drive the remeshing process, this option has the effect to force the use of Meshadapt instead of Yams during step i). Note that in this case Meshadapt is run only 1 time during step i) (option -O1 for surface adaptation) and nb\_iter times during step ii) (option -O3 for surface+volume adaptation),
- \*refinement\_origin this command is used to define the name origin-nset of an nset taken as a base to calculate the metric map according to function func(x,y,z) (see next command),
- \*refinement this command defines the function func(x,y,z) used to compute the metric map (see explanations in the Metric map calculation paragraph). Note the only variables allowed in this function are x, y or z and that the definition should end with a ";" character.
- \*refinement\_file with *fname* the name of a metric map file (the file syntax is explained in the Metric map calculation paragraph).
- \*absolu is a Yams option and is only used during step i). In this case edge sizes (arguments of the \*min\_size or \*max\_size commands, and/or values given in the metric map) are *absolute* values. The default alternative is to use *relative* values (ie. sizes are scaled by size of the bounding box of the initial mesh, see the Yams user manual).
- \*gradation grad is a Yams only option (step i) that may be used to control the element size variation (please refer to the Yams user manual).
- \*tolerance tol is a Yams only option (step i) used to set the value tol of the maximum chordal deviation tolerance (please refer to the Yams user manual).
- \*optim\_style opt is a Yams only option (step i) and can be used to specify the type of optimization style (coearsening, enrichment etc... please refer to the Yams user manual). Default is opt=2 for geometrical mesh enrichment.

- \*preserve\_XXset used to preserve set(s) during surface remeshing. This can only be applied to set(s) located on the mesh skin and it does not mean that the *exact* set(s) topology should be kept (set(s) is/are rebuilt after the remeshing process: position and element type could have changed).
- \*yams\_options , \*ghs3d\_options and \*meshadapt\_options commands are meant for advanced DISTENE tools users, and allow to specify manually the DISTENE softwares command line options (see user manuals in the Zmesh/ folder). Default values are in general appropriate for typical Zébulon applications.

#### ${\bf C}{\it alculation}$ of the metric map

The metric map is defined by a set  $\mathcal{P}$  of base points with the target element sizes near those points:

 $\mathcal{P} = \{ P_i(x_i, y_i, z_i), d_i ; i = 1, ..., n_b \}$ 

where  $n_b$  if the number of base points,  $P_i$  a particular point in  $\mathcal{P}$  with coordinates  $(x_i, y_i, z_i)$ and a prescribed size of  $d_i$ .

 $\mathcal{P}$  may be defined either by commands \*refinement\_origin or \*refinement\_file:

\*refinement\_origin origin-nset fills  $\mathcal{P}$  with points corresponding to nodes in nset origin-nset,

\*refinement\_file fname defines the name of a file containing  $P_i$ ,  $d_i$ . Syntax of this ASCII file is the following one:

- first line:  $n_b$  (number of base points: 1 integer)
- line 2 to  $n_b + 1$

 $x_i \ y_i \ z_i \ d_i$  (definition of  $P_i$  coordinates and prescribed size: 4 double values)

Then for each node N in the current mesh, the prescribed size d(N) of element edges built from node N is calculated in the following way:

- first, find closest point  $P^*$  from N in set  $\mathcal{P}$ :  $P^*$  such that  $distance(P^*, N) = Min \{ distance(P_i, N) , i = 1, ..., n_b \}$
- then, if a function func(x,y,z) is given by means of command \*refinement:

$$d(N) = func(x^*, y^*, z^*)$$

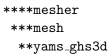
with  $(x^*, y^*, z^*)$  the coordinates of point  $P^*$ ,

• else:

$$d(N) = d^*$$

with  $d^*$  the prescribed size given for point  $P^*$  in  $\mathcal{P}$ .

- finally, if either commands \*min\_size min or max\_size max are specified:
  - if  $d(N) < \min$  then  $d(N) = \min$
  - if d(N) > max then d(N) = max



Note that the option \*absolu is advised in this case to insure that d(N) values calculated do correspond to actual edge sizes in the finite element mesh.

#### Example:

```
See tests in folder $Z7PATH/TESTS/Distene_test/INP/.
```

#### \*\*\*\*mesher

```
***mesh output
**open input.geof
**yams_ghs3d
*force_meshadapt
*nb_iter 3
*absolu
*min_size 0.03
*max_size 0.2
*preserve_faset haut-ext bas-ext U1=0 U3=0
*refinement_origin bas-ext
*refinement x*x+y*y+z*z;
****return
```

# \*\*yams\_by\_elset

# **Description**:

This command is derived from **\*\*yams\_ghs3d** mesher, and can be seen as a "mesher modifier". It allow to preserve elsets topology during **yams\_ghs3d** remeshing process. This is achieved by internally providing a "color" for each named elset. The colored areas borders topology are then "protected".

#### Syntax:

The command has the following syntax:

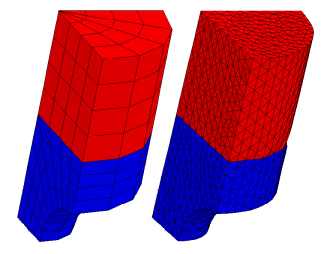
```
**yams_by_elset
[ *elset elset-name1 elset-name2 ... ]
[ current "**yams_ghs3d" options]
```

**\*elset** names of elsets to be preserved.

#### Example:

See tests in folder \$Z7PATH/TESTS/Distene\_test/INP/.

```
***mesher
***mesh cyl4
**open cyl.geof.ref
**yams_by_elset
*elset center attach
%%% yams_ghs3d options
*force_meshadapt
*nb_iter_surf 1
*nb_iter 3
*absolu
*min_size 0.1
*max_size 0.1
*refinement 0.1;
****return
```



# \*\*refine\_elset

#### **Description**:

This command is derived from **\*\*yams\_ghs3d** mesher, and can be seen as a "mesher modifier". it allow to remesh a single elset using **yams\_ghs3d** remeshing process, without modifying neighboring element. This is achieved by extracting the target elset in a separate mesh and imposing the conservation of the interface ( interface : the skin shared between the elset and the remaining of the mesh).

#### Syntax:

The command has the following syntax:

\*\*yams\_by\_elset
[ \*elset elset-name ]
[ \*material\_elset material\_elset\_name ]
[ current "\*\*yams\_ghs3d" options]

**\*elset** name of elset to be remeshed.

- \*material\_elset this command can be used for multi-material models. It will add all elements created by remeshing elset "elset\_name" to elset "material\_elset\_name".
- \*maxsize (inherited from \*\*yams\_ghs3d) note that by default the element max size is automatically calculated as the maximum size of element at the interface of remeshed elset note that

#### Example:

See tests in folder \$Z7PATH/TESTS/Distene\_test/INP/.

```
****mesher
***mesher
***mesh cyl3
**open cyl.geof.ref
**refine_elset
*elset center
*material_elset ALL_ELEMENT
%%% yams_ghs3d options
*nb_iter 3
*absolu
*min_size 0.03
*preserve_faset haut-ext bas-ext U1=0 U3=0
*refinement_origin bas-ext
*refinement y*y;
****return
```

#### \*\*remesh\_from\_results

#### **Description**:

This mesher allow to build a metric map (for Disten tools) based on the results of a previous computation. The metric map definition can be found in **\*\*yams\_ghs3d** section or in the Disten manuals. It can be useful to refine a mesh according to the gradient of given field or an a posteriori error estimator.

In this mesher, the metric map can be partially computed from results over a defined elset, the remaining of the map (corresponding to elset complement to the mesh) is simply filled with the actual mesh metric. We can also in this case choose a conservative remeshing based on refine\_elset (outside elset will remain unchanged) or a less conservative remeshing based on yams\_ghs3d (outside elset will change but keeping the same characteristic element length).

#### Syntax:

The command has the following syntax:

```
**remesh_from_results
[ *result_name result-file-name ]
[ *Z8 ]
[ *var var-name ]
[ *card card-number ]
[ *power power-coefficient ]
[ *free_interface ]
[ current "**refine_elset" options]
```

\*result\_name results file name

- \*Z8 add it to specify that results are in Z8 format (.zres folder). if omitted Z7 format is assumed
- \*var name of the variable in the results database to be used for metric computation.
- \*card index of the results card to be used for metric computation.

\*power modify the value of the variable var such that  $var = var^{power}$ 

\*free\_interface if set, activate the "less conservative remeshing" otherwise the
 refine\_elset strategy is used

#### Example:

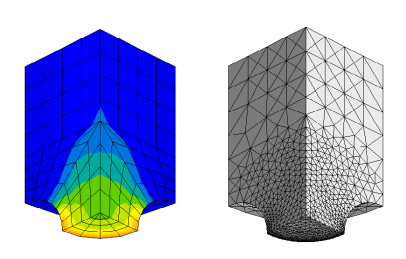
See tests in folder \$Z7PATH/TESTS/Distene\_test/INP/.

```
***mesher
***mesh cylr4
**remesh_from_results
```

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\*\*\*mesher
\*\*\*mesh
\*\*remesh\_from\_results

```
*result_name CYL/cyl
*var sig22
*card 2
*power 0.5
*free_interface
%%% refine_elset/yams_ghs3d options
*elset center
*absolu
*min_size 0.01
*max_size 0.15
*nb_iter 3
*preserve_faset haut-ext bas-ext U1=0 U3=0
****return
```



# Chapter 3

# Finite Element (.inp file)

# Introduction

#### Description:

This chapter concerns the definition of finite element problem solutions with the Zebulon solver, and is indicated under the main command **\*\*\*\*calcul**. The FEM problem definition will be given by a primary file with name *problem.inp*, and also several auxiliary files, as required. The prefix *problem* will be used in other input and output file names, and is henceforth called the problem name. Generally, a geometry specification file is required with the default name *problem.geof*, where the nodes, elements, and other geometrical information is given. It should be noted that the geometry file does *not* describe the type of element (formulation), which will be defined in the .inp file. Frequently a separate material file is given as well, but this information may be added to the .inp file just as easily.

The FEM calculation mode of Z-set reads and interprets all the non-commented instructions in the main .inp file between the commands **\*\*\*\*calcul** and **\*\*\*\*return**. Other problem modes may therefore be, in the same file, demarcated by commands starting with four asterisks.

In the file .inp we may give references to the names of other files for the material files, geometry files, or other data files such as external temperature fields, initial values for the material variables, etc. These files will be discussed throughout the command summary here, and in the section on file formats (page 5.10).

An overview of the functions to which the .inp file serves is given below:

- Define the nature of the calculation: mechanical, thermal steady-state, thermal transient, etc.
- To give complementary information of the geometry or formulation of the calculation. For example, if a 2D problem is in plane stress or plane strain conditions, in finite strain formulations, etc.
- Give the boundary conditions. These include specification of the degrees of freedom and the associated forces (e.g. displacement and point forces), as well as calculated conditions such as pressure or centrifugal forces. With the exception of eigen frequency analysis, the boundary conditions must all be prescribed in the time scale of the calculation. In the case of a static mechanical calculation, the time may not have a physical significance, but is rather used as a fictitious measure of the loading processes. For example, in order to calculate a static structure in three loading cases, we can write the boundary condition evolutions as a function of dimensionless times 1, 2, and 3.
- Assign groups of elements (element sets or *elset*) a material file specifying the material law and its coefficients. Each material file is also assigned an integration method if necessary. The separation of the finite element model and numerical integration allows one to work with a small base of material files for many calculations with different numerical methods.
- Define the method(s) of global solution, and define the solution steps which must be made in a non-linear calculation. This data will include the algorithm used (Newton-

Raphson, BFGS,...) the convergence criteria, and automatic time stepping methods. The time used here defines the time measure used for all other parts of the problem definition such as the boundary conditions.

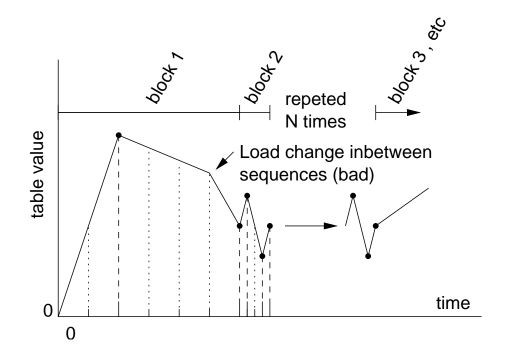
- To define tables as a function of time. Specification of numerous parameters in the calculation such as the boundary condition amplitudes will be defined by these tables.
- Specify output for whole-model fields, and directly computed curves which are generated during the calculation. Multiple output "blocks" may be specified, with different frequencies of output, etc.

## Linear solution:

Linear solutions typically involve a number of "loading cases" which are normally superimposed to a structures response to many different external solicitations. There is now a special key given in the output section **\*\*linear\_solution** (see page 3.149).

#### Time stepping:

The success of a nonlinear calculation depends on definition of appropriate loading sequences. The following figure summarizes the basic elements of loading sequences:



This example shows the waveform desired for a particular parameter in the calculation (such as a loading level). On the x-axis is the time scale of the calculation, broken into segments (long dashed lines) and increments (dotted lines). The y-axis represents the loading level described by a hypothetical table. The sequences are put into groups any of which can be repeated to give cyclic loadings.

The function of sequences is to demarcate segments of a particular solution strategy in the linear progression of time from  $t_1$  to  $t_2$  with  $t_2 > t_1$ . During the sequence the solution parameters such as convergence criteria, method of stiffness matrix resolution, etc are fixed.

Increments which divide the sequence are used to refine the incremental solution in a linear manner (normally through steady changes in loading). The increment in time during each loading increment is therefore  $\Delta t = \frac{t^2 - t^1}{n}$  for *n* increments. For nonlinear solutions, within each increment convergence may require several trial solutions which will be called *iterations*.

An alternative to the linear division is to give progressively increasing or decreasing steps, or automatic calculation of the increments. This later allows stepping as a function of the problem variables for accuracy control, as well as in function to the convergence rate, etc for divergence control.

As the loading magnitudes are given in terms of time, and not directly in terms of the segment numbers, incompatibility in the load waveforms with the sequences is available. This is schematically shown in the above figure within the second load sequence. Here the load changes at the end of the third increment of the second sequence, which is one increment away from the sequence end. This type of loading is **not** advised, and should be cautiously verified in the case of complex loadings (e.g. cyclic or otherwise lengthy).

With these terms defined, we may progress with the command summary for the FEM problem loading. To clarify their meaning however, it may be very useful to follow several of

the example problems given in the example section of the manual.

# Specifying node sets:

Node sets are defined in the .geof file, which is created by meshing operations. In commands which require a **nset**, give the character name for the node set desired, as appears in the .geof file. An additional shortcut exists for the case of specifying *all* the nodes in the mesh. In this case, the keyword ALL\_NODE may be used for the node set name.

# Specifying element sets:

Element sets are defined in the .geof file as well, defining lists of element numbers which may be identified together. These sets will be used to assign particular element formulations to regions in the mesh, material behaviors, boundary conditions, etc. As for node sets, element sets are identified by their full name given in the .geof file. The entirety of elements in a mesh may be identified with the shortcut keyword ALL\_ELEMENT to be used in the place of a user-defined set name.

## Specifying magnitudes:

The magnitudes of commands (particularly boundary conditions) will be controlled using either tabular or functional values. The definition of tables and functions are treated as a major element of the user input file definition (at the **\*\*\*** level). The commands to assign tables is **\*\*\*table** and functions **\*\*\*function**, each of which will create an object or objects with a name attribute. Giving names to the functions / tables provides a means for access in other objects which will use them to calculate magnitudes in time. The program structure is such that tables or functions may be referenced before their definition, so the **\*\*\*table** and **\*\*\*function** commands may appear anywhere in the valid three asterisk command level under **\*\*\*\*calcul**.

The code now accepts pre-defined magnitude names. The most useful of these<sup>1</sup> is time which provides a magnitude equal to the current time.

<sup>&</sup>lt;sup>1</sup> in fact, the only current one

# Specifying command parameters:

Many of the commands (especially boundary conditions) take particular parameters to define the actual command applied. These may be character string names, integers, floating point values, vectors, etc. The program is rather strict about typing these data, in order to ensure that what is read is indeed what was intended. The following summary is thus provided to define the syntax data types:

- character values character values refer to alphanumeric character strings, which may include some symbol characters such as = +, but not delimiting characters such as spaces or commas. The string is terminated at the next newline, space, tab, comma, or comment character % or #. Character values may be up to 255 characters long, but will never be shortened.
- *integer values* these are numeric fields separated by the standard delimiters (as in character strings), and must only be composed of the digits 0-9 with no decimal points. Decimal values found for integers will produce an error message.
- real values real values are used to specify decimal numbers. They may be positive, or negative, and *must* include a decimal point. Exponential notation is allowed. Some examples are 1.29 5.e+6.1 0.314159E-1.
- vector values vectors define either a point in 2D or 3D space, or the components of a vector direction in space. The size of the vector should be coherent with the spatial dimension of the problem. Vectors are always surrounded by parenthesis () and take only real data for their components. example: (1...5..5).

# \*\*\*\*calcul

#### **Description**:

This command marks the beginning of a FEM calculation definition. The Z-set program in FEM mode will search this command and interpret all the sub-commands until the termination token **\*\*\*return** is reached. A keyword following the **\*\*\*calcul** token will indicate the type of calculation which is to be made.

#### Syntax:

The calculation will be defined using the following syntax:

****calcul	type			
options				
****return				

The calculation *types* which are possible are listed below:

CODE	DESCRIPTION
mechanical	static mechanical (no inertial effects)
eigen	eigen frequency analysis
dynamic	mechanical with inertial effects (implicit)
explicit_mechanical	explicit solver for structural dynamics
$thermal\_steady\_state$	stationary thermal calculation
$thermal_transient$	transient thermal calculation
diffusion	Ficks Law diffusion analysis with multiple phases
weak_coupling	Generalized coupled analysis

In the absence of a type, the default will be mechanical.

The basic components of the allowable input data after \*\*\*\*calcul are summarized below<sup>2</sup>:

 $<sup>^2\</sup>mathrm{More}$  options may exist. Later versions of the code output all available command names with the  $-\mathrm{H}$  switch.

CODE	DESCRIPTION
***mesh	used to specify the types of elements in a mesh, or give an alternate geometry file name
***restart	requests that the calculation be continued from a previous stored result
***resolution	used to declare the solution procedures including the load- ing sequences
***equation	declare relationships between entities (multi-point con- straints) within the calculation
***impose_kinematic	imposes a geometrical evolution for problems which do not have displacement variables but do have an integration vol- ume which changes (thermal, diffusion).
***sub_problem	used to define a sub-problem in the sequential weak cou- pling algorithm; may be post calculations or re-meshing operations as well.
***parameter	allows specification of externally calculated (given) parame- ters which may be used to alter the material characteristics during the calculation
***contact	defines surfaces of possible contact and the method of en- forcement in the event there is contact
***bc	specify both the geometrical and force boundary conditions

***table	specify the tabular loading magnitudes for parameters and boundary conditions
***function	specify functional loading magnitudes for parameters and boundary conditions in terms of the time
***material	gives the information for material files as attached to el- ement sets, local integration methods, material rotations, and initial variable values
***output	used to specify the desired output from the analysis; mul- tiple output sections can be given to optimize the solution storage

## \*\*\*\*calcul dynamic

#### **Description**:

This option of the **\*\*\*\*calcul** command indicates that dynamic effects should be taken into account. The solution procedure is either an implicit d-form of the Newmark time integration scheme or the  $\alpha$ -method form Hilber, Hughes and Taylor. These methods are compatible with all mechanical element formulations. Most of the following explanations can be found in [hughes87] and [belytschko00]

The semi-discrete equation of motion (discretized in space, continuous in time) is written as:

$$\mathbf{Ma}(t) + \mathbf{F}_{damp}(\mathbf{v}(t)) + \mathbf{F}_{int}(\mathbf{d}(t)) = \mathbf{F}_{ext}(t)$$
(1)

where  $\mathbf{d}(t)$ ,  $\mathbf{v}(t) = \mathbf{\dot{d}}(t)$  and  $\mathbf{a}(t) = \mathbf{\dot{v}}(t)$  are the vectors of nodal displacement, velocity and acceleration respectively. **M** is the mass matrix and  $\mathbf{F}_{damp}$  and  $\mathbf{F}_{ext}(t)$  are the nodal vectors of damping and external forces respectively. The following initial conditions hold:

$$\mathbf{d}(0) = \mathbf{d}_0 \tag{2}$$

$$\mathbf{v}(0) = \mathbf{v}_0 \tag{3}$$

The linear (or linearized) form of equation (1) reads:

$$\mathbf{M}\ddot{\mathbf{d}} + \mathbf{C}(\dot{\mathbf{d}}) + \mathbf{K}(\mathbf{d}) = \mathbf{F}_{ext}$$
(4)

where  $\mathbf{C}$  and  $\mathbf{K}$  are the damping and rigidity (or tangential rigidity) matrices respectively.

#### Newmark schemes:

...

The methods of the Newmark family consist in discretizing equation (4) in time in the following way:

$$\mathbf{Ma}^{t+\Delta t} + \mathbf{Cv}^{t+\Delta t} + \mathbf{Kd}^{t+\Delta t} = \mathbf{F}_{ext}^{t+\Delta t}$$
(5)

$$\mathbf{d}^{t+\Delta t} = \widetilde{\mathbf{d}}^{t+\Delta t} + \Delta t^2 \beta \mathbf{a}^{t+\Delta t}$$
(6)

$$\mathbf{v}^{t+\Delta t} = \widetilde{\mathbf{v}}^{t+\Delta t} + \Delta t \gamma \mathbf{a}^{t+\Delta t} \tag{7}$$

where the predictors (known from the previous increment)  $\widetilde{\mathbf{d}}^{t+\Delta t}$  and  $\widetilde{\mathbf{v}}^{t+\Delta t}$  are defined by:

$$\widetilde{\mathbf{d}}^{t+\Delta t} = \mathbf{d}^t + \Delta t \mathbf{v}^t + \frac{\Delta t^2}{2} (1 - 2\beta) \mathbf{a}^t$$
(8)

$$\widetilde{\mathbf{v}}^{t+\Delta t} = \mathbf{v}^t + \Delta t \left(1 - \gamma\right) \mathbf{a}^t \tag{9}$$

 $\mathbf{d}^t$ ,  $\mathbf{v}^t$  and  $\mathbf{a}^t$  are approximations of  $\mathbf{d}(t)$ ,  $\dot{\mathbf{d}}(t)$  and  $\ddot{\mathbf{d}}(t)$  respectively. Parameters  $\beta$  and  $\gamma$  determine the stability and accuracy of the algorithm.

There are several possible implementations of the Newmark algorithm. The one used in Z-set is the d-form, meaning that the equations are solved in terms of  $\mathbf{d}$  (*i.e.* not in terms of  $\mathbf{a}$  or  $\mathbf{v}$ ). The linear system to solve then reads:

$$\left(\frac{1}{\beta\Delta t^2}\mathbf{M} + \frac{\gamma}{\beta\Delta t}\mathbf{C} + K\right)\mathbf{d}^{t+\Delta t} = \mathbf{F}_{ext}^{t+\Delta t} + \left(\frac{1}{\beta\Delta t^2}\mathbf{M} + \frac{\gamma}{\beta\Delta t}\mathbf{C}\right)\widetilde{\mathbf{d}}^{t+\Delta t} - C\widetilde{\mathbf{v}}^{t+\Delta t} \quad (10)$$

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Note that the choice  $\beta = 0$  (corresponding to the explicit Newmark algorithm if **M** and **C** are diagonal) is not suitable for the d-form.

#### $\underline{\alpha}$ -method (HHT):

The  $\alpha$ -method introduced by Hilber, Hughes and Taylor owns to a more general class of integration schemes called linear multisteps methods (LMS). The  $\alpha$ -method is only slightly different from the Newmark one: the update equations (6-9) are conserved, the difference lies in the time-discrete equation which now reads:

$$\mathbf{M}\mathbf{a}^{t+\Delta t} + (1-\alpha)\mathbf{C}\mathbf{v}^{t+\Delta t} + \alpha\mathbf{C}\mathbf{v}^{t} + (1-\alpha)\mathbf{K}\mathbf{d}^{t+\Delta t} + \alpha\mathbf{K}\mathbf{d}^{t} = (1-\alpha)\mathbf{F}_{ext}^{t+\Delta t} + \alpha\mathbf{F}_{ext}^{t}$$
(11)

By setting  $\alpha = 0$ , the Newmark family of time integration methods is recovered. The  $\alpha$ -method is usually used with the following set of parameters:

$$\gamma = \frac{1}{2} + \alpha \qquad \beta = \frac{(1+\alpha)^2}{4} \qquad \alpha \in \left[0, \frac{1}{3}\right]$$
(12)

The reasons of this choice will be explained in the following section.

#### Linear problems:

For linear problems, the behaviors of the Newmark and  $\alpha$ -methods are well known. To select the appropriate set of parameters  $(\alpha, \beta, \gamma)$ , three points are of particular importance:

- stability (conditional or unconditional)
- order of convergence
- controllable algorithmic dissipation of the high-frequency modes

The last attribute if often desirable in structural dynamics problems. High-frequency modes are poorly approximated by the spatial finite element discretization. By employing algorithms with high-frequency dissipation, spurious high-frequency response is damped out. Properties of some classical methods are summarized in Table 1. Generally, the stability condition reads:

unconditional 
$$2\beta \ge \gamma \ge \frac{1}{2}$$
 (13)

conditional 
$$\gamma \ge \frac{1}{2}, \quad \beta < \frac{\gamma}{2} \quad \Rightarrow \quad \omega^h \Delta t \le \Omega_{\text{crit}}$$
 (14)

Where  $\omega^h$  corresponds to the highest pulsation of the spatially discretized problem (*h* is related to the spatial discretization, *i.e.* to the size of the finite elements) and  $\Omega_{\rm crit}$  depends on the physical damping parameter. It can be shown that  $\omega^h$  is bounded by the maximum element pulsation  $\omega^h \leq \omega_e^h$  which increases when *h* decreases. As a consequence, for conditional stability, the critical time step decreases with decreasing element size. Moreover, selecting  $\gamma = \frac{1}{2}$  ensures a second order accuracy but adds no damping of high-frequency modes. Selecting  $\gamma > \frac{1}{2} = \frac{1}{2} + \alpha$  allows artificial (purely numerical) damping of spurious high frequency modes. This damping is maximized for  $\beta = \frac{(1+\alpha)^2}{4}$ . Within the Newmark framework, this choice leads to a first-order accuracy whereas a second-order accuracy is achieved with the  $\alpha$ -method, and this is the main advantage of the  $\alpha$ -method compared to the Newmark one. Note that damping

Method	β	$\gamma$	Stability condition	order of ac- curacy	
Central difference	0	$\frac{1}{2}$	$\omega^h \Delta t \leq 2$	2	not suitable for the d-form
Linear acceleration Fox-Goodwin	$\frac{\frac{1}{6}}{\frac{1}{2}}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\begin{aligned} \omega^h \Delta t &\leq 2\sqrt{3} \\ \omega^h \Delta t &< \sqrt{6} \end{aligned}$	$\frac{2}{2}$	no HF damping no HF damping
Average acceleration (trapezoidal rule)	$\frac{2}{4}$	$\frac{2}{1}$	unconditional	2	no HF damping
Newmark modified average acceleration ( $\alpha > 0$ )	$\frac{(1+\alpha)^2}{4}$	$\frac{1}{2} + \alpha$	unconditional	1	$\begin{array}{ll} \text{HF} & \text{damping} \\ \text{proportional} & \text{to} \\ \alpha \end{array}$
$\alpha$ -method ( $\alpha > 0$ )	$\frac{(1+\alpha)^2}{4}$	$\frac{1}{2} + \alpha$	unconditional	2	$\begin{array}{ll} \text{HF} & \text{damping} \\ \text{proportional to} \\ \alpha \end{array}$

Table 1: Properties of some classical methods for linear dynamics. Stability conditions are given for physically undamped problems.

is proportional to  $\alpha$  and that low-frequency modes are affected more strongly for higher values of  $\alpha$ .

#### Nonlinear problems:

The notion of stability and accuracy developed for linear dynamics are not sufficient for nonlinear problems. The use of the previous rules does not guarantee stability.

### Syntax:

Dynamic calculations accept a subset of the **\*\*\*\*-level** commands, with an additional **\*\*\*init\_velocity** command used to specify initial velocity conditions. The dynamics specific commands are found in the following table:

CODE	DESCRIPTION
***resolution	same as for the static case with specification of the time
	integration scheme parameters $\alpha$ , $\beta$ and $\gamma$ within the sequence definition (see **sequence)
***init_velocity	set up initial velocities

The command **\*\*\*init\_velocity** has the following syntax:

***init_velocity				
$dof\_name$ elset	$elset\_name$	value		

By default, the initial velocity is set up to 0 everywhere.

The selection of the integration scheme (Newmark or  $\alpha$ -method) and the parameter definition can be done within the **\*\*sequence** block (see page 3.190) through optional commands

\*alpha, \*beta and \*gamma. All other \*\*sequence sub-commands are available and remain unchanged.

	**sequence	Γ	N	]			
[	*alpha	val1	[	val2,	valN	]	]
[	*beta	val1	[	val2,	valN	]	]
Γ	*gamma	val1	Γ	val2.	valN	]	]

These commands can be used with the following increasing level of description:

- none of these command is used. The  $\alpha$ -method is selected with  $\alpha = 0.05$ ,  $\beta = \frac{(1+\alpha)^2}{4}$  and  $\gamma = \frac{1}{2} + \alpha$ .
- only \*alpha is used. This selects the coefficients of the  $\alpha$ -method with  $\beta = \frac{(1+\alpha)^2}{4}$  and  $\gamma = \frac{1}{2} + \alpha$ .
- only \*alpha and \*gamma are used. This selects the coefficients of the  $\alpha$ -method with  $\beta = \frac{(\frac{1}{2} + \gamma)^2}{4}$ .
- \*alpha, \*beta and \*gamma are used. The three parameters are defined independently.

To select the Newmark integration scheme,  $\alpha$  must be set to 0. Due to its unconditional stability and good convergence rate, it is recommended using the  $\alpha$ -method in accordance with relations (12), even for non-linear problems.

#### Example:

An example implicit dynamic calculation follows. In the first sequence a Newmark average acceleration scheme is selected. In the second one, an  $\alpha$ -method with  $\alpha = 0.1$  has been chosen. Initial velocity of nset INIT1 is set up to 0.1. Note the expiring boundary condition to move and then release a load point.

```
****calcul dynamic
***mesh updated_lagrangian_plane_strain
***resolution
 **sequence
                 1.0 1.0
   *dtime
   *increment
                 10 10
   *alpha
                 0.
                     0.1
   *ratio absolu 1.e-6
   *algorithm
                 p1p2p3
 ***bc
 **impose_nodal_dof
                U2
                     0.0
      wall
                U1
                     0.0
      wall
           exp U2 -1.0 tab
      load
***table
 **name tab
   *time
            0.0
                  1.
   *value
            0.0
                  1.
 ***init_velocity
```

```
U1 elset INIT1 0.1
***output
**curve dynam_ul.test
*node_var 6 U2
***material
*file ../MAT/dynam_ul
****return
```

### \*\*\*\*calcul mechanical\_explicit

#### **Description**:

This option of the **\*\*\*\*calcul** command is used to activate the explicit solver. Such a solver can be used to model and capture short time scale phenomena such as waves propagation. Explicit solvers can although be used for rough problems like crash, impact or metal sheet forming where contact occurs on large surfaces (contact is very rough from a computational point of view).

#### Central difference explicit schemes:

The central difference explicit scheme can be derived from the Newmark integration scheme (eq. (5)–(9) of the **\*\*\*\*calcul dynamic** section) with  $\beta = 0$  and  $\gamma = 0.5$ . The a-form of the integration scheme then reads:

$$\left(\mathbf{M} + \frac{\Delta t}{2}\mathbf{C}\right)\mathbf{a}^{t+\Delta t} = \mathbf{F}_{ext}^{t+\Delta t} - \mathbf{C}\mathbf{v}^{t+\Delta t/2} - K\mathbf{d}^{t+\Delta t}$$
(15)

with

$$\mathbf{v}^{t+\Delta t/2} = \frac{1}{\Delta t} \left( \mathbf{d}^{t+\Delta t} - \mathbf{d}^t \right) \tag{16}$$

The following expression follows from equations (6) and (9):

$$\mathbf{a}^{t} = \frac{1}{\Delta t^{2}} \left( \mathbf{d}^{t+\Delta t} - 2\mathbf{d}^{t} + \mathbf{d}^{t-\Delta t} \right) \quad \text{and} \quad \mathbf{v}^{t+\Delta t/2} = \mathbf{v}^{t-\Delta t/2} + \Delta t \mathbf{a}^{t}$$
(17)

hence the name "central difference" scheme. Note that  $\mathbf{d}^{t+\Delta t}$  and  $\mathbf{v}^{t+\Delta t/2}$  are known from the previous step at time t (equations (17) and (16)) so that the right hand side of (15) is known.

In order to make this scheme explicit, a diagonalization of  $\mathbf{M}$  and  $\mathbf{C}$  is performed (also called a lump) so that the solution of equation (15) is trivial. Each time step is therefore solved very quickly through a simple matrix vector product and there is no linear system to solve. The basic algorithm is described below:

1. initiate  $\mathbf{d}^0$  and  $\mathbf{v}^0$ 

2. compute 
$$\mathbf{a}^0 = \mathbf{M}^{-1} \left( \mathbf{F}_{ext}^0 - \mathbf{K} \mathbf{d}^0 - \mathbf{C} \mathbf{v}^0 \right)$$
 and  $\mathbf{v}^{1/2} = \mathbf{v}^0 + \frac{\Delta t}{2} \mathbf{a}^0$ 

- 3. enforce velocity boundary conditions
- 4. increment time from  $t \Delta t$  to t and compute  $\mathbf{d}^t = \mathbf{d}^{t-\Delta t} + \Delta t \mathbf{v}^{t-\Delta t/2}$
- 5. compute  $\mathbf{a}^t = \mathbf{M}^{-1} \left( \mathbf{F}_{ext}^t \mathbf{K} \mathbf{d}^t \mathbf{C} \mathbf{v}^{t-\Delta t/2} \right)$
- 6. compute  $\mathbf{v}^{t+\Delta t/2} = \mathbf{v}^{t-\Delta t/2} + \Delta t \mathbf{a}^t$
- 7. if computation not finished, go to 3

The drawback of explicit algorithms lies in the stability condition that imposes that the time step  $\Delta t$  is bounded by a critical time step  $\Delta t_{crit}$ . The stability criterion for explicit central difference method reads (damping has no effect on stability):

$$\omega^h \Delta t \le 2 \quad \Rightarrow \quad \Delta t_{crit} = 2/\omega^h \tag{18}$$

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3.15

where  $\omega^h$  is the highest natural frequency of the discretized structure. Computing  $\omega^h$  is very expensive since it requires solving a large eigen values system. It can be shown that  $\omega^h$  is bounded by the maximum frequency of individual elements:

$$\omega^h < \omega^h_{el} \tag{19}$$

Let us take the example of a linear beam element with stiffness  $\mathbf{K}$  and lumped mass matrix  $\mathbf{M}$  such that:

$$\mathbf{K} = \frac{ES}{h} \begin{pmatrix} 1 & -1 \\ -1 & 1 \end{pmatrix} \qquad \mathbf{M} = \frac{\rho h S}{2} \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}$$
(20)

where h, S, E and  $\rho$  are the element length, section, Young's modulus and mass density respectively. The non-zero solution of the eigen value problem  $K - \omega_{el}^{h^2} M$  leads to  $\omega_{el}^{h} = 2/h\sqrt{E/\rho} = 2c/h$  where c is the wave speed. The critical time step can therefore be written:

$$\Delta t_{crit} = h/c \tag{21}$$

This corresponds to the time for the wave to go through the element. This interpretation can be verified for every kind of finite element. In practice, this last remark is used in Z-set to evaluate  $\Delta t_{crit}$ . The characteristic size of the element is taken as the minimum distance between two nodes of the element. The wave speed is taken to be the longitudinal wave speed (the fastest one) which reads :

$$c_L = \sqrt{\frac{\lambda + 2G}{\rho}} \tag{22}$$

where  $\lambda$  and G are the Lamé coefficients. An important property of finite elements applied to hyperbolic problems is that  $\Delta t_{crit}$  is O(h) whereas it is  $O(h^2)$  for parabolic problems. This makes explicit methods difficult to apply to parabolic problems such as heat conduction or diffusion in general. It is important to note that the critical time step is governed by the size of the smallest element in the mesh. If there is only one spurious small element (due to a bad meshing for instance), the computational cost would increase.

Explicit algorithms are very interesting in the sense that no linear system need to be solved. However, a major limitation lies in the O(h) critical time step. Explicit methods are therefore well suited to model problems where small time steps are imposed by the physics. If there is no need for small time steps, implicit methods may be economically competitive.

#### Non linear problems:

Extension to non-linear problems is direct and equation (15) is replaced by:

$$\left(\mathbf{M} + \Delta t\mathbf{C}\right)\mathbf{a}^{t+\Delta t} = \mathbf{F}_{ext}^{t+\Delta t} - \mathbf{F}_{int}^{t+\Delta t}(\mathbf{d}^{t+\Delta t}, \mathbf{v}^{t+\Delta t/2})$$
(23)

In the algorithm presented above, the computation of  $\mathbf{F}_{int}^{t+\Delta t}(\mathbf{d}^{t+\Delta t}, \mathbf{v}^{t+\Delta t/2})$  is performed between steps 3 and 5.

The stability condition (18) emanates from an analysis of linear equations. At this time, there is no stability theorem that covers the range of nonlinear phenomena such as contactimpact. Instability cannot be overlooked in linear problems since the solution tends to grow exponentially. However, this is not always true for nonlinear equations. A good way to detect instabilities is to check the energy balance and make sure that no spurious energy is created. The default stability criterion used in Z-set is defined as:

$$|W_{\rm kin} + W_{\rm int} + W_{\rm dmp} - W_{\rm ext}| \le \varepsilon \max\left(W_{\rm kin}, W_{\rm int}, W_{\rm ext}\right) \tag{24}$$

where  $W_{\rm kin}$  is the kinematic energy,  $W_{\rm int}$  and  $W_{\rm ext}$  are works done by internal and external forces respectively, and  $W_{\rm dmp}$  is the energy dissipated by damping. An absolute criterion can also be used :

$$|W_{\rm kin} + W_{\rm int} + W_{\rm dmp} - W_{\rm ext}| \le \varepsilon \tag{25}$$

 $\varepsilon$  is a small value defined with the **\*\*sequence \*ratio** command. The absolute criterion is selected by using the *absolute* option (see below). If the energy balance is not verified, the sub-step is recomputed with half time step.

#### Syntax:

The resolution procedure is specified by the usual way using the **\*\*\*resolution** command. Specific commands are therefore added to control some parameters. Here are the different options available within the resolution block:

```
***resolution
 **sequence [ N ]
[ *ratio [absolute] val1 [ val2 ... valN ] ]
[ *beta val1 [ val2 ... valN ] ]
[ *fixed_dt val1 [ val2 ... valN ] ]
[ *max_dt val1 [ val2 ... valN ] ]
[ *min_dt val1 [ val2 ... valN ] ]
[ *damping val1 [ val2 ... valN ] ]
[ *max_successive val1 [ val2 ... valN ] ]
[ *show_gauge]
```

**\*\*sequence** has the same syntax as in **\*\*\*\*calcul** (see page 3.190) with additional commands. Note that the time stepping is not defined by the number of increments set in **\*\*sequence** but is either automatically computed by Z-set (by default) or explicitly specified using **\*fixed\_dt**. If the time increment is larger than the algorithm time step, sub-stepping occurs. The number of sub-steps performed within each sequence increment is displayed in the standard output.

\*ratio [absolute] value, defines  $\varepsilon$  in (24) (or in (25) if the keyword absolute is used).

- \*beta value. If the critical time step is computed by Z-set, the effective time step is taken to be  $\Delta t_{crit} \times value$ . Values between 0.8 (default) and 0.9 are usual.
- \*fixed\_dt value, specifies the time step to use. This value is not affected by \*beta.
- \*max\_dt value, specifies the upper bound of the time step
- \*min\_dt value, specifies the lower bound of the time step. If the time step is lower than
  value, an error message is sent and the computation is stopped.
- \*damping value, add constant diagonal components in the damping matrix **C**. This command is similar to the command \*\*\*explicit \*\*damp (see below).

- \*max\_successive value, specifies that if the energy balance is not verified after value successive sub-steps (dividing each successive time step by 2), Z-set will stop and send an error message. The default value is set to 50 which is a relatively large value.
- **\*\*show\_gauge** allows to display the sub-stepping evolution within an increment. A percent value is displayed in the standard output. This can be useful to get an idea on how fast the computation goes when a lot of sub-steps are performed.

Additional options are also available through the **\*\*\*explicit** command.

***explicit				
**damp [constant	] value			
**every_update value				

- **\*\*damp** [constant] value, is similar to **\*damping**. Each **\*\*damp** command is cumulative. At the moment, the only damp type is "constant" (default).
- \*\*every\_update value, specifies that the critical time step is computed every value time steps. The default value is set to 20. Recomputing the critical time step is useful within a finite strain calculation, where large deformation of some elements can indeed modify  $\Delta t_{crit}$  (*i.e.* the time for the wave to go through the element can decrease or increase). However, this can be time consuming, please check CPU time at the end of the computation.

#### Example:

An example explicit dynamic calculation follows.

```
****calcul explicit_mechanical
 ***mesh plane_strain
 ***resolution
 **sequence
   *time 10. 20.
   *increment 10 30
   *fixed_dt 0.05 0.01
   *ratio
           1.e-2
   *max_successive 10. 20.
   *beta 0.9 0.8
 **show_gauge
 ***explicit
 **every_update 1
 **damp constant 1.e-3
 ***bc
 **impose_nodal_dof
    left
                 U2
                      0.0
    left
                 U1
                      0.0
```

```
**pressure
right 0.00001 echelon
***table
**name echelon
*time 0.0 1.e-10 10000000.
*value 0.0 1.0 1.0
***material
**elset R1
*file dynam_unit
***return
```

# \*\*\*\*calcul eigen

#### **Description**:

This option of the **\*\*\*\*calcul** command indicates that eigen value solution of the natural frequencies are to be calculated. The solution of the eigen-value problem is controlled using the command **\*\*\*eigen** (see page 3.107).

Eigen mode calculations accept a sub-set of the above commands, with an additional command **\*\*\*eigen** used to give parameters of the eigen problem solution. The allowable commands are found in the following table:

CODE	DESCRIPTION
***mesh	same as above
***eigen	give eigen solution parameters and resolution method
***equation	same as above
***bc	same as above; only fixed conditions are allowed
***material	same as above
***output	specifies what output to save at the end of an eigen solution; this is a sub-set of the above

#### Example:

An example file for Eigen solution follows.

```
****calcul eigen
***mesh
  **elset solid small_deformation
  **elset springs spr1
***eigen lanczos
    6 0.01 2.
***bc
   **impose_nodal_dof
     base
            U1 0.00
            U2 0.00
     base
            U3 0.00
     base
***output
***material
   *file building.mat
****return
```

## \*\*\*\*calcul thermal\_transient

#### **Description**:

This option of the **\*\*\*\*calcul** command indicates that the problem is transient thermal analysis.

## Example:

```
****calcul thermal_transient
  ***resolution
    **sequence
      *time
              10. 8010.0
      *increment 10 100
      *iteration 50 50
      *ratio 0.01 0.01
      *algorithm p1p1p1
  ***bc
    **fluconv exte
      h 232.5
      Te 1000.0 tab1
  ***init_dof_value
      TP uniform 20.
  ***material
      *file ../MAT/TTLL_02_89
  ***table
    **name tab1
      *time 0.
                   50000.
      *value 1.
                   1.
  ***output
    **curve
      *precision 4 *small 1.e-4
      *node_var 1 TP
      *node_var 16 TP
****return
```

# \*\*\*\*calcul diffusion

# **Description**:

This option of the **\*\*\*\*calcul** command indicates that the problem is a diffusion one, obeying Ficks law.

CODE	DESCRIPTION
***impose_kinematic	used to allow the geometry to evolve which can be very
	useful for coupled analysis

# \*\*\*\*calcul weak\_coupling

#### **Description**:

The weak-coupling implementation uses an iterative approach to an arbitrary coupled problem. Any number of sub-problems are solved (thus keeping the individual systems small) with the coupling taking place through appropriate transfer of results between problems. For example, a mechanical problem can transfer internal heat generation due to mechanical dissipation to a thermal problem which calculates the resulting temperature field evolution which is re-transferred to the mechanical problem to allow coefficient alteration.

#### Syntax:

Some special commands are of interest for the coupled problem. These define the sub-problem files (standard .inp files), and the method of determination for convergence of the coupled problems.

- **\*\*\*resolution** define the global time steps to be run; convergence parameters for the sub-problems are determined in their input files.
- **\*\*\*coupled\_resolution** additional convergence parameters specific to the coupled solution.
- **\*\*\*sub\_problem** specify a sub-problem to be added; these will be run in the order they are entered.

### Example:

The following is the top level command file for the test \$Z7PATH/test/Coupled\_test/INP/MechTherm.inp

```
****calcul weak_coupling
   ***resolution
    **sequence
                4.0 8.0
     *time
     *increment 10 10
                1.e-4
     *ratio
   ***coupled_resolution
     **iteration 2
   ***sub_problem fem MechTherm/plastic
    **transfer integ_nodeparam
     *variable q_dot
     *file
               MechTherm/heat_out
    **transfer node_kinematic
     *file
               MechTherm/kine_out
   ***sub_problem fem MechTherm/thermal
    **transfer node_nodeparam
     *variable TP
               MechTherm/temp_out
     *file
***return
```

This section list all three stars commands available under **\*\*\*\*calcul**, you may find an exhaustive list of these commands in the index at the end of this handbook page 7.2.

# $***linear_solver$

### **Description**:

This keyword specifies the solver used to solve the linear system of equations involved in the global step of the Newton-Raphson algorithm. Both direct and iterative solvers are implemented in Z-set.

### Syntax:

#### **\*\*\*linear\_solver** type

Where type is a solver type, which can take the following values.

- frontal is a direct frontal solver based on a Cholesky factorization, so matrices have to be symmetric, definite and positive. This solver is the default solver.
- sparse\_direct is a direct solver using sparse storage, i.e. only non-zero terms of the
  factorized matrix are stored. So it is less memory consuming than the frontal solver.
  This solver is based on a Crout factorization, so matrices have to be symmetric, definite,
  but not necessary positive.
- sparse\_dscpack is an optimized direct solver using sparse storage. It is based on a multifrontal algorithm using a Cholesky factorization, so matrices have to be symmetric definite and positive. To be very efficient, this solver uses the BLAS optimized mathematical library working on full matrices, so it is locally more memory consuming than the sparse\_direct solver. This solver is developed by Padma Raghavan of the Pennsylvania State University, and used by permission. Information on the DSCPACK solver is available on Web at the link : http://www.cse.psu.edu/~raghavan/Dscpack/dscpack.ps
- sparse\_iterative includes all the available iterative solvers. These solvers are less memory consuming than direct ones, because matrices have never to be assembled, but if matrices have a bad condition number, it can be very hard to achieve convergence. Two iterative solvers can be used in Z-set, a Conjugate Gradient algorithm (cg), or a Global Minimum RESidual one (gmres). The main difference between these two solvers, is the assurance to achieve convergence if matrices are non-positive using the GMRES algorithm. This solvers type needs some complementary subkeywords described in section \*\*\*linear\_solver sparse\_iterative page 3.27.

### Example:

The following table gives memory needed, total CPU time and number of iterations to solve using different solvers a traction problem for a cube meshed with 13824 linear elements corresponding to 46875 dofs.

	Total CPU time	Memory needed (Mb)	Number of iterations
Frontal	48.22 mm	730 Mb	
Sparse_direct	27.82 mn	489 Mb	
Sparse_dscpack	$3.52 \mathrm{~mn}$	$538 { m ~Mb}$	
CG + lumped	1.92 mn	146 Mb	238
CG + Cholesky	2.17 mn	160 Mb	94
GMRES + lumped	$5.9 \mathrm{mm}$	161 Mb	750
GMRES + Cholesky	3.12  mm	$175 { m ~Mb}$	169

# \*\*\*linear\_solver sparse\_iterative

### **Description**:

This keyword specifies the iterative solver used to solve the linear system of equations involved in the global step of the Newton-Raphson algorithm.

# Syntax:

[**precond type]		
[**full_output]		
[**solver solver]		

Where

- **\*\*precond** specifies the *type* of pre-conditioning used to accelerate convergence. The two available types are lumped (e.g. diagonal) and cholesky. Default value is lumped. Using cholesky preconditioner can significantly reduce the number of iterations even if it needs to evaluate and factorize an supplementary matrix, unlike lumped preconditioner.
- **\*\*full\_output** enables to print convergence informations. Default value is FALSE. If FALSE, nothing is said about solver iterations.
- **\*\*solver** specifies the type of iterative solver used. *solver* can take these two values cg (i.e. Conjuguate Gradient) or gmres (i.e. Global Minimum RESidual). Default value is cg. According to the chosen solver, the following keywords are different.

### Syntax:

For the Conjuguate Gradient algorithm, syntax is the following:

[\*max\_iteration max\_iter ]
[\*precision eps ]
[\*output\_every\_iter nb\_iter]
[\*output\_to\_file file]
[\*keep\_direction dir]
[\*max\_standing max]
[\*min\_iter min\_iter]
[\*reprojection]

- \*max\_iteration max\_iter, where max\_iter is the maximum number of allowed iterations when solving the problem. Default value is 1000.
- \*precision eps, where eps is a real value defining the relative precision required for convergence when solving the problem with the CG method. Default value is 1e-08. For a system of equation:

### $\mathbf{K}\mathbf{q} = \mathbf{F}$

this relative ratio is defined by:

$$ratio = \frac{||\mathbf{F} - \mathbf{K}\mathbf{q}||}{||\mathbf{F}||}$$

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and convergence occurs when:

ratio < eps

- \*output\_every\_iter *iter*, where *iter* is the frequency of the iteration information output. This option is only active if the keyword \*\*\*full\_output is *TRUE*. Default value is 10.
- \*output\_to\_file file, where file is a character string specifying the file where iteration information are written. This option is only active if the keyword \*\*\*full\_output is TRUE. Default value is iterative\_solver\_it.
- \*keep\_direction *dir*, where *dir* is the the integer value specifying the number of orthogonal descent directions retained during the CG iterations. Increasing *dir* leads to faster convergence but is more memory consuming. Default value is *max\_iter*+2.
- \*max\_standing max, where max is an integer specifying the maximum number of CG iterations allowed without any significant decrease of the convergence ratio. Default value is 50.
- \*min\_iter min\_iter, where min\_iter is the minimum iterations when solving the problem.
  Default value is 1.
- \*reprojection This subcommand can significantly reduce the number of CG iterations, when used in conjunction with quasi-Newton schemes of tangent matrix update (such as eeeeee or p1p1p1, see the \*\*algorithm command). With this option the descent directions calculated during previous load increments are reused, leading to convergence in just a few iterations when the tangent matrix and the load increment stay constant over several Newton increments.

### Syntax:

For the Global Minimum RESidual algorithm, syntax is the following :

```
[*max_iteration max_iter ]
[*precision eps ]
[*output_every_iter nb_iter]
[*output_to_file file]
[*krylov_space krylov_dim ]
```

Where \*max\_iteration, \*precision, \*output\_every\_iter, \*output\_to\_file are the same that for CG solver.

\*krylov\_space krylov\_dim, where krylov\_dim is an integer value specifying the dimension of the krylov\_space built for each cycle of the GMRES algorithm. Increasing this value leads to faster convergence.

# Example:

```
***linear_solver sparse_iterative
    **full_output
    **precond cholesky
```

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\*\*\*\*calcul
\*\*\*linear\_solver
sparse\_iterative

```
**solver cg
*output_to_file iterations.hist
*precision 1.e-12
*output_every_iter 1
*max_iteration 1000
```

# \*\*\*linear\_solver rigid

## **Description**:

This keyword specifies a "wrapper" solver which encapsulate a real solver. It behaves exactly like the underlying solver, except that the kernel of the operator is computed. If requested, a boundary condition is automatically added to fix rigid body motions.

The way rigid body motions are computed is explained in the theory manual, at the linear solver chapter.

# Syntax:

[\*\*local\_solver type ]
 options for the local solver
[\*\*create\_bc]
[\*\*verbose solver]

Where

\*\*verbose asks the wrapper to print detailed informations about rigid body motions found,

- **\*\*local\_solver** allow to choose the underlying linear solver. Any linear solver may be used, but note that iterative one sometimes may exhibit weird behavior (especially depending on the convergence ratio used for these solvers),
- **\*\*create\_bc** specifies that a boundary condition has to be added to fix all body motions. Note also that in sequential computations the presence of body motions often means that there is an error in the input file.

# \*\*\*auto\_remesh

# **Description**:

The **\*\*\*auto\_remesh** section is derived from **\*\*\*initialize\_with\_transfer** and allows automatic remeshing during a computation without stopping it. This is a backbone feature for crack propagation or computations using adaptive mesh.

Basically the **auto\_remesh** bloc of commands is divided into:

- mesher commands controlling the remesh process.
- timing commands that trigger the automatic remeshing.
- transfer commands that specify how to transfer data between original and modified meshes.

## <u>Note</u>:

- The use of the  $\mathbf{Z8}$  output database is mandatory.
- auto\_remesh transfer only variables that have been saved to the output database. To transfer all variables, one should use **\*\*save\_all** command in **\*\*\*output** command bloc
- The graphical interface scripts **Zcrack** and **Zxfem** may be used to handle crack propagation problems and automatically generate input data for this command.

### Syntax:

**\*\*\*auto\_remesh** takes a number of **\*\*\*initialize\_with\_transfer** controls, and some additional specific commands.

```
***auto_remesh
  [**output_after_remesh]
  [**no_deform_mesh]
  [**each_incr]
  [**frequency]
         . . .
  [**remeshing_criterion criterion ]
   **mesher_commands
% below options are inherited from initialize_with_transfer, see 3.130\%
  [**quiet]
  [**reequilibrium
    *algo algorithm
    *ratio convergence
    *iter max_iterations ]
  [**skip_nodal_transfer]
  [**skip_integ_transfer]
  [**nodal_var_transfer
    [*mapping mapping_method ] ]
  [**integ_var_transfer
    [*integ_transfer transfer_method ] ]
```

- **\*\*output\_after\_remesh** save the results of the transfer to the database. With this option, Z8 database will contain two copies of the same results: the original data on the original mesh, and the transferred data on the new mesh obtained as a result of the specified mesher commands.
- **\*\*no\_deform\_mesh** use initial configuration of the old mesh to locate the nodes/IP of current mesh (if not specified, the default behavior is to use deformed mesh)
- **\*\*each\_incr** force remeshing on each increment of current computation.
- **\*\*frequency** used to trigger the remeshing process, for more sub options see 3.148
- \*\*remeshing\_criterion a more general remeshing trigger, The possible criterion are sum-

CODE	DESCRIPTION	
at_time		
quadratic_curv:	ing_criterion	
element_shape_c	luality	
for_crack_propa	gation	
error_based		

\*\*mesher\_commands mesh manipulators commands see 2.4 for usable commands

### Example:

```
***auto_remesh
%% AUTO_REMESH options
  **output_after_remesh
  **no_deform_mesh
  **frequency
   *at_time 0.02
%% INITIALIZE_WITH_TRANSFER options
  **integ_var_transfer default
   *locator bb_tree
   *integ_transfer nearest_gp_corrected
%% TRANSFORMERS
  **yams_ghs3d
   *no_deform_mesh
   *min_size 0.2
   *absolu
  **nset enc
   *use_bset enc
```

This example can be found in Crack\_test/INP/cube\_2cracks.inp

```
***global_parameter
  Solver.OutputFormat Z8
  Zmaster.OutputFormat Z8
***auto_remesh
 **frequency
  *d_cycle 1
  *cycle_period 2.0
  *at 2.0
 **no_deform_mesh
 **skip_integ_transfer
 **skip_nodal_transfer
 **save TO_REMESH.geo
 **drive_crack
  *mesher cube_2cracks.inp
  *geo_in TO_REMESH.geo
  *geo_out REMESHED.geo
  *advance cube_2cracks.adv
  *separate
 **open REMESHED.geo
```

```
***output
 **save_parameter
 **save_all
```

#### \*\*\*bc

### **Description**:

This procedure and its options define the boundary conditions of a problem. The definition of boundary conditions groups both conditions acting on the degrees of liberty and those acting on the associated forces.

## Syntax:

There may be any number of sub-options defining the different conditions to impose, and also any number of **\*\*\*bc** instances.

The different types of boundary conditions are selected by using different values for *bc*type. These different BC commands are the subject of the following pages. The pages of the sub commands are sorted according to their applicability to specific problem types. The following tables are included as a quick-directory to the BCs.

If the optional arguments from t1 to t2 are specified, all the bounday conditions declared in the current \*\*\*bc block will be active only between times t1 and t2. You may declare several \*\*\*bc to partition the activity intervals of your boundary conditions.

### General purpose BCs:

Some BCs are general-purpose, being specified in terms of any degree of freedom in the problem, and their attached location (nodal or elemental).

CODE	DESCRIPTION	
impose_nodal_dof	used to directly impose any DOF value which is located at a node - $p.3.45$	
<pre>impose_nodal_dof_rate</pre>	used to impose any DOF value in a rate form - $\mathrm{p.}3.47$	
impose_nodal_reaction	sets a node's associated force - $\mathrm{p.}3.47$	
<pre>impose_nodal_reaction_rate</pre>	rate of the nodes reaction - $\mathrm{p.3.50}$	
<pre>impose_element_dof</pre>	Fix DOFs which exist at element Gauss points - p.3.41	
<pre>impose_element_dof_reaction</pre>	Set DOF reactions which exist at element Gauss points - $p.3.42$	
<pre>impose_elset_dof</pre>	for DOFs defined over an element set - $\mathrm{p.}3.43$	
<pre>impose_elset_dof_reaction</pre>	reactions associated to DOFs defined over an element set - $\mathrm{p.3.44}$	
<pre>impose_nodal_dof_density</pre>	Density of a nodal dual force over element faces or edges - $p.3.51$	
impose_nodal_energy	Fix DOF until a specified energy is reached	
<pre>impose_nodal_dof_and_release</pre>		
release_nodal_dof	Time based release of fixed DOFs (can be used for crack growth) - $p.3.52$	
submodel	used to directly impose any DOF value at sub- model boundary nodes from a master compu- tation - p.3.53	

# Mechanical BCs:

There are many mechanical specific boundary conditions in Zebulon. A summary follows.

CODE	DESCRIPTION
deformation	used to impose displacements with an
	intermediate (strain) tensor
$deformation\_cosserat$	imposes a "Cosserat" type strain
strain_gradient	displacements as $u = Er + 1/2D \ r \times r$
crack_release	propagate crack (releasing nodes) according to a material variable criterion
radial	submit all or part of a structure to a radial expansion
radius	enforces "rolling" part of a structure on a radius (for example holding the radius of a joint)
pressure	impose a surface pressure on a liset or a faset
shear	impose a shear pressure on a liset (only valid in 2D)
impedance	impose an impedance boundary condition (only valid in dynamics).
hydro	impose a surface pressure on the deformed ge- ometry; this simulates a fluid pressure
curvature	Curvature on a boundary $u_i = \epsilon_{ijk} K_{jl} X_l X_k$
centrifugal	centrifugal loading for all the elements of a structure due to a rotational frequency
gravity	applies a uniform acceleration force to the en- tire structure
rotation	rotate nodes about a given axis
free_rotation	Rotation about an axis with one direction free.
linear_rotation	Rotation with small angle $(\theta \approx \sin \theta)$
linear_free_rotation	Free rotation with linear approx.
K_field	Impose the linear elastic crack tip solution to a node set
static_torsor	Impose a static torsor (resultant and momen- tum) on a nset. It is associated to a rigid body motion of the nset.

# Thermal BCs:

Thermal boundary conditions are provided for a variety of heat flux transfer options. Surface to surface radiation transfer is not yet possible.

CODE	DESCRIPTION
surface_heat_flux	applies a constant heat flux on a liset or a faset - p.3.78
convection_heat_flux	convective flux on a liset or a faset - $\mathrm{p.3.79}$
interface_heat	imposes an inter-facial thermal resistance between two lisets or fasets - $p.3.80$
volumetric_heat	applies a volumetric heat flux on all the elements of a structure - $p.3.81$
volumetric_heat_from_parameter	applies a volumetric heat flux on all the ele- ments of a structure. The values come from a parameter field - p.3.82
volumetric_heat_in_file	Import an internal heat generation from a me- chanical problem (coupled) - $p.3.83$
radiation	heat flux by radiation on a liset or faset - $\mathrm{p.}3.84$

Some of these keywords replace deprecated keywords as follows :

flucons	by	<pre>surface_heat_flux</pre>
fluconv	by	convection_heat_flux
fluconv_interface	by	interface_heat
fluvol	by	volumetric_heat
fluvol_in_file	by	volumetric_heat_in_file

# Mass Diffusion BCs:

CODE

DESCRIPTION

surface\_flux

DESCRIPTION

Impose surface flux of concentration

### Scaling of boundary values:

The magnitude of boundary conditions are normally specified in two parts. The first is a "base value" real number, function or file data (see below) which acts as a multiplicative scale factor. In the event that this value is zero, the BC will be a fixed condition, not requiring a loading table. The second part is a table name which refers to a corresponding defined table (using the **\*\*\*table** command). The table is used to describe the magnitudes in time.

Table specifications may usually take more than one table name. This is very useful in the case of cyclic loadings, or other complex lengthy waveforms. The tables will be sequentially searched for valid times. That is if the first table is defined from 0. to 250., and a second from 200. to 1000., times until 250. will be calculated with the first table, while the remaining time is taken from the second table. If the time exceeds 1000. there will be an error.

## Base value:

As mentioned earlier, the "base value" acts as a multiplicative scale factor for the following table. The standard base value is simply a real number. It may also be a more elaborated object, such as a function or file, which allows space-dependent boundary conditions.

The following example presents all four types of base values currently available:

#### \*\*\*bc

```
**impose_nodal_dof
left U1 0.  % real value
right U1 file right.dat table1 % binary file
bottom U2 ascii_file bottom.dat table2 % ascii file
**pressure
top function sin(x); time % function
```

Note that some multi-point-constraints (see 3.110) also use this concept of base value.

### **Duration of application**:

Normal use will involve defining the value of a boundary condition throughout the time scale of the problem. It is very important to include the value of the BC at zero time.

Boundary conditions can also be applied over a limited time during the problem, with the condition either "expiring" or coming into action after a specified period of time. This is very useful for dynamic problems where an initial movement is given and then released, or if a condition is applied to a certain point, and then continued with a different type condition. An example of the later is to apply force control for a first sequence, followed by a rate of displacement. Note after the load is applied the absolute value of displacement is unknown (for non-linear problems) so a displacement rate must be imposed (e.g. with impose\_nodal\_dof\_rate).

These cases are handled by having the BC's table defined over a limited duration of the problem time scale, and including the keyword **exp** to indicate that the expiration or activation of the BC was intentional.

```
***resolution
 **sequence
 *dtime 1.0 0.95
***bc
 **impose_nodal_dof
 wall U2 0.0
```

\*\*\*\*calcul \*\*\*bc

```
wall U1 0.0
load exp U2 -1.0 tab
***table
**name tab
*time 0.0 1.
*value 0.0 1.
```

### Association to geometry:

Most BCs may also be localized to certain portions of the geometry, through the use of node, element, line and face sets. Specification of these entities are discussed in the meshing chapters *file .inp: 2D meshing* and *file .inp: 3D meshing*. The following general statements can be made:

- Boundary conditions to be applied in order to directly impose the value of nodal degrees of freedom are applied to *nsets*.
- Boundary conditions which are distributed over a surface, or have a mean value for a surface (e.g. pressure) will be applied to an *liset* in 2D or a *faset* in 3D.
- Conditions distributed through the body will be applied to an *elset*.

Line or face set surface conditions can be difficult to get right the first time because the sense of the set must be well defined.

When it is desired to impose a condition to a single node or to a single element, the node or element number may always be substituted for an nset or elset.

There are also pre-existing nsets and elsets for every node or element in the mesh. These sets are named respectively ALL\_NODE and ALL\_ELEMENT.

\*\*\*\*calcul
 \*\*\*bc
 \*\*impose\_element\_dof

## \*\*impose\_element\_dof

### **Description**:

This command is used to impose degrees of freedom which exist at element integration points. These DOFs are commonly part of mixed formulations, such as pressure-displacement, and plane-stress elements.

### Syntax:

\*\*impose\_element\_dof
 elset dof value table

### Example:

Making a 2.5 D case of planar displacements and an imposed  $\epsilon_{33}$  strain. formulation.

```
***mesh plane_stress
***bc
**impose_nodal_dof
bottom U2 0.
top U2 1. time
left U1 0.
**impose_element_dof 1 EZ 1.e-3 time
```

Element DOFs are also imposed for the RVE elements, with a full strain tensor being the element DOFs.

#### \*\*\*bc

```
**impose_element_dof
ALL_ELEMENT E12 1. time
```

# \*\*impose\_element\_dof\_reaction

### **Description**:

This command imposes the conjugate reaction to an element degree of freedom.

#### Syntax:

The syntax takes an element set name, the name of the degree of freedom for which reaction will be fixed, a scale value, and a table name (required for non-zero value).

\*\*impose\_element\_dof\_reaction
 elset dof value table

### Example:

The following example apples mixed mode loading to an RVE element with the impose\_element\_dof\_reaction conditions directly imposing the stress. This demonstrates the use of an element number for the element set. Note that E22 is given to fix  $\sigma_{22}$ . This is because the reaction is specified by the degree of freedom name, which is the strain in RVE elements.

#### \*\*\*bc

```
**impose_element_dof
1 E33 0.0
**impose_element_dof_reaction
1 E22 25. time
1 E11 400. time
```

# \*\*impose\_elset\_dof

## **Description**:

The boundary condition allows one to impose the value of a degree of freedom over an element set. Principally this condition applies to the elements  $2_5D$  (see **\*\*\*mesh**). The DOFs imposed by this condition must be located at element integration points (Gauss points), and not nodal unknowns.

# Syntax:

\*\*impose\_elset\_dof
 name\_elset dof\_name value name\_table

- name\_elset character name of the element set. This must be the name of a valid elset defined in the geometry file. The DOF will be imposed at every Gauss point in this element set.
- dof\_name The character name of the degree of freedom to be imposed. For the 2\_5D elements the choices are t1 t2 t3 w1 w2 w3.

value base value which scales table values (real).

table character name or list of names for the tables to be used (see **\*\*\*table**).

# Example:

```
**impose_elset_dof
    str t1 .01 tab1
    str t2 0.
    str t3 0.
    str w1 0.
    str w2 0.
    str w3 0.
```

# \*\*impose\_elset\_dof\_reaction

#### **Description**:

This command allows application of a force on the group of nodes within an element set. Note that application of a force is *not* the same as application of a pressure.

# Syntax:

\*\*impose\_elset\_dof\_reaction name\_elset dir value name\_table

name\_elset character name of the element set within which the reaction will be applied.

*dir* real value for the direction of application for the force.

*value* base value (real) for the condition. This value is a multiplier of the current table value thereby establishing the magnitude of the condition.

table character name for a valid loading table input with the command **\*\*\*table**.

# Example:

\*\*impose\_elset\_dof\_reaction
ALL\_ELEMENT w1 1. tabmx
ALL\_ELEMENT w2 1. tabmy

# \*\*impose\_nodal\_dof

#### **Description**:

This boundary condition imposes degrees of freedom located at nodes to defined values in time. The condition is general and therefore applies to *all* types of DOF for all types of problem.

# Syntax:

The syntax required to impose nodal DOFs is:

\*\*impose\_nodal\_dof
 nset\_name dof\_name value table\_name

- nset\_name This is the name of a valid node set (nset) which gives all the nodes where the DOF is to be imposed.
- dof\_name The character name of the DOF to be imposed. This must be one of the defined DOF types given by the problem, and as indexed in the appendix. DOF names are also listed in the *problem.ut* reference file.
- *value* This real value acts as a multiplier on the current applied load table value. If the value is zero, no table will be required.
- table\_name A character name for the loading table which describes the DOF value in time. A corresponding table must therefore be given elsewhere (see the procedure **\*\*\*table**).

#### Example:

Supposing that a mesh has been created with a node set composed of a single node, node 1. This example displaces that node a fixed amount in the directions  $u_1$  and  $u_2$ . The  $u_1$  magnitude will be 1.2 times the table value specified by table3 while  $u_2$  will be only 0.2 times that value. Using the same table assures synchronization between the loadings.

\*\*\*bc

```
**impose_nodal_dof node1 U1 1.2 table3
**impose_nodal_dof node1 U2 0.2 table3
```

Note that a structure will normally have more displacement conditions than the fixing of a single node. Other conditions such as pressure and force are also likely.

The same command may be used to specify the temperature in a thermal problem.

\*\*impose\_nodal\_dof node1 TP 200.0 tab1

#### <u>Note</u>:

Some variants exist, which allow the *value* to vary in space:

\*\*impose\_nodal\_dof
 nset\_name dof\_name function function(x,y,z); table\_name
 nset\_name dof\_name file binary\_file table\_name
 nset\_name dof\_name ascii\_file ascii\_file table\_name

The function variant allows the DOF value to depend on nodal coordinates. The function may also depend on time; this is however discouraged, because in that case incremental values are not properly computed. The time dependency should rather be specified through the table.

Both file variants read the value of each DOF from a file; thus the file should contain as many entries as nodes in the nset, values being ordered as in the nset. Note that the binary form expects entries as "floats" (not "doubles").

# \*\*impose\_nodal\_dof\_rate

#### **Description**:

This boundary condition imposes degrees of freedom located at nodes on a rate basis. The loading table will be scaled by the base value at any time to give the loading rate.

#### Syntax:

The syntax required to impose nodal DOF rates is:

\*\*impose\_nodal\_dof\_rate
 nset\_name dof\_name value table\_name

- nset\_name This is the name of a valid node set (nset) which gives all the nodes where the DOF is to be imposed.
- dof\_name The character name of the DOF to be imposed. This must be one of the defined DOF types given by the problem, and as indexed in the appendix. DOF names are also listed in the *problem.ut* reference file.
- *value* This real value acts as a multiplier on the current applied load table value. If the value is zero, no table will be required.
- table\_name A character name for the loading table which describes the DOF value in time. A corresponding table must therefore be given elsewhere (see the procedure **\*\*\*table**).

#### Example:

This example is for an axisymmetric specimen with applied hydrostatic pressure of 52 MPa, followed by strain rate controlled compression cycling along one axis, while the hydrostatic pressure is maintained. Note the use of zero in \*dtime to apply step loading of the strain rate. Expiring (exp) is also applied to the DOF rate condition because the DOF is not active at all times.

```
***bc
```

```
**pressure
    press
              -1. ptab
          exp -1. ptab2
    top
  **impose_nodal_dof
             U2 0.0
    v=0
             U1 0.0
    x=0
  **impose_nodal_dof_rate
    v=1
        exp U2 1.0 seg
***table
  **name ptab
    *time 0.0
                100. 5000.
    *value 0.0
                52.0 52.0
  **name ptab2
    *time 0.0
                100.
    *value 0.0
                52.0
  **name seg
```

\*\*\*\*calcul
\*\*\*bc
\*\*impose\_nodal\_dof\_rate

\*dtime 100.0 0.0 3.1579e+01 0.0 7.8947e+01 0.0 5.5263e+01 \*value 0.0 -3.8e-05 -3.8e-05 3.8e-05 3.8e-05 -3.8e-05 -3.8e-05

# \*\*impose\_nodal\_reaction

#### **Description**:

This option sets the "force" reactions which act on a node set. Note that applying a force to a node set is very different than applying a pressure (using the BC **\*\*pressure**). The condition is general so it may be used for all types of problem. For the simple mechanical case, the nodal reactions are in units of force.

# $\underline{Syntax}$ :

\*\*impose\_nodal\_reaction nset dir val table

- nset Character name of the node set where the condition is applied.
- *dir* Direction in which the reaction is applied. This will be the DOF name whose associated reaction is affected. See the appendix for DOF names.
- val Real number for the multiplicative base value of the condition.

table Character table name of a valid table, or a list of character table names.

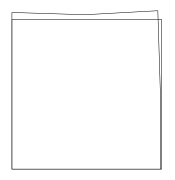
# Example:

The current example demonstrates the incorrect use of this option to impose a surface pressure. The difference is a common misconception of boundary conditions in the FEM. The problem uses a plane-stress eight node square element with fixed U1 on the left border and fixed U2 on the bottom. Material behavior is linear elastic.

The top node set is defined using the right top, middle top, and left top nodes in sequence. As the square has unit sides, one could suppose that the three forces applied should be 1/3 the stress. The BC is therefore applied as:

```
***bc
    **impose_nodal_reaction
    top U2 0.3333 tab
# **pressure
# press 1. tab
```

which results in the following distorted deformation:



Proper application of a pressure requires knowledge of the surface integration along the boundary. The options **\*\*pressure** and **\*\*shear** must therefore be used.

# $**impose\_nodal\_reaction\_rate$

#### **Description**:

This is an associated reaction version of **\*\*impose\_nodal\_dof\_rate**. The condition is useful when the initial reaction state is unknown, which can be generated with a mix of displacement and reaction BCs using the **exp** keyword.

# Syntax:

The syntax is analogous to  $\texttt{**impose_nodal_dof_rate}.$ 

\*\*impose\_nodal\_reaction\_rate
 nset\_name dof\_name value table\_name

# \*\*impose\_nodal\_dof\_density

#### **Description**:

The command imposes the flux (associated reaction) density across a surface (nset or faset). It is a general boundary condition, which can mean imposed pressure in the X-Y-Z directions for mechanical problems. The condition **\*\*fluconv** is a special case of this condition as well.

Note that the **\*\*pressure** command adjusts itself for **updated** elements according to the surface normal, and for 3D meshes there is no **\*\*shear** option because the face tangent is not fixed. This option differs because it is related to the density of the DOF reactions and therefore rests in their coordinate space.

# Syntax:

\*\*impose\_nodal\_dof\_density
 bset dof value table

where *bset* is a boundary set name (liset or faset), *dof* is a degree of freedom name (e.g. U1), *value* is a scaling value (decimal point number), and *table* is the name of a loading table input elsewhere.

# Example:

Pressure defined in the U1-U2 directions.

```
***pc
```

```
**impose_nodal_dof
left U1 0.0
bottom U2 0.0
fix U3 0.0
**impose_nodal_dof_density
face.2 U1 0.25 tab1
face.2 U2 0.25 tab2
```

# \*\*release\_nodal\_dof

#### **Description**:

This command allows a time-based progressive release of a fixed condition. The crack length is given by the loading table of the condition. Using a function or table of values one can specify complex crack growth rates. For example crack growth measurements can be input as a table, and the crack propagated as actually seen. Then loading parameters such as  $\Delta K$  or maximum crack tip opening can be correlated to the growth.

#### Syntax:

\*\*release\_nodal\_dof nset dof base-val table

#### <u>Note</u>:

The crack length is measured as the distance sum of distances between nodes with the initial tip being at the first node of the given node set (i.e. its the arc length s of the line drawn by the node set). Curved crack paths are therefore possible. The node set should be ordered beforehand.

# \*\*submodel

#### **Description**:

Assume that the current computation is running on a part of a bigger problem already solved. we call *submodel* the current part and *Master* the bigger problem. This boundary condition imposes degrees of freedom located at boundary nodes in a submodel to values computed from the Master. The submodel boundary nodes have not to match with Master's one, this Bc is able to locate them in the Master mesh and interpolate their values from Master's dofs. The condition is general and therefore applies to *all* types of DOF for all types of problem.

# Syntax:

×

The syntax of submodel is:

**submodel
[*format format]
*global_problem Master_name
*dofs dof_1_name dof_2_name dof_n_name
*driven_nsets nset_name

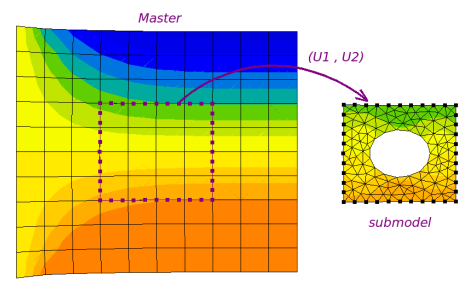
format This is the results format of the Master computation. If **\*format** is omitted, the default Z7 format is assumed.

Master\_name The name of the Master computation.

- dof\_i\_name The name of the DOF to be imposed. This must be one of the defined DOF types given by the *Master* and *submodel* problems. DOF names are also listed in the *problem*.ut reference file.
- nset\_name This is the name of a valid node set (nset) which gives all the *submodel* nodes where the DOFs are to be imposed.

# Example:

This example can be seen in test database, we use the displacement U1, U2 on a simple square to drive a submodel with a hole.



\*\*\*\*calcul \*\*\*bc \*\*submodel

\*\*\*bc
 \*\*submodel
 \*format Z7
 \*global\_problem master
 \*dofs U1 U2
 \*driven\_nsets external

# \*\*K\_field

#### **Description**:

This boundary condition is used to impose on a NSET the displacement field given by the linear solution at a crack tip. The crack is loaded under mode I or II. This boundary condition is limited to 2D plane problems. It will work on axisymmetric problems , however its physical meaning will be uncertain.

The dispalcement field is given by:

Mode I:

$$u_1 = \frac{K_I}{2\mu} \sqrt{\frac{r}{2\pi}} \cos\frac{\theta}{2} \left(\kappa - 1 + 2\sin^2\frac{\theta}{2}\right)$$
$$u_2 = \frac{K_I}{2\mu} \sqrt{\frac{r}{2\pi}} \sin\frac{\theta}{2} \left(\kappa + 1 - 2\cos^2\frac{\theta}{2}\right)$$

Mode II:

$$u_1 = \frac{K_{II}}{2\mu} \sqrt{\frac{r}{2\pi}} \sin\frac{\theta}{2} \left(\kappa + 1 + 2\cos^2\frac{\theta}{2}\right)$$
$$u_2 = -\frac{K_{II}}{2\mu} \sqrt{\frac{r}{2\pi}} \cos\frac{\theta}{2} \left(\kappa - 1 - 2\sin^2\frac{\theta}{2}\right)$$

with

$$\kappa = \frac{3 - 4\nu}{(3 - \nu)/(1 - \nu)}$$
 plane strain plane stress

where  $\nu$  is the Poisson's ratio. Let M be the point where the displacement field is computed and T the crack tip.  $r = ||\vec{MT}||, \theta = (\vec{Ox_1}, \vec{MT}).$ 

#### Syntax:

The syntax is as follows: \*\*K\_field nset center young poisson plane\_state mode value [table] nset name of the NSET center crack tip position (2D vector) young Young modulus (double) poisson Poisson coefficient (double) plane\_state plane\_stress or plane\_strain mode specifies loading mode: I or II value basic boundary condition value (double). This represents the basic value of  $K_{I,II}/2\mu$ . table table name

#### Example:

\*\*\*\*calcul \*\*\*bc \*\*K\_field

\*\*K\_field border % nset name (0.,0.) % crack tip position 0.3 % Poisson's ratio plane\_stress II % mode II 10. % basic value Ktab % table name

# \*\*T\_field

# **Description**:

This boundary condition is used to impose on a NSET the displacement field given by the linear solution at a crack tip of and applied T–field.

Under plane strain conditions, the displacement field is given by:

$$u_1 = T \frac{1 - \nu^2}{E} R \cos \theta$$
$$u_2 = -T \frac{\nu(1 - \nu)}{E} R \sin \theta$$

where  $\nu$  is the Poisson's ratio, E the Young modulus.

# Syntax:

```
The syntax is as follows:

**T_field nset center young poisson value [table]

nset name of the NSET

center crack tip position (2D vector)

poisson Poisson coefficient (double)

young Young module (double)

value basic boundary condition value (double). This represents the basic value of T.

table table name
```

# Example:

# \*\*centrifugal

#### **Description**:

The centrifugal boundary condition is used to apply body forces  $\rho\omega^2 r$  to a defined set of elements (elset) due to rotational acceleration, with  $\rho$  the volumetric mass,  $\omega$  the rotational rate and r the distance to the rotational axis.

#### Syntax:

\*\*centrifugal [updated] [square]
elset\_name (v) dir value table ... [tableN]

- *elset\_name* Character name for the set of elements which are subjected to the centrifugal loading.
- updated Uses the current configuration rather than the initial configuration for determining the distance r to the rotational axis as well as the direction of the centrifugal force.
- square Applies a force proportional to the rotational rate  $\omega$  instead of  $\omega^2$ .
- v Vector giving a point position through which the rotational axis passes. Axisymmetric problems require that the origin is input as  $(0.0 \ 0.0)$ .
- dir Direction of the rotational axis (d1, d2, or d3). Axisymmetric 2D problems are required to use the d2 axis direction. Planar 2D geometries require that the axis is d3.
- value Base value (real) which scales a table value to determine the applied rotational rate  $\omega$ .
- table Character name for a valid loading table or tables which, after multiplication with value, will describe the time evolution of the rotational rate.

To apply a centrifugal force, it is required to give the volumetric mass  $\rho$  for the material in the material file. This is specified with the **\*\*\*coefficient** command.

Units required depend of course on the consistency of units within the problem. For example, if the dimensional unit is millimeter, the forces are given in Newtons, and the rotational rate  $\omega$  is in radians per second, the volumetric mass must be in tons (metric)/mm/mm/mm.

#### Example:

```
**centrifugal
str (0.0 0.0) d2 1.e7 tab1
```

# \*\*crack\_release

#### **Description**:

This BC is used to propagate a crack (release DOFs) according to node-extrapolated material values (note there can be some significant error due to the extrapolation error, especially with the high variable gradients found around crack tips).

# Syntax:

\*\*crack\_release nset dof var-value-list

where *nset* is the node set defining the ligament through which the crack propagates, and *dof* is the fixed DOF name (e.g. U2). Following these entries, a list of variable-value combinations are to be entered to specify the crack growth criterion.

# Example:

For a plasticity problem a maximum allowable plastic strain equivalent of 20% could be used (remember the name of p - here epcum depends on the material law used).

# \*\*crack\_release middle U2 epcum 0.20

Or one could use a secondary material variable such as the von Mises stress as a ultimate tensile strength.

```
**crack_release
    ligament U2 sig::mises 520.0
    ligament U1 sig::mises 520.0
```

# \*\*deformation

#### **Description**:

This boundary condition imposes the displacement vector  $\vec{v}$  of a node by the following formula:

 $\vec{u} = \boldsymbol{\epsilon} \vec{r}$ 

where  $\epsilon$  is a symmetric tensor, and  $\vec{r}$  the position vector of the node considered. The origin of  $\vec{r}$  may be a fixed point or a node of the mesh.

For the case where the origin is a fixed point the node set considered must be given, as well as the position of the origin point in the form of a vector, and the components of the tensor  $\epsilon$ . In 1D  $\epsilon_{11}$  et  $\epsilon_{22}$  may be imposed; in 2D  $\epsilon_{12}$  may be added; in 3D the remaining  $\epsilon_{33}$ ,  $\epsilon_{13}$ , and  $\epsilon_{23}$  terms may be added. The tensor components are respectively named E11, E22, E12, E33, E13, and E23. If a component is not defined its value is set to zero. The values may be set by using a table for all of the components, or a table for each individual component (only for non-zero components).

# Syntax:

```
**deformation [ node ]
    nset_name (vec | num_node)
    compos_name value [ table ]... compos_name value
[ table ]
```

node is an optional keyword indicating that the origin of  $\vec{r}$  is a node of the mesh.

nset\_name Character name of the nset group upon which the displacement will be imposed.

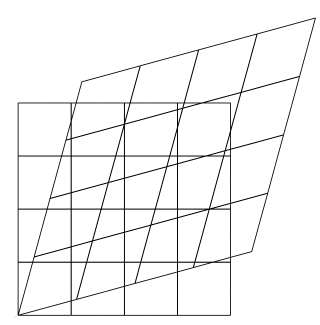
- vec Position of the origin point in the form of a vector. This is in the case where the center is not a node of the mesh. Recall the vector form is 2 or three real values enclosed in parenthesis such as (1.0.0.).
- num\_node Integer number of a node in the case where the node keyword was given.
- compos\_name value Character string giving the name of a component of the  $\epsilon$  tensor as defined above, followed by a real giving the components value. Any number of these components may be given in any order thereby specifying the deformation tensor.
- table character name of the table(s) used to describe the field amplitude. See the command \*\*\*table.

# Example:

This example shows a simple uniform deformation field to an initially square multi-element mesh. The required BC input data is the following (see test deform in Z7test/Static\_test/):

```
**deformation
boundary (0. 0.) E22 0.1 E11 0.1 E12 .3 tab
```

where **tab** is a unit table definition given after. The resulting deformed structure is displayed below. This type of condition is useful for testing the mechanical behavior of materials under complex strain loadings.



Another imposed strain field is given below:

\*\*deformation node
 nsetdef 2 E11 2. E33 4. tab

# \*\*free\_rotation

#### **Description**:

This rotation condition allows a boundary to be rotated about one of the coordinate axes (2 displacements are imposed), while the DOFs along the third axis remains free. The usefulness of this condition is obviously only for 3D problems.

#### Syntax:

The syntax is the following:

\*\*free\_rotation nset dir [origin] value table

nset Name of the node set upon which the condition is active.

- *dir* an axis about which the node set will rotate (d1 d2 or d3). The DOF acting along the given direction is free in the absence of other conditions acting on it.
- origin specifies an arbitrary point on the rotation axis (use vector notation, see 1.3); it defaults to  $(0. \ 0. \ 0.)$ .

value Base value (scaling factor, decimal number).

table Name of a valid loading table.

# \*\*linear\_free\_rotation

# **Description**:

A linearized version of the **\*\*free\_rotation** condition for use in linear small deformation analysis. See page 3.67 for more discussion on this.

# \*\*gravity

# **Description**:

The gravity condition applies a body force due to uniform acceleration such as gravity.

To apply a gravitational force, it is required to give the volumetric mass for the material in the material file. This is specified with the **\*\*\*coefficient** command.

# Syntax:

The gravitational BC is specified with the following syntax:

**\*\*gravity** elset\_name dir value [table]

- *elset\_name* Character name for the pre-defined element set where the gravitational acceleration is applied.
- dir Direction of the gravitational force (d1, d2 or d3). Axisymmetric 2D problems require that the direction be d2. Plane 2D geometries require that the direction is d1 or d2.
- *value* Real base value scaling the current table magnitude to determine the acceleration magnitude.
- table Character name for a valid loading table or tables which will describe the acceleration in time.

Units required depend of course on the consistency of units within the problem. For example, if the dimensional unit is millimeter, the forces are given in Newtons, and the gravitational acceleration is in mm/sec/sec, the volumetric mass must be in tons (metric)/mm/mm.

# Example:

\*\*gravity
str d2 -20.0 tab1

# \*\*hydro

#### **Description**:

This boundary condition is used to impose a surface pressure calculated based on the updated (deformed) geometry, i.e. based on the geometry at the *end* of each increment (compare this to **\*\*\*bc \*\*pressure** at page 3.68). Hydro pressure will be defined according to the geometrical normal of a line set in 2D or a face set in 3D.

The sense of the normal with respect to the plane will of course be determined by the order of the node definitions in the surface definitions. For 2D problems if one walked along the nodes in the order of the **liset** command definition, the normal will be oriented to the right hand side. By convention, the normal is oriented outwardly from a closed body. In 3D geometries, this convention will be assured by using the standard Z-set meshing utilities.

The sign convention for the pressure is in the sense of mechanics and not hydrolics. That is to say that the pressure sign will be positive for pressure in the direction of a surface normal. A cube submerged in fluid will therefore have negative pressure. Note that if the normal is defined in a non-standard sense which is toward the interior of the body the sign of the pressure will be reversed.

#### Syntax:

\*\*hydro bset value [table]

- bset This is the character name of a boundary set, i.e. a liset for 2D problems or a facet in 3D problems.
- *value* Real value for the base or scaling value of the hydrostatic pressure. The table value is scaled by this factor to calculate the pressure at a given time.
- table Character name for a valid loading table or tables which will describe the pressure in time.

# Example:

\*\*hydro face1 -100.0 tab1

#### \*\*impedance

#### **Description**:

The impedance boundary condition can be useful, for example, for modeling the presence behind an interface of a material having different wave propagation properties. It can be used to tune the part of reflexion and transmission of an incident wave on an interface. The characteristic impedance Z of a medium is a material property defined as :

$$Z = \rho c$$

where  $\rho$  is the density of the medium and c is the longitudinal wave speed.

The impedance bc is a Robin condition that links the stress on the boundary with the velocity :

$$\underline{\sigma}\underline{\vec{n}} = Z\left(\vec{v}.\vec{n}\right)\vec{n} \quad \text{on } \Gamma_i$$

Where  $\vec{n}$  is the unitary outward normal vector to the boundary  $\Gamma_i$ .  $\underline{\sigma}$  and  $\vec{v}$  are the stress tensor and the velocity respectively. Numerically, impedance is handled through additional terms in the damping matrix (see calcul dynamic page 3.10).

A classical situation where an impedance bc is needed is the hopkinson experiment. An input bar impacts a sample from a side, a wave is generated and propagates though the sample in contact with an output bar on the other side. To simulate the sample/output bar interface without meshing the output bar, an impedance bc can be used with Z equal to the output bar characteristic impedance.

Here are two special cases to illustrate the role of the impedance in the reflexion/transmission behavior of a wave at the interface between two media. Lets take two media having impedances  $Z_1$  and  $Z_2$  respectively and lets consider a wave coming from material 1:

- if  $Z_2/Z_1 = 0$  (medium 2 is void for instance), the wave is totally reflected
- if  $Z_2/Z_1 = 1$ , the wave is totally transmitted to medium 2

#### Syntax:

\*\*impedance bset value

Where *bset* is the name of the bset where the impedance bc is applied and *value* is the impedance factor.

#### Example:

```
***bc
**impedance
interface 40.e6
```

# \*\*linear\_rotation

#### **Description**:

This condition is a linearized version of the **\*\*rotation** boundary condition (page 3.72). For small deformation problems (not updated), this condition should be used.

This version uses the approximation that angles will be small, so  $\sin \theta \approx \theta$ . Large rotations will of course result in an expansion of the boundary.

# Syntax:

The syntax is the same as for **\*\*rotation**. For 2D problems we have:

```
**linear_rotation [ node ]
    nset_name ((origin) | num_node )
    angle [ table ]
```

For 3D problems, the syntax is slightly changed:

```
**linear_rotation [ node ]
    nset_name [ num_node ] (axe)
    [ (origin) ] angle [ table ]
```

See the discussion on page 3.72 for more detail on the input structure.

# \*\*pressure

#### **Description**:

The pressure mechanical boundary condition is used to impose a surface pressure normal to a valid surface defined by the liset or faset geometry commands. The sense of the normal with respect to the plane will of course be determined by the order of the node definitions in the surface definitions. For 2D problems if one walked along the nodes in the order of the liset command definition, the normal will be oriented to the right hand side. By convention, the normal is oriented outwardly from a closed body. In 3D geometries, this convention will be assured by using the standard Z-set meshing utilities.

Pressure calculation is always calculated based on the *initial geometry* for small deformation/small displacement formulations, and on the *updated* (deformed) geometry at the *start* of the increment for small deformation/large displacement formulations In *large deformation* problems therefore this option is not convenient for the fluid pressure on a surface. The **\*\*hydro** command (page 3.65) may be used to apply a pressure on the deformed surface to simulate hydrostatic fluid pressure.

The sign convention for the pressure is in the sense of mechanics and not hydrolics. That is to say that the pressure sign will be positive for pressure in the direction of a surface normal. A cube submerged in fluid will therefore have negative pressure. Note that if the normal is defined in a non-standard sense which is toward the interior of the body the sign of the pressure will be reversed.

#### Syntax:

\*\*pressure liset value table ... [tableN] faset value table ... [tableN]

- liset Character name of the line set upon which pressure is applied in 2D problems.
- faset Character name of the face set upon which pressure is applied in 3D problems.
- value Base value (real number, function or file) which scales a table value to determine the pressure magnitude.
- table Character name for a valid loading table or tables which will describe the pressure in time.

# \*\*pressure\_from\_function

#### **Description**:

This boundary condition extends **\*\*pressure** (3.68): the pressure may depend on time and coordinates.

# Syntax:

\*\*pressure\_from\_function
\*nset nset\_name
\*function function(t,x,y,z);

nset\_name is the name of a valid node set (nset) where the pressure is imposed. Note that the corresponding best *must* also exist (and have the correct orientation).

function specifies a function, used to compute the pressure value.

# Example:

From \$Z7TEST/Static\_test/INP/bc\_pressure\_from\_function.inp:

```
**pressure_from_function
*nset top-ext
*function 1.+ (x/50.);
```

# Note:

```
The syntax
```

\*\*pressure top-ext function 1.+ (x/50.); my\_table
may also be used, if table loading desired.

# \*\*radial

# **Description**:

This boundary condition allows imposing radial displacements which simulate the expansion or contraction around a group of nodes. An example is the shrinking or expanding of tubes.

# Syntax:

\*\*radial nset\_name origin direction value [table]

nset\_name Character name of the nset where the condition is applied.

- origin Vector form for the origin where the expansion axis passes. A vector form will give the two or three coordinates necessary in parenthesis. An example is (1. 0. 1.).
- direction Direction of the axis of expansion. These directions must coincide with the problem coordinate axis, and are defined with the tokens d1, d2, and d3.
- *value* Real number giving the base value for the conditions magnitude. A positive value results in expansion and negative contraction.
- table Name of a table describing the load magnitude through time (see **\*\*\*table**).

# Example:

\*\*radial str (0.0 0.0) d3 3.0 tab1

In 2D the expansion axis is necessarily d3.

# \*\*radius

#### **Description**:

The radius condition imposes an indentation profile of constant radius. The condition is valid only in 3D mechanical problems with the radius profile in the U2 direction.

The radius profile is applied as the inverse of the radius during deformation. This allows a zero radius (flat surface) to be deformed in sequence to the given radius. The surface is therefore tightened into the radial form. The exact expression used to calculate the radius at the increment i between the start of a sequence s and the end e is:

$$\frac{1}{R_i} = \frac{1}{R_s} + \frac{i}{N_{inc}} \left( \frac{1}{R_s} - \frac{1}{R_e} \right)$$

with  $N_{inc}$  the number of increments in the sequence.

If the radius value is positive, the center of the deformation profile will be located in the negative z space. The inverse statement is of course true as well.

# Syntax:

The radial condition takes the following syntax:

```
**radius nset_name value [table]
```

- nset\_name The character name of the node set upon which the condition is applied. This node set must be defined in the plane z = 0.
- value Base value (real) for the radius. The base value is a multiplicative scale for the table values at all times.
- table Character name for the table which describes the condition's magnitude.

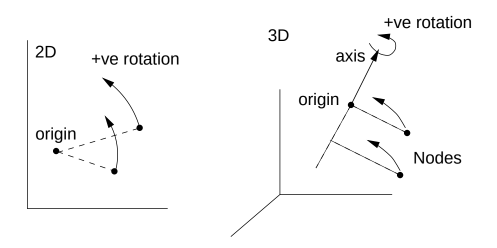
# Example:

\*\*radius
 surface 100.0 tab1

# \*\*rotation

#### **Description**:

This condition rotates all or part of a structure about (1) a fixed point, (2) a node of the mesh. This condition functions in 2D or 3D mechanical problems. For the 2D case, the rotation is always about the third direction axis.



# Syntax:

The syntax for 2D problems is the following:

```
**rotation [ node ]
    nset_name ((origin) | num_node )
    angle [ table ]
```

For 3D problems, the syntax is slightly changed:

```
**rotation [ node ]
nset_name [ num_node ] (axe)
[ (origin) ] angle [ table ]
```

- **node** Optional keyword which specifies that the condition's center of rotation will be defined about a node of the mesh.
- nset\_name Name of the node set upon which the condition is active.
- origin Origin of rotation from which the displacement field is calculated. This point is given using vector syntax.
- *num\_node* Integer node number defining the center of rotation in the event that the **node** keyword was given.
- angle Real value in degrees acting as the multiplicative factor for the condition magnitude. At any time, the rotation angle will be this value times the given table value.
- table Name of a valid loading table.

axe Vector giving the axis of rotation. Note that the Cartesian axes are defined by X:(1. 0. 0.), Y: (0. 1. 0.), Z: (0. 0. 1.).

# Example:

Several examples of the rotation syntax are given below. These each give the corresponding test name user the **Static\_test** directory.

% ztole3m \*\*rotation bord (0. 0.) 360.0 tab1

% hook\_ld2
\*\*rotation node sommet 1 180. tab

% cuberot \*\*rotation rot (1. 0. 0.) (0. 0. 0.) 360. table

# \*\*shear

#### Description:

The shear BC imposes a tangential pressure along a pre-defined line set in an analogous fashion as in the **\*\*pressure** command. it is impossible to impose a shear along a face set, thereby limiting its applicability to 2D problems.

# Syntax:

\*\*shear liset value table ... [tableN] ...

*liset* Character name of the line set where the shear pressure is applied.

value Real base or scaling value for the condition.

 $table\$  Character name for a valid loading table or tables which will describe the pressure in time.

# Example:

This example is presented as a useful template for testing material behavior in shear. The single square element is defined as follows:

```
***geometry
**node 8 2
                0.000000
 1 0.0000000
 2 1.0000000
                0.000000
 3
    1.0000000
                1.0000000
    0.000000
 4
                1.000000
 5
    0.5000000
                0.000000
    1.0000000
                0.500000
 6
    0.5000000
 7
                1.0000000
 8 0.000000
                0.500000
**element
          1
 1
     c2d8
             15263748
***group
**liset s1
               1
                     5
                           2
 quad
               3
                     7
                           4
 quad
**liset s2
 quad
               2
                     6
                           3
               4
                     8
                           1
 quad
*nset bottom
 1 5 2
**nset corner
 1
***return
```

\*\*\*\*calcul \*\*\*bc \*\*shear

A shear stress loading may be applied with the following boundary conditions in the input file:

# \*\*\*bc \*\*shear s1 exp 1. tab s2 exp -1. tab \*\*impose\_nodal\_dof bottom U2 0.0 corner U1 0.0

# $**static_torsor$

#### **Description**:

The static\_torsor BC imposes a static torsor (resultant and momentum) on a nset. This means that the sum of the action of every nodal forces of the nset is equivalent to the specified static torsor. It is an ill-posed problem since we add 3N unknowns (nodal forces in the three directions) and only 6 equations (3 equations for the resultant and 3 for the momentum). This BC is therefore associated to a rigid body motion of the nodes of the nset through a **\*\*mpc\_rb** relationship (see page 3.123). Note that the user doesn't have to define the **\*\*mpc\_rb** relationship, it is done automatically.

When a **\*\*mpc\_rb** is applied, only master 6 dofs over every dofs of the nset remain in the linear system. A small system of equations is then solved where the 6 remaining external forces associated to the 6 master dofs are computed.

#### Syntax:

**static_tors	r
*nset	nset
*resultant	vector
*momentum	vector
*point	vector
*table	value table_name

Where *nset* is the name of the nset where the torsor is applied. **\*resultant** and **\*momentum** define the resultant and the momentum at the point defined by the **\*point** command. *value* and *table\_name* define the multiplicative factor of the torsor.

#### Example:

```
***bc
 **static_torsor
  *nset
               top
  *resultant
               (5.
                       1.
                             0.)
  *momentum
               (0.01)
                       0.0
                             0.)
  *point
               (1.
                       0.5
                             0.5)
  *table
               1.
                   time
```

\*\*\*\*calcul
 \*\*\*bc
 \*\*strain\_gradient

# \*\*strain\_gradient

# **Description**:

The syntax is as follow :

# Syntax:

\*\*strain\_gradient nset dof value table

# \*\*surface\_heat\_flux

#### **Description**:

The **\*\*surface\_heat\_flux** condition is used to impose a surface heat flux on a line set in 2D or a face set in 3D (i.e. a Neumann boundary condition).

This keyword replaces the now deprecated keywords **\*\*flucons** and **\*\*surface\_heat**.

# Syntax:

\*\*surface\_heat\_flux liset value table faset value table

- liset Character name for the line set upon which the heat flux is applied in 2D.
- faset Character name for the face set upon which the heat flux is applied in 3D.
- *value* Real base value for the heat flux. This value scales the current table value to obtain the heat flux magnitude.
- table Character name of a pre-defined loading table or list of tables (see the option **\*\*\*table** p. 3.199). The table value determines the exterior temperature.

# Example:

\*\*surface\_heat\_flux
myliset -10. table1

# \*\*convection\_heat\_flux

#### **Description**:

The **\*\*convection\_heat\_flux** BC imposes a convective heat flux on a line set in 2D or a face set in 3D.

 $q_c = a(T - T_e)$ 

This keyword replaces the now deprecated keyword **\*\*fluconv**.

#### Syntax:

\*\*convection\_heat\_flux group
 h a
 Te Te table

- group Character name for the group of line sets in 2D or face sets in 3D where the flux condition is to be applied.
- a Real value for the coefficient a.

Te Real value for the exterior temperature  $T_e$ .

table Character name of the table describing the flux value in time (see **\*\*\*table**).

The coefficient *a* may depend on the temperature for this condition. The syntax required for a convection heat flux condition with this dependence is given below:

```
**convection_heat_flux group
   h temperature
   h0 T0
   h1 T1
   h2 T2
   ...
   Te Te table
```

The keyword **\*\*convection\_heat\_flux** and the two following lines must be repeated as many times as there are convective flux boundary conditions.

# Example:

```
**convection_heat_flux myliset
    h 100.0
    Te -10. table1
```

# \*\*interface\_heat

# **Description**:

The **\*\*interface\_heat** BC option is used to impose an inter-facial heat transfer with thermal resistance between two line set boundaries in 2D, or two face set boundaries in 3D. The inter-facial flux will be calculated as:

 $flux = a(T_1 - T_2)$ 

where  $T_1$  and  $T_2$  are the temperature at each side of the interface.

The two interfaces must be in geometrical correspondence, which may be summarized as follows:

- A line or face at one side of the interface must correspond to exactly one line or face on the other side of the interface.
- The nodes in the lines or faces in correspondence must be paired exactly one to one. For this to be true, it is necessary that the distance between two corresponding nodes be inferior than the distance  $\epsilon$ . In result of this requirement is the fact that all nodes which are *not* in correspondence have a separation distance greater than  $\epsilon$ .

This keyword replaces the now deprecated keyword **\*\*fluconv\_interface**.

# Syntax:

\*\*interface\_heat group1 group2  $\epsilon$ h a

group1 Character name for the first line set group in 2D, or face set group in 3D.

group2 Character name for the second line set group in 2D, or face set group in 3D.

- $\epsilon$  Real value for the critical distance between nodes.
- a Base value (real) for the value of the coefficient a.

The coefficient a may depend on the temperature for this condition. The syntax required for an inter-facial convection resistance condition with this dependence is given below:

rface_heat group
temperature
T0
T1
T2
Te table

The keyword **\*\*interface\_heat** and the two following lines must be repeated as many times as there are convective flux boundary conditions.

# Example:

```
**interface_heat liset1 liset2 0.001
```

```
h 100.0
```

# \*\*volumetric\_heat

#### **Description**:

This BC is used to impose a volumetric heat flux within a given element set. This keyword replaces the now deprecated keyword **\*\*fluvol**.

#### Syntax:

**volumetric_heat		
elset value table		

- *elset* Character name for the element set (elset) within which the heat generation is applied.
- *value* Real base value for the heat flux. This value scales the current table value to obtain the heat flux magnitude.
- table Character name of a pre-defined loading table or list of tables (see the option **\*\*\*table** p. 3.199). The table value describes the magnitude of the flux in time.

### Example:

\*\*volumetric\_heat
 str 20.0 tab1

# $**volumetric\_heat\_from\_parameter$

#### **Description**:

This BC is used to impose a volumetric heat flux within a given element set. The time-space values are taken from a parameter field.

#### Syntax:

\*\*volumetric\_heat\_from\_parameter
 elset value parameter\_name

- *elset* Character name for the element set (elset) within which the heat generation is applied.
- *value* Real base value for the heat flux. This value scales the given parameter field to obtain the heat flux magnitude.

parameter\_name Name of the parameter.

### Example:

\*\*volumetric\_heat\_from\_parameter
 str 20.0 param1

# \*\*volumetric\_heat\_in\_file

#### **Description**:

The **\*\*volumetric\_heat\_in\_file** BC is used to impose a volumetric heat flux within a given element set, using input from a file.

This keyword replaces the now deprecated keyword **\*\*fluvol\_in\_file**.

#### Syntax:

\*\*volumetric\_heat\_in\_file
 elset value file-prefix

- *elset* Character name for the element set (elset) within which the heat generation is applied.
- *value* Real base value for the heat flux. This value scales the current table value to obtain the heat flux magnitude.
- file-prefix pre-fix for the input file. The first file of concern is file-prefix.catalog, which lists the initial and final times for the file inputs (presumably from another time step of a mechanical problem, see below). Binary files for the beginning of the problem, start of an increment and end of an increment are: file-prefix.first file-prefix.initial file-prefix.final The current increment should be within the time bounds given in the catalog file.

# Example:

```
***sub_problem fem MechTherm/plastic
 **transfer integ_nodeparam
 *variable q_dot
 *file heat_out
```

In thermal.inp the heat is imported using the current BC:

```
**volumetric_heat_in_file
ALL_ELEMENT 1.0 heat_out
```

An example of what's in the catalog file is:

Time\_ini 7.600000 Time 8.000000

### \*\*radiation

#### **Description**:

This BC for thermal problems applies a radiation heat flux on a line set (liset) in 2D, or a face set (faset) in 3D problems. The expression for the heat flux is:

 $flux = a(T^4 - Te^4)$ 

Syntax:

\*\*radiation group a Te table

group Character name of the previously defined line set in 2D or face set in 3D.

- a Real value for the coefficient a.
- Te Real value for the exterior temperature,  $T_e$ , scale factor. This value will be a multiplicative scale of the current table value to determine the external temperature.
- table Character name of a pre-defined table name (see **\*\*\*table**). This table describes the magnitude of the external temperature

The coefficient *a* is constant. There is currently no provision for temperature or time dependence in this parameter. We recall that the coefficient is the product of the two terms:

#### $a=\sigma\epsilon$

with  $\sigma$  the Stefan's constant (5.73  $10^{-8}$   $W/m^2$   $K^4$ ) and  $\epsilon$  the grey body constant (no dimension).

The formula of radiation is not valid except for temperatures defined in on the absolute scale (Kelvin). This requires therefore that the units throughout the problem be defined consistently in Kelvin when radiation boundary conditions are applied.

#### Example:

\*\*radiation myliset 1.0e8 273.0 table1

# $^{***} coupled\_resolution$

### **Description**:

This command is used to specify special controls relating to the convergence of weak coupled problems.

# Syntax:

\*\*\*coupled\_resolution
\*\*iteration iter

# Example:

An example control for a coupled problem follows (from \$Z7PATH/test/Coupled\_test/INP/MechTherm.inp):

```
***resolution
**sequence
*time 4.0 8.0
*increment 10 10
*ratio 1.e-4
***coupled_resolution
**iteration 2
```

# $***compute_G_by_gth$

### **Description**:

This command is used to calculate the energy release rate G along a 3D crack front using the the energetic G- $\theta$  method. Various options may be activated to:

- smooth the crack front by defining the number of points of the spline curve used to build the variational problem solved to obtain the energy release rate (G) and crack virtual extension ( $\theta$ ) at each control point of the underlying spline,
- output results in terms of conventional stress-intensity factors (SIF)  $K_I$ ,  $K_{II}$ ,  $K_{III}$ ,  $K_{III}$  instead of the default G values,
- define propagation laws based on the previous SIF calculations, that will generate crack advance values that may be used to drive crack propagation by means of the **\*\*\*auto\_remesh** command,
- define various crack bifurcation criteria for out-of-plane crack propagations.

Note that this command may be applied either to conforming cracks FE models (ie. the crack is explicitly introduced in the mesh, and the crack front is defined by a liset) or xfem ones (in this case the discontinuity is defined by means of level-sets in a **\*\*xfem\_crack\_mode** block of commands, see 3.205). Note also, that the input for this command may be built automatically by the interactive **Zcracks** and **Zxfem** commands.

# Syntax:

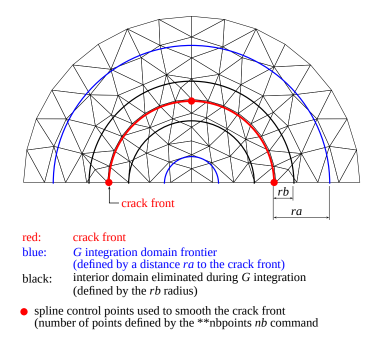
The syntax is as follows:

```
***compute_G_by_gth component-name
[ **crack_front liset-name ] | [ **xfem ]
    **nbnodes nbnodes
[ **theta ra [ rb ] ] | [ **elem_radius nbe ]
[ **elset elset_name ]
[ **exclude_BE ]
[ **lip lipname ]
[ **compute_K ]
[ **compute_K ]
[ **fatigue preload period ]
[ **behavior btype ]
[ **behavior btype ]
[ **belta_N deltan ]
[ **h hmax ]
[**optim_dir prec scan ]
[ **vectorial ]
```

The different option-keys are

- *liset-name* is the liset (line set of nodes) defining the crack front for conforming mesh modelizations. This liset is usually automatically created in the FE mesh by the Zcracks interactive script, and maintained during propagation analysis.
- alternatively the command **\*\*xfem** should be used for xfem models.

• as defined on the next figure, the parameter *nbnodes* (\*\*nbnodes command) defines the number of points on the spline curve built internally to smooth out the crack front. Note that a negative values can be given for the *nbnodes* parameter, in which case the number of spline control points is calculated as a fraction of the total number of crack front nodes (number of nodes in liset *liset-name* in the conforming mode).



- the **\*\*theta** command defines the size of the integration domain used to build the G- $\theta$  variational problem. As shown on the figure, this domain has the shape of a ring, centered around the crack front, where *ra* is the radius of the circular section of the ring. The optional parameter *rb* may also be entered, and correspond to the size of an inner volume, close to the crack front: in this case, the mechanical fields on the first rows of integration points/elements near the front are not considerered precise enough, and are eliminated from the integration domain. Note that in practice, the outer radius *ra* of the integration volume, should be chosen to stay within the volume (ie. it should not reach the outer free surface) to avoid erroneous  $G \theta$  solutions.
- the command **\*\*elem\_radius** is an alternative way to specify the size of the integration domain. In this case the outer radius *ra* is calculated to correspond to *nbe* times the average size of crack front elements.
- the optional command **\*\*elset** can be used to speed up the G- $\theta$  volume definition, ie. only candidate elements in elset *elset\_name* will be tested for addition in the volume integration.
- in general, values near free surfaces (at both ends of opened crack fronts) are less precise than inside the volume along the crack front. The optional command **\*\*exclude\_BE** may be used to reset those values, and calculate SIFs at the front ends by extrapolation of values obtained on inner points.
- command \*lip is not used in the xfem mode. *lipname* is a name of an nset in the FE mesh, containing nodes on the crack lips. In the conforming mode this nset is used to

calculate the tangential plane at each crack front point, an the orientation of the front normals at this point.

• when the optional **\*\*compute\_K** command is specified, the default G energy release rate output, is replaced by the mode I  $K_I$  stress intensity factor. This  $K_I$  value is obtained from G using the following equation:

$$K_I = \sqrt{\frac{E G}{1 - \nu^2}}$$

Note that in order to evaluate the previous equation, the  $G - \theta$  module needs to retrieve the elasticity coefficients of elements close to the current crack front node, and that the influence of external parameters (eg. temperature) on those parameters is accounted for. Note also, that when this option is used, the propagation law behavior automatically takes K values as input, instead of the default G energy release rate, such that coefficients of those laws should be modified accordingly.

• the optional **\*\*compute\_Ki** command activates the calculation of the  $K_I$ ,  $K_{II}$ ,  $K_{II}$ stress intesity factors by means of an interaction integral computation based on the Westergaard analytical solution. This choice has an impact on the branching criterion for out of plane propagation, that automatically switches to a modal mixity angle  $\alpha$ calculated from those stress intensity factors instead of the default *Gmax* based criterion. In this case,  $\alpha$  is computed using the following equation:

$$\alpha = 2 \arctan\left(\frac{-K_{II}/K_I + \sqrt{(K_{II}/K_I)^2 + 8}}{4}\right)$$

- command **\*\*fatigue** preload period is mandatory for fatigue crack propagation, and defines the cycle period (parameter period) and a preloading initial time value at the beginning of the calculation during which crack propagation is ignored (parameter preload). Depending on the above definitions, crack propagation will occur at times  $t = preload + n_{cyc}$  period, where  $n_{cyc}$  is the cycle number.
- command **\*\*behavior** btype defines the model btype selected to compute crack advance, that will drive crack propagation and automatic remeshing procedures (by means of the **\*\*\*auto\_remesh** command). Currently the only model available is the Paris law for fatigue crack propagation:

$$\theta = \frac{da}{dN} = C \left(\Delta G\right)^m$$

where  $\Delta G$  is the G variation during the cycle (or K if the \*\*compute\_K keyword is activated) computed at each point along the crack front.

• command **\*\*Delta\_N** dn defines a multiplication factor dn applied to the crack advance  $\theta$  calculated with the propagation law specified by the **\*\*behavior** command. This option may be important to insure that a significant crack advance will occur at each simulated cycle in the FE analysis, and that automatic remeshing is indeed pertinent.

• command **\*\*h** hmax is another way to control the value of the multiplication factor dn applied to the  $\theta$  values. When a positive value is given, hmax is the target crack advance when remeshing occurs, and the multiplication dn is calculated accordingly:

$$dn$$
 such that  $hmax = \max_{i} \left\{ dn \; \frac{\theta_i}{\theta_{max}} \right\}$ 

where  $\theta_i$  is the crack advance at point *i* on the crack front, and  $\theta_{max}$  the max value of  $\theta_i$  on all crack front points. When a negative values is specified, *hmax* defines a target crack advance of *hmax* times the mean size of elements along the crack front.

• the command **\*\*optim\_dir** prec scan triggers out-of-plane propagation. Without this keyword, the crack advance direction always lie in a plane corresponding to the initial crack definition: in the xfem mode this plane is characterized by the first levelset definition, and in the conforming mode the **\*\*lip** nset allows this tangential plane calculation at each crack front point.

When a  $G_{max}$  bifurcation criterion is used (which happens when no keyword **\*\*compute\_Ki** has been specified, see above), 2 methods are available to compute the angle  $\alpha_{max}$  (at each crack front point) for which  $G = G_{max}$ .

First one (default case) is a dichotomy search where parameter *prec* is the precision, and *scan* the angle variation used during the scanning of angle values needed to intiate the dichotomy algorithm. Angle values should be entered in degrees, and a typical definition is:

\*\*optim\_dir 1. 15.

for a precision of 1 degree in the calculation of the modal mixity angle, and a scan of values every 15 degree to calculate the initial dichotomy interval.

The second method is activated with the **\*\*vectorial** keyword, and is based on the calculation of G in 2 orthogonal directions. The first direction, along the tangential crack direction, gives a so-called  $G_I$  value. Then a second  $G_{II}$  value is calculated in a direction orthogonal to this tangential plane. The branching angle  $\alpha$  for the new propagation direction is then defined as:  $\alpha = \arctan\left(\frac{G_{II}}{G_I}\right)$ .

#### Example:

```
% Conforming mode example:
***compute_G_by_gth FRONTO
% G-theta problem definition
**crack_front FRONTO
**nbnodes -8
**elem_radius 3
**elset NEW
**lip lip
% SIF output options
**compute_K
**compute_Ki
**excludeBE
```

\*\*\*\*calcul
 \*\*\*compute\_G\_by\_gth

% crack propagation law definition \*\*fatigue 0.000000e+00 2.000000e+00 \*\*behavior paris C 1.000000e-06 m 3.000000e+00 \*\*Delta\_N 100 \*\*h -2.0 % out-of-plane propagation \*\*optim\_dir 1.000000e+00 1.500000e+01

Example:

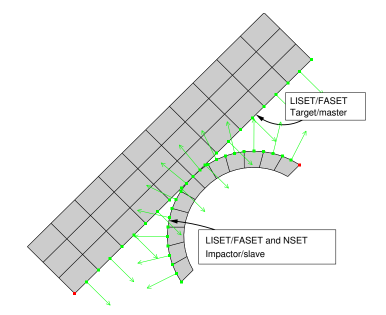
```
% XFEM mode example (note, no out-of plane propagation)
***compute_G_by_gth gtheta_a
   **xfem
   **theta 0.3 0.0
   **nbnodes 3
   **behavior paris
   C 335.298
   m 3.74
   **Delta_N 2000
```

# \*\*\*contact

#### **Description**:

The **\*\*\*contact** section defines zones of contact in static and dynamic mechanical problems. Typically, applying contact is one of the more troublesome aspects of nonlinear finite element analysis, because of increased discontinuities and sometimes unavoidable over- and underconstraints. This is above the obvious increase in modeling and input file complexity. After Z-set version 8.2, major updates to the contact system have been made which hopefully increase the robustness and quality of contact mechanics. The algorithms of Z-set are particularly good at avoiding global convergence issues with chatter, and in performing well when there are large differences in the relative stiffnesses of the contacting parts.

Contact is enforced with a group of *contact zones*. Zones are modeled by a node set acting as the *impactor* (*slave*) surface which may come in contact with a *target* (*master*) surface area. Each zone may have a different contact model (law) with or without frictional behaviors, and can include a variety of control parameters. The contact condition is defined in terms of a contact stress, but can alternatively be defined directly in terms of nodal reaction forces.<sup>3</sup>



To define contact zones, one must identify target/master surface areas, and corresponding impact/slave sets of nodes. Under normal use, i.e. stress-based contact, a node set (*nset*) is required for the impactor side, and a boundary set (*liset* or *faset*) is required for *both* target and impactor.<sup>4</sup> The boundary sets *must* be set up so that the normals point outward (see page 5.7), which can be verified graphically in Zmaster using the Bset normals option of the Mesh Bset... command. An example view of a properly set up two-dimensional example is given in the above image, where the target nset is not visible because it coincides with its corresponding liset/faset (as it should).

 $<sup>^{3}</sup>$ Note that for second order interpolation functions the magnitude and even direction of the nodal reactions is not intuitive. The fact is that treating nodal reactions directly assumes a discrete force acting over a zero area, which of course results in a singular stress in a continuum. Thus for these higher interpolations the only real choice is to deal with stresses distributed over an area.

<sup>&</sup>lt;sup>4</sup>This is in contrast to older versions where a bset was required for the target only. A facility is available to automatically generate the impactor bset from its nset, but in general it is preferable to set this up ahead of time. One can use the **\*\*\*mesher \*\*bset** command (page 2.16) with the **\*use\_nset** sub-command.

Note that if contact is occurring on only one node, or if there is a zero radius corner in contact, the contact stress will still be distributed over the adjacent boundaries. If the purpose of this modeling is to impose a boundary condition, or if the frictional behavior is not important, it is possible to use the **\*\*force\_basis** option, or to define the impactor boundary set with a dot set. In the latter case, each node has unit area and does not have any additional connectivity.

### Solution method:

Contact introduces an additional system of constraints on the displacement field. These are implicitly defined in terms of the contact stresses. The contact strategy in Z-set consists in solving a sub-problem for the contact stresses and relative displacements between surfaces, from which the additional global constraints are computed (a process very similar to a substructuring problem). The nonlinear contact solution is thus done by running "sub-space iterations" depending on the chosen contact law, and a *flexibility matrix* found from the global stiffness. The flexibility matrix is a (possibly) full matrix describing the deformation of each node on a contacting surface due to a unit force on each other node on the contact surface. Note that these terms will only be zero if the nodes are on separate bodies. The contact-modified solution intervenes in the global iterative procedure in several steps:

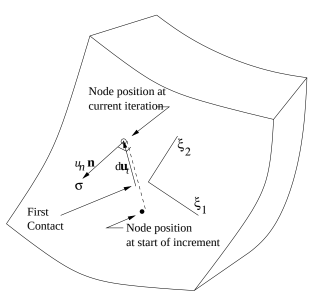
- 1. An estimate of the global field is made *without* enforcing the contact in the current iteration, by applying the classic Newton method using the last converged residual. This residual consists of both the internal reactions and the added contact reactions from the last iteration.
- 2. The flexibility matrix is computed from the global stiffness matrix by using repeated back solves of unit loads over all contact nodes. Because back solves are being made, it is very advantageous to have all the contact nodes numbered at the end of the global unknowns. However, this will in general increase the total bandwidth of the problem. Very often the computation of the flexibility matrix will be the most important part of the solution time of a contact problem. Currently the sparse matrix solvers include a compromise renumbering scheme for best performance.
- 3. The next step is then proceeding with the "inner" iterations by using this flexibility matrix to solve for the contact stresses and displacements. There are both direct solvers (involving the inverse of the flexibility matrix) and block diagonal iterative solvers in case that direct solution becomes too costly. The solver can be chosen with the **\*\*solve\_method** option. By default, the currently implemented contact laws use a direct solver. Running in verbose mode (either using the **-v** command line switch, the **\*\*verbose** output option, or by using the **\*\*verbose\_contact** contact option) will show a great deal of additional output on the progress of the local contact solution. This option should be enabled if there are any difficulties with the contact solution.
- 4. The last stage updates the set of global residual equations to include the contact reactions, which will in turn be used for the next dof prediction in the global Newton iterations.

The contact algorithm outlined above has many advantages, at the cost of computing the flexibility matrix. Firstly the method isolates the severe nonlinearities involved in solving the contact state from the global material and geometry nonlinearities. In particular, the effect of "chattering" is almost totally isolated to the local contact iterations, where it is very much more efficiently treated. In addition, the local system defining the contact is nonsymmetric

in the case of sliding friction, but because these terms are solved only in the local iterations, the global matrix can remain symmetric even under high friction (as long as other equations have led to a symmetric tangent before the contact has been considered).

#### Contact behavior:

The contact behavior is defined in terms of the quantities schematically represented in the following figure:



As shown, there is a normal mode of contact (to be enforced to null within a given limit), and a shear mode with a displacement vector defined in local coordinate axes  $(\xi_1, \xi_2)$  of the face element. The following variable definitions are used in the descriptions of the contact laws:

- $\sigma$  the normal contact stress (scalar), defined with compressive values being positive.
- au the tangent stress vector.
- $u_n$  the normal displacement (scalar) between potentially contacting surfaces. Values greater than zero have a gap between the surfaces and values less than zero indicate a penetration.
- $d\mathbf{u}_t$  the incremental tangent displacement (vector), measuring the projected slip on the element face during the increment.
- $k_n, k_t$  intrinsic contact stiffnesses in the normal and tangent directions (units of stress/length), as estimated by the code from the global matrix.
- $\mu$  (not drawn, dimensionless scalar) the friction coefficient as a fraction of the normal stress.

With these variable definitions we describe the following contact models:

normal contact with the following auxiliary variables  $\hat{\sigma} = \sigma - k_n u_n$  and  $\hat{\tau} = \tau - k_t d\mathbf{u}_t$ ,

the normal contact law is expressed as

$$\begin{array}{ll} \text{if } \hat{\sigma} > 0 \\ \text{then} \\ k_n u_n = 0 \\ \text{if } |\hat{\tau}| < \mu \sigma \\ \text{then } k_t d\mathbf{u}_t = 0 \\ \text{else } \boldsymbol{\tau} = \mu \sigma \mathbf{n}_t \quad \text{with} \quad \mathbf{n}_t = \hat{\tau}/|\hat{\tau}| \\ \text{else } \sigma = 0, \boldsymbol{\tau} = 0 \end{array}$$

The normal and tangential stiffnesses  $k_n > 0$  and  $k_t > 0$  are normalizing parameters which are automatically set by the program.

penalty contact is identical to normal contact, except that it allows for the contact constraints to be violated using the penalty parameters in the normal and tangential directions  $\gamma_n$  and  $\gamma_t$ . The larger the penalty, the more the constraint is satisfied. Numerically, these penalty parameters tend to make the local contact matrix diagonally dominant and can condition the matrix under situations where other constraints conflict with the contact constraints. When the penalty parameters approach infinity, the penalty contact model behaves exactly as the standard normal contact. In addition, in problems with dynamic impact, it is useful to include a term  $\beta$  which effectively damps any oscillations introduced by severe impact, while introducing only a small penalty under less impulsive conditions. Defining the auxiliary forces  $\hat{\sigma} = \sigma - [\gamma_n^{-1}\sigma + k_n(u_n + \beta\gamma_n^{-1} du_n)]$ and  $\hat{\tau} = \tau - [\gamma_t^{-1} d\tau + k_t d\mathbf{u}_t]$ , the penalty contact model is expressed as

```
if \hat{\sigma} > 0

then

\gamma_n^{-1}\sigma + k_n(u_n + \beta\gamma_n^{-1} du_n) = 0

if |\hat{\tau}| < \mu\sigma

then \gamma_t^{-1} d\tau + k_t d\mathbf{u}_t = 0

else \boldsymbol{\tau} = \mu\sigma\mathbf{n}_t with \mathbf{n}_t = \hat{\boldsymbol{\tau}}/|\hat{\boldsymbol{\tau}}|

else \sigma = 0, \boldsymbol{\tau} = 0
```

isotropic coulomb is almost identical to penalty contact, with the exception that the friction stress is limited by a user-defined maximum shear strength  $\tau_{max}$ . Thus, the normal contact model is a subset of the penalty contact model, and the penalty contact model is a subset of the isotropic Coulomb contact model. Defining the auxiliary forces  $\hat{\sigma} = \sigma - [\gamma_n^{-1}\sigma + k_n(u_n + \beta\gamma_n^{-1}du_n)]$  and  $\hat{\tau} = \tau - [\gamma_t^{-1}d\tau + k_t d\mathbf{u}_t]$ , the isotropic

Coulomb contact model is expressed as:

if 
$$\hat{\sigma} > 0$$
  
then  
 $\gamma_n^{-1}\sigma + k_n(u_n + \beta\gamma_n^{-1} du_n) = 0$   
if  $|\hat{\tau}| < \min(\mu\sigma, \tau_{max})$   
then  $\gamma_t^{-1} d\tau + k_t d\mathbf{u}_t = 0$   
else  $\boldsymbol{\tau} = \min(\mu\sigma, \tau_{max}) \mathbf{n}_t$  with  $\mathbf{n}_t = \hat{\tau}/|\hat{\tau}|$   
else  $\sigma = 0, \boldsymbol{\tau} = 0$ 

orthotropic coulomb is a generalization of the isotropic coulomb model.

# Syntax:

Contact takes a number of main control commands, and sub-blocks defining the different zones and behavior models. One can alternately define the contact model to apply to all zones, or zone-by-zone.

The **\*\*soft\_param** and **\*\*penalty\_param** commands are no longer supported in versions 8.2 and newer. This is now specified using the **\*\*zone \*behavior\_coef** command, as described below for each specific contact model. A warning will be issued, but these commands will be ignored.

```
***contact [ contact model ]
    **zone [ contact model ]
    zone subcommands
[ **conv precision max_iter [ force_precision ] ]
[ **stable eps [ precondition|damp ] ]
[ **stable eps [ precondition|damp ] ]
[ **force_basis ]
[ **solve_method direct|iterative ]
[ **solve_method direct|iterative ]
[ **limit_activity t1 t2 ]
[ **spy_node node [ node ] ... ]
[ **verbose_contact [ filename ] ]
```

- **\*\*zone** defines a contact zone, as well as the law defining its contact model. Current (as of version 8.2) contact models are normal, penalty, coulomb and ortho\_coulomb. The keyword soft is deprecated and automatically converted to penalty instead. More than one zone may be specified. If multiple zones exist which all use the same contact model, then the contact model may be specified immediately after the **\*\*\*contact** command. Available sub-options are described on page 3.100.
- **\*\*conv** defines the convergence parameters. The first (real) parameter *precision* defines a non-dimensional tolerance defining the *reduction* of the contact residual relative to the residual at the beginning of the inner contact iterations (i.e. the global problem residual *without* the additional contact influences of this global iteration included). A moderate relative value is therefore quite acceptable, even for problems needing a high *absolute final* residual.

The default value of 1.e-5 is quite small, so in most cases this command will be useful. A recommended value of 1.e-2 indicates that the contact convergence is two orders of magnitude smaller than the current global convergence state. Unless it is believed that the influence of contact is very very significantly altering the global behavior, this value should be ok.

The second (integer) parameter is the number of iterations before contact divergence will be signaled (default 200). The last optional (real) parameter *force\_precision* indicates the desired magnitude of the contact residual relative to the magnitude of contact stress (default 1.e-5). Only one of the two convergence criteria needs to be satisfied in order to achieve convergence.

Unlike previous versions, the contact solution algorithm in 8.2 always assigns the initial guess of the contact stresses based on the contact stress determined from the previous global iteration. As the global iteration converges, the initial absolute error in the local

contact iteration will also decrease. It is recommended that *force\_precision* be set only as tight as the expected convergence rate of the global iteration. When the global iterations converge, the absolute error in the contact law will be satisfied to the same degree as the global iterations.

\*\*stable takes one real value eps (default 1.e-2) and the choice between precondition (default) or damping. If the static model contains multiple bodies and at least one of these bodies has a rigid body mode (if the contact constraints are not considered), then this command is absolutely required. For dynamic problems it is not needed. The command instructs Z-set to add a term on the diagonal of the stiffness matrix of the contact nodes, in effect adding a grounded incremental spring to each node involved in contact. The value of eps then denotes the relative magnitude of the spring constant.

The specifier **precondition** instructs Z-set to add *only* the incremental spring to the stiffness matrix. It does *not* add the force of the spring to the external forces. This option stabilizes the contact solution algorithm, but in no way affects the converged solution.

With the specifier damping, not only the incremental spring is added to the stiffness matrix, but also the force of the incremental spring is added to the external forces. This actually affects the converged solution. This specifier can be conveniently used to avoid quasi-static singularity problems when a free body is not initially in contact in the initial loading stages. A small value relative to the problem stiffness is recommended, as is a convergence study to ensure that this numerical technique does not significantly alter the structural behavior.

- **\*\*init\_d\_stress** causes the algorithm to make an initial guess for the stress increments based on the previous solution, thereby possibly accelerating convergence. This has advantages especially for monotonic loading paths. For non-monotonic loading paths, the option sequence disables this guess for the first increment of each sequence, so that the contact algorithm is not confused too much by sudden changes in loading direction. This command is somewhat similar in spirit to the **\*\*\*resolution \*\*init\_d\_dof** command (see page 3.188). Without this command, the initial guess for the stress increment is set to zero, that is, the *initial* guess of the contact stress at the end of the current increment is assigned a value equal to the contact stress at the end of the previous increment.
- **\*\*force\_basis** sets the algorithm based on forces instead of stresses. Synonyms: **\*\*lumped** and **\*\*forced\_based**.
- **\*\*solve\_method** specifies the type of solver used for the contact solution. The option **direct** (default value for all currently implemented contact laws) uses the recommended direct solver. A second option **iterative** uses less memory, but requires more iterations to converge.
- \*\*limit\_activity only activates the contact conditions between times  $t_1$  and  $t_2$  (taken as two real parameters).
- **\*\*spy\_node** causes Z-set to print out more detailed information of the contact stresses at the specified list of nodes. These nodes must lie in the node sets given under **\*\*zone** impactor.

**\*\*verbose\_contact** gives additional information about the current state of all contact nodes involved. This information is written to an output file *filename*, or in absence of a user-supplied filename to a file called *problem.contact\_info*.

### Advice when problems occur:

The following hints may help out with common issues of setting up contact problems.

- Use the **\*\*verbose\_contact** contact option to examine the progress details.
- Check the target (master) surface normals to make sure that they point outward from the target body. This can be easily checked in Zmaster's Mesh mode, under the Bsets.. popup to observe best normals. In the case that some normals are inversed, one can use the inverse\_liset (page 2.68) or faset\_align mesher commands (page 2.59).
- Check to see that the target is the stiffer and/or less refined surface.
- Check the contact behavior. All contact models from versions before 8.3 are deprecated. The preferred "standard" model is **coulomb** which includes penalty parameters for "soft" behavior in the case of overconstraint, can optionally capped friction behavior.
- Check if any of the bodies are unconstrained in any of the degrees of freedom if not for contact. In this case, the global matrix will be singular and will no doubt cause problems during the first prediction of new displacement fields. In this case use the **\*\*stable** precondition command with a small value (e.g. 1.e-4). This command adds a small value to the global matrix pivots for the contact nodes, stabilizing the global matrix conditioning, but not actually altering the residual computation (i.e. not changing the physics at all). Of course a large stabilization value makes the global matrix less correct, and eventually convergence will be inhibited by this parameter. This method is only needed for static problems.
- Check to see if a body is underconstrained in some dofs, and may not always be in contact. If this is the case one can use the **\*\*stable damp** command which will include *incrementally* defined springs on the contact nodes with a given stiffness. Please be certain (by doing an appropriate convergence study) that the value of this parameter does not actually alter the results.
- Check to see if there are impactor nodes near the point of expected contact. If the slave surface is too coarse or the warning distance is too large, the contact may not be properly detected. In particular, if the impactor is too coarse there may be cases where the impactor does not penetrate the target, but the target does penetrate the impactor. The definition of the master/slave relationship of contact zones allows the master nodes to penetrate the slave.
- Check to see if there are sharp convex points in the contact zone. This can cause problems because the surface normals vary tremendously on the master surface, or by redistributing the contact stress erroneously around corners of slave surfaces.
- Check the integration order of the elements. Generally underintegrated elements will produce less oscillations (spatial and temporal) in the solution. This is especially the case under highly hydrostatic constraint or in locations where there is a rapid change in the

contact condition (e.g. Hertzian contact problems). This will be especially important for isochoric plasticity and viscoplasticity problems, or otherwise close to incompressible elements.

• Check for odd convergence and 'chattering" when the contact residual rocks back and forth between two residual values. Sometimes there is a problem with a node alternating between sticking and sliding contact states. In such an instance increase the slipping penalty parameter.

# \*\*zone

#### **Description**:

Contact zones define different contact pairs (impactor/target), zone by zone control parameters, and contact material behaviors.

#### Syntax:

The zone syntax is summarized below:

```
**zone [ contact model ]
 *impactor impactor_zone
 *target target_zone
 *behavior_coef
  key-value-pairs
[ *dont_check_bcs ]
[ *variable_friction file [ position ] ]
[ *warning_distance warning_distance ]
```

- \*\*zone specifies a contact zone with its specific contact model. Details of each model can be found in the following pages. Currently (as of Z-set 8.2), valid choices are normal, penalty, coulomb and ortho\_coulomb. Multiple \*\*zone commands may exist, each with its own contact model. If all contact zones have the same contact model, this model may be specified immediately after the \*\*\*contact command, instead of here. There is no default behavior. Parameters of the contact model are given with the \*behavior\_coef command.
- \*impactor (or alternatively \*slave) needs the nset impactor\_zone. The associated bset (liset or faset) has to exist as well. The bset has to be oriented with normals pointing away from the interior of the associated body.
- \*target (or alternatively \*master) takes the bset target\_zone. Again, this bset must be oriented properly.
- **\*behavior\_coef** sets the parameters of the specific contact model. Details can be found in the following pages.
- \*dont\_check\_bcs specifies that the code will *not* check whether contact conditions are compatible with other boundary conditions that may exist at the same node, as it does by default. With the default behavior, the code will try to fix any incompatibilities, or if that is not possible, the contact condition will be ignored. A warning message will be given if this is the case. This option disables the verification for the current zone.
- \*variable\_friction (or \*file) specifies a variable friction coefficient  $\mu$ , depending on the current and/or cumulated value of tangential slip. This variable friction cannot (yet) depend on any external parameter, such as temperature. A precise description of this behavior can be found in the material manual under \*\*\*behavior variable\_friction. Not implemented for the ortho\_coulomb contact model.
- \*warning\_distance specifies the maximum distance warning\_distance for which the algorithm will try to detect if contact occurs between target and impactor of the current zone. In other words, if the distance between target and impactor is *larger* than this

warning distance, the algorithm will *not* try to detect contact. This distance should be chosen larger than the decrease in distance between target and impactor during the increment of interest, otherwise there is a risk of penetration without the contact algorithm coming into action.

The subcommands **\*friction** and **\*gap** are now deprecated, as they are replaced with their equivalents in the **\*behavior\_coef** subcommand. Z-set will issue a warning, and in the case of **\*friction** automatically convert to the new format. The **\*gap** command will be entirely ignored.



# \*\*zone normal

#### **Description**:

This is the normal contact model as described a few pages back. For the meaning of its coefficient the reader is referred to that description.

# Syntax:

The normal behavior zones take all the standard \*\*zone commands, with the following specific behavior coefficient:

\*\*zone normal
[ other zone subcommands ]
 \*behavior\_coef
 friction mu

The default value is mu = 0.

# \*\*zone penalty

#### **Description**:

This is the penalty contact model as described a few pages back. For the meaning of its coefficients the reader is referred to that description.

#### Syntax:

The penalty behavior zones take all the standard \*\*zone commands, with the following specific behavior coefficients:

\*\*zone penalty
[ other zone subcommands ]
 \*behavior\_coef
 friction mu
[ penalty\_normal gamma\_n ]
[ penalty\_slip gamma\_t ]
[ damp\_normal beta ]

Default values are mu = 0,  $gamma_n = 1.e12$ ,  $gamma_t = 1.e2$  and beta = 1.

# \*\*zone coulomb

#### **Description**:

This is the "standard" contact model for contact with Coulomb friction, as described a few pages back. For the meaning of its coefficients the reader is referred to that description. Note that this contact model effectively contains the normal and penalty contact models.

### Syntax:

The Coulomb behavior zones take all the standard **\*\*zone** commands, with the following specific behavior coefficients:

```
**zone coulomb
[ other zone subcommands ]
 *behavior_coef
 friction mu
[ max_shear tau_max ]
[ penalty_normal gamma_n ]
[ penalty_slip gamma_t ]
[ damp_normal beta ]
```

Default values are mu = 0.,  $tau_max = 1.e12$ ,  $gamma_n = 1.e12$ ,  $gamma_t = 1.e2$  and beta = 1.

# \*\*zone ortho\_coulomb

### **Description**:

This is the orthotropic Coulomb contact model. It is similar to the isotropic Coulomb model, except that some parameters now have two be supplied with two values instead of one, and that the initial material orientation (vector) with respect to the master or slave surface now has to be specified.

# Syntax:

The orthotropic Coulomb behavior zones take all the standard **\*\*zone** commands, with the following behavior coefficient key-value pairs:

```
**zone ortho_coulomb
[ other zone subcommands ]
 *behavior_coef
 friction mu_1 mu_2
[ max_shear taumax_1 taumax_2 ]
[ penalty_normal gamma_n ]
[ penalty_slip gamma_s1 gamma_s2 ]
[ damp_normal beta ]
[ direction nx ny nz ]
[ master | slave ]
```

Default values are  $mu_1 = mu_2 = 0$ ,  $taumax_1 = taumax_2 = 1.e12$ ,  $gamma_n = 1.e12$ ,  $gamma_s1 = gamma_s2 = 1.e2$ , beta = 1., (nx ny nz) = (1. 0. 0.), and the last keyword has default value slave. Note that this behavior is currently not compatible with the \*variable\_friction zone subcommand.

### \*\*\*dimension

#### **Description**:

Because of numerical noise, Z-set sometimes has to make decisions about when a quantity is very small, small, large, or huge. For instance, when a typical time increment during a calculation is of the order of 10 s, output is not written to a file at t = 1000 s (if so requested through the **\*\*\*output** command) when the increment ends at t = 999.9999999 s.

In order to remedy this kind of problem, Z-set has predefined typical values that occur very often, and defines small as "multiplier  $\times$  typical value". The example above will pass with the default values: the default value for time is 1.0 and the default multiplier associated with small is  $10^{-6}$  (so small =  $1.0 \times 10^{-6} = 10^{-6}$ ), and since

1000 - small = 999.9999990 < 999.9999999 < 1000.000001 = 1000 + small,

output will be written. However, for certain other cases these default values need to be modified, for instance for impact problems where time increments may come down to the order of  $10^{-7}$  s or less. This can be done through the **\*\*\*dimension** command.

The **\*\*dimension** command lets the user modify the typical values for **stress**, **deformation**, **displacement**, **time** and **undimensional**. The multipliers associated to **tiny**, **small**, **large** and **huge** may also be changed. Default values are listed below.

#### Syntax:

The syntax is as follows:

\*\*\*dimension
[ \*\*unit unit typical ]
[ \*\*size size multiplier ]

unit may be stress, deformation, displacement, time or undimensional.

typical specifies the typical order of magnitude (a positive double value) that will occur for the quantity unit. The default values are 100. for stress, 10<sup>-6</sup> for deformation, 0.1 for displacement, 1. for time and 1. for undimensional. Note: this does not specify the actual units (despite the name of the \*\*unit command that suggests otherwise). For example, giving \*\*unit time 1.0e-6 does not mean that all times are measured in microseconds.

size may be tiny, small, large or huge.

multiplier gives the multiplier (a positive double) associated with the size keyword. Default values are  $10^{-12}$  for tiny,  $10^{-6}$  for small,  $10^{6}$  for large and  $10^{12}$  for huge. Their respective values should satisfy tiny < small < 1.0 < large < huge. Note that the multiplier given here affects all units.

#### Example:

The following example may be useful for impact problems, where very small time increments often occur. In absence of these commands, output will not be generated at the proper instants for time increments smaller than  $10^{-6}$  s.

```
***dimension
    **unit time 1.0e-6
```

Z-set — Non-linear material & structure analysis suite

### \*\*\*eigen

### **Description**:

The procedure **\*\*\*eigen** allows to specify the modes and resonant frequencies to extract in eigen-value problems.

The eigen frequencies are normalized in the following manner:

 $Max(\text{over the nodes})(\sqrt{U_1^2 + U_2^2 + U_3^2}) = 1$ 

And the associated energy for a frequency is calculated as:

E = U.K.U with U normalized as given above

Eigen frequencies are stored in the file named *problem.eigen* which can be visualized with the post-processor program. The frequency values are also stored with their associated energies in an ASCII text file named *problem.eigen\_info*.

# Syntax:

\*\*\*eigen method nb\_freq d\_freq freq\_max

There are currently two methods implemented to extract eigen values: lanczos and inverse\_vector\_iteration (this is the default method).

*nb\_freq* is the number of eigen frequencies to find.

 $d\_freq$  is the search interval for the resonant frequencies. If several frequencies are located in an interval smaller than  $d\_freq$  the calculation may fail to find all the frequencies. The lanczos method doesn't need this parameter.

 $freq\_max$  is the maximum value of the frequencies to search for.

# Example:

For this example the calculation will search for 8 frequencies and then stop. The calculation will stop also if the last frequency extracted is greater than 10000 ( $freq\_max$ ).

\*\*\*eigen lanczos 8 10000.

Alternaltively, one can use the inverse\_vector\_iteration method (note that this method is slower):

\*\*\*eigen 8 100. 10000.

The extraction of the resonant modes is made with the method of inverse powers and with a shifting of frequencies. After having obtained a frequency f, the next frequency is searched about  $f + \Delta f$  where  $\Delta f$  is given by the parameter *d\_freq*. The choice of this frequency is therefore very important.

- If d\_freq is small the calculation may fall back on a frequency already calculated. In this case the search will be shifted to  $f + 2\Delta f$  and so on until a new frequency is found. This succession of searches will inevitably add to the cost of the calculation.
- If *d\_freq* is large the search process risks to omit intermediate resonant frequencies.
- Lastly if it is desired to have the first resonant frequency it will be necessary to give a small value for the parameter *d\_freq*. If the value is small enough the calculation will be sure to not skip the first frequency. Searches for subsequent frequencies will then be very slow however.

### \*\*\*elastic\_energy

#### **Description**:

This option indicates that a calculation of the global elastic strain energy should be made and output. This calculation gives the following energy calculation for the structure:

$$\frac{1}{2} \int_V \tilde{\sigma} : D^{-1} : \tilde{\sigma} \quad dV$$

where D is the elastic matrix extracted from the the material behavior.

The result of this the elastic energy is stored in a formatted ASCII file named *problem.elastic\_energy*. The output is sequential, with one line per increment of the calculation.

### Syntax:

\*\*\*elastic\_energy elset-name
[ \*\*frequency frequency ]
[ \*\*precision precision ]

- *elset-name* is a character name for the element set over which the strain energy is calculated. The predefined element set ALL\_ELEMENT may be used to declare the entire structure.
- frequency may be used to specify when the energy is computed (by default, it is made at each increment). This option has the same syntax as output frequency (see p.3.150).
- precision is an integer specifying how many significant digits are used in the output file (default is 6).

# Example:

From \$Z7TEST/Energy\_test/INP/elastic\_energy1.inp

\*\*\*elastic\_energy ALL\_ELEMENT

# \*\*\*equation

#### **Description**:

This option is used to add linear relationships (equations) between different degrees of freedom in the problem. This allows the standard set of linear equations to include constraint conditions (MPC = multi-point constraint). In the current version of the code, these supplemental relationships are resolved by the direct elimination of degrees of freedom.

### Syntax:

***equation			
$**MPC_type$			
•••			

The relationship types which are currently defined are summarized in the following table:

CODE	DESCRIPTION
**free	impose an arbitrary relation between a single DOF and any number of others (p. 3.111)
**mpc1	sets nodal DOFs of a given type to be equal within a node set (p. $3.112$ )
**mpc2	sets two groups of nodes on a paired basis. A linear relation is enforced between the <i>i</i> th DOF in one node set and the <i>i</i> th node on the other (p. $3.113$ )
**mpc2x	is similar to mpc2, and provides an extended syntax (p. $3.114$ )
**mpc2_dof_elset	Sets a condition similar to $**mpc2$ between two DOFs belonging to ELSETs (p. 3.115)
**mpc3	models sectorial symmetry of the 3-axis $(p. 3.116)$
**mpc4	sets the DOFs of an node set equal to a surface function (line set in 2D or face set in 3D); this condition allows incompatible meshes to be attached at the surface (p. $3.117$ )
**mpc2d3d	combines mixed dimension meshes $(p. 3.119)$
**mpc_periodic	periodic boundary conditions (p. 3.120)
**nul_div_u	this procedure imposes a constant volume in the interior of a line set in 2D or a face set in 3D (p. $3.121$ )
**mpc_rb	imposes a rigid body motion on a nset (p. 3.123)

\*\*free

# **Description**:

This MPC imposes any relation between a single DOF and other DOFs. There are multiple ways to define the DOFs.

The equation applied is:

 $s = c_1 m_1 + \dots c_n m_n + C$ 

# Syntax:

This command has a slightly free form, non-standard format.

```
**free
slave is
master1 coef1
...
masterN coefN
coefficient
```

The terms above slave and master# are some special definitions of DOFs specific to this MPC condition. The syntax is given below. The coef# values are real number for the  $c_i$  terms above. And *coefficient* is a real value for the coefficient C. Note that the **is** above is a necessary keyword.

node:id:XX nodal dof, id is the node id, XX the dof type.
node:120:U2.

element:id:rk:XX element dof, id is the element id, rk the rank of the dof in the element
 (counted from 1 to the number of element dof of type XX), XX the dof type.

element:25:3:EZ.

elset:name:XX elset dof, name is the elset name, XX the dof type.

elset:metal:E33

# \*\*mpc1

### **Description**:

This equation type imposes that a given degree of freedom at all the nodes of a valid node set are equal in value.

# Syntax:

\*\*mpc1 nset\_name dof\_name

nset\_name Character name for a valid node set.

dof\_name Character name for a DOF keyword. The keywords available for different problem types are described in the chapter *DOF*.

# Example:

The following example could be used to assure that the top surface of a flat topped structure remains flat during deformation.

% from cisap.inp
\*\*\*equation
 \*\*mpc1 top U2

### \*\*mpc2

#### **Description**:

This equation type indicates that groups of nodes are tied on a paired basis. The  $i^{\text{th}}$  DOF in one node set will be equal to the  $i^{\text{th}}$  node on the other factored by a coefficient, plus a possible translation.

Alternatively, one can specify element sets to link DOFs belonging to elements. Note that node set names are search first, which could lead to some problems if element sets and node sets have similar names.

#### Syntax:

There are two possible syntaxes. A standard one and an extended one. The standard syntax is used to simply apply a factor between two degrees of freedom; the default value of *ratio* is equal to 1, and the relationship is such as  $dof2_{in}name2 = ratio$ .  $dof1_{in}name1$ 

\*\*mpc2 name1 dof1 name2 dof2 [ratio]

The extended one is used to introduce the affine relationship

 $dof2_in_name2 = ratio . dof1_in_name1 + translation:$ 

```
**mpc2 name1 dof1 name2 dof2
 *ratio ratio
 *translation translation
 *inversion
```

The **\*inversion** keyword may be used to prescribe an inversion of the nset order.

#### Example:

This could be used to model in 2D a sectorial symmetry following an angle of 45 degrees. The nodes of the bisection line of the 1 and 2 axis should then be imposed to have the same displacement following the axis 1 and 2.

This condition also allows modeling of axisymmetric conditions (symmetry in relation to a point) by taking two symmetrical node sets in relation to a point and tying similar typed DOFs of the same type with a coefficient of -1.

```
% from aube.inp
***equation
**mpc2 n_fuite U1 n_fuite U2 0.4431622
```

On the other hand, the following syntax allows the user to impose a shift between two lips of a crack.

```
% from tsqu1.inp
***equation
**mpc2 lipleft U1 lipright U1 *ratio 1. *translation 0.15
**mpc2 lipleft U2 lipright U2 *ratio 1. *translation -0.1
```

# \*\*mpc2x

### **Description**:

This equation type indicates that groups of nodes are tied on a paired basis. The  $i^{\text{th}}$  DOF in one node set will be equal to the  $i^{\text{th}}$  node in the other factored by a coefficient, plus a possible translation:

$$\mathrm{DOF}_Y^B = \alpha \cdot \mathrm{DOF}_X^A + \tau$$

It generalizes the standard mpc2, by allowing both the translation and the ratio to be given by a basic value (i.e. the product of a value and a time-dependent table, see p.3.39 for more details).

# Syntax:

```
**mpc2x
    nsetA dofX
    nsetB dofY
[ *translation value table ]
[ *ratio value table ]
[ *cumulative ]
[ *inversion ]
```

nsetA nsetB character name of the two paired node sets.

translation value<sup>5</sup> and table specifying the translation  $\tau$ .

ratio value<sup>5</sup> and table specifying the proportionality coefficient  $\alpha$ .

cumulative allows subsequent invocations of this mpc to be cumulated. It is otherwise forbidden, to avoid accidentally appying twice an MPC on the same DOF.

inversion may be used to prescribe an inversion of the nset order.

# Example:

The first example (from \$Z7TEST/Hyperelastic\_test/INP/ring-3-degrees-mpc.inp) prescribes  $U_{\theta} = 0$  in a sectorial mesh:

```
**mpc2x % U3 = z/x * U1 (ensures U_theta = 0)
face.2 U1
face.2 U3
*ratio function z/x ; tab_constant
```

The second example imposes the relative slide of both faces of a crack, where the translation is stored in an file (one value per node in the nset):

```
**mpc2x
faceA U1
faceB U1
*translation ascii_file slide_U1.dat tab_crack
```

<sup>&</sup>lt;sup>5</sup>As for boundary conditions, the value may either be a real value, a function or a file.

# \*\*mpc2\_dof\_elset

### **Description**:

The syntax is the same as for  $\tt mpc2$  but the condition is used to link DOFs belonging to elements sets.

#### Example:

This could be used in the case of periodic elements. Here in the case of large strains periodic elements:

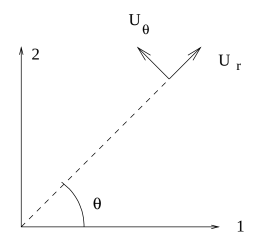
\*\*mpc2\_dof\_elset ALL\_ELEMENT E12 ALL\_ELEMENT E21

## \*\*mpc3

### **Description**:

The mpc3 equation type models sectorial symmetry about the 3<sup>rd</sup> axis. The following relation is applied to all the nodes of the given node set:

 $U_{\theta} = 0.0 \iff U_2 = tg(\theta).U_1$ 



### Syntax:

For this relationship it is required to give a node set name situated on the symmetry plane, and the angle of the sector in degrees.

**\*\*mpc3** nset\_name theta

nset\_name Character name for the node set in the plane of symmetry.

theta Real value for the sectorial angle.

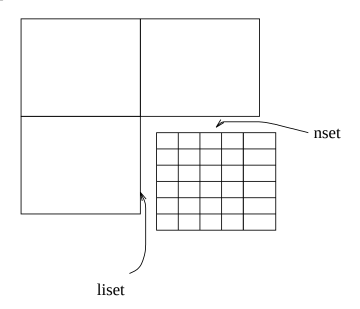
### Example:

```
% from zcentrifuge3.inp
***equation
    **mpc1 haut U3
    **mpc3 p0 1.0
```

# \*\*mpc4

### **Description**:

The mpc4 relationship is used to interface incompatible meshes. The DOFs of a given node set will be set to the DOF value on a corresponding boundary. This boundary will generally be taken on the coarser mesh, and is entered as a line set in 2D or a face set in 3D problems. Satisfaction of the mpc4 condition is made by DOF elimination on the node set.



The position of each node in the node set is first searched in the corresponding line or face set. Two cases may present themselves during this search process. The first is that the node has no projected position on the surface in which case no constraint is applied to that node. The second case is that the node does have a projected position on the surface. In this case the DOF type will be verified for existence on the surface and then subjected to the following relation:

$$U_{\rm nset nodes} = \sum_{\rm surface nodes} N_i U_i$$

If the DOF is not found on the surface during the verification no condition will be placed on that DOF.

It may be noted here that the global solution size (front) will be reduced if the numbering scheme is such that the element numbers attached to the node set are lower than those attached to the line or face sets.

... continued

\*\*\*\*calcul
 \*\*\*equation
 \*\*mpc4

# Syntax:

There are two possible syntaxes. A standard one and an extended one. The standard syntax is:

**mpc4	nset liset (faset)
and the extend	led one is:
**mpc4	
*nset	nset
*bset	liset (faset)
*delt	a $(dx,dy)$ $((dx,dy,dz))$

The **\*delta** keyword may be used to prescribe a fixed spatial offset between the nset and the liset. This is useful for instance to impose periodic conditions on a unit cell which exhibits incompatible pair sides (e.g. for a 2D square unit cell, left and right sides do not have the same number of nodes).

# \*\*mpc2d3d

### **Description**:

This MPC condition is used to "glue" a 3D mesh to a 2D mesh for mixed dimension analysis. The masters are the DOFs of the 2D mesh controlling the 3D slave DOFs. DOFs are attached (equal) if they have the same  $x_1$  and  $x_2$  coordinates.

# Syntax:

\*\*mpc2d3d nset2d nset3d dof dx

nset2d the name of a node set on the 2D mesh.

nset3d the name of a node set on the 3D mesh.

dof the DOF type to fix (e.g. U1, U2, ...).

dx a real value for the distance criteria for node position equivalence.

# \*\*mpc\_periodic

## **Description**:

This procedure enables one to prescribe an average deformation with periodic boundary conditions, according to

u = Ex + v(x)

where v(x) is periodic; the components are denoted E11, E22, E12, E21 in the 2D case and E11, E22, E12, E21, E33, E13, E31, E23, E32 in the 3D case. The actual value of the prescribed average deformation is equal to the product of the basic value *i* and the value in the table *tab*. Concerning the *nsets* it is important to distinguish faces, edges and corners.

# Syntax:

\*\*mpc\_periodic [inv]
dof nset1 nset2 component1 value1 component2 value2 ... tab

### \*\*nul\_div\_u

#### **Description**:

This relationship imposes a constant volume inside a surface (line set in 2D or face set in 3D) which is linearized by increment. The linearization cumulates an error of the same order as the incremental deformations. The condition may be strictly respected by demanding error correction, but only when the volume is calculated as follows: In 2D:

$$\int x \, dy \quad \text{or} \quad \int y \, dx$$

In axisymmetric:

$$\int 2\pi r z \, dr$$

In 3D:

$$\iint x \, dy dz \quad \text{or} \quad \iint y \, dz dx \quad \text{or} \quad \iint z \, dx dy$$

#### Notes:

The surface must be closed, or at the least its limits must have fixed displacements.

This procedure may be introduced in the middle of a calculation by using **\*\*\*restart**. If the keyword correction is not present after **\*\*\*restart** the volume is maintained constant from the instant of restart. In the contrary case the volume is maintained constant, but by taking the reference volume as the volume at time t = 0.

#### Syntax:

**\*\*nul\_div\_u** surf\_name [correction dir]

surf\_name Character name of the line set in 2D or the face set in 3D.

correction Keyword indicating that the corrections are to be activated.

dir Direction indicating the manner which the volume is calculated: In 2D plane geometries:

$$dir=\mathbf{d1} \quad \text{if} \quad V = \int x \, dy$$
  
$$dir=\mathbf{d2} \quad \text{if} \quad V = \int y \, dx \tag{26}$$

In 2D axisymmetric geometry:

$$dir=\mathbf{d2}$$
 if  $V = \int 2\pi r z \, dr$ 

In 3D geometry:

$$dir=\mathbf{d1} \quad \text{if} \quad V = \iint x \, dy dz$$
  

$$dir=\mathbf{d2} \quad \text{if} \quad V = \iint y \, dz dx$$
  

$$dir=\mathbf{d3} \quad \text{if} \quad V = \iint z \, dx dy$$
(27)

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3.121

\*\*\*\*calcul \*\*\*equation \*\*nul\_div\_u

# Example:

\*\*\*equation
\*\*mpc1 bottom U1
\*\*mpc2 bottom U1 top U2
\*\*mpc3 nset1 20.
\*\*mpc4 nset1 surface1
\*\*nul\_div\_u faset1

# \*\*mpc\_rb

### **Description**:

This command imposes a rigid body motion relationship between all nodes of a nset. Mpc can only define linear relationships between dofs, so that mpc\_rb is only valid under the small displacement assumption. The relationship is reactualized at every increment, so that mpc\_rb can be reasonably used with finite transformation if the increments are small enough.

In 3D, 6 "well chosen" master dofs are sufficient to define the rigid body motion (*i.e.* 3 translation components and 3 rotation components). They are automatically selected so that the three rotation parameters can be determined.

Under the small displacement assumption, the displacement  $\vec{U}_i$  of every node *i* (at location  $O_i$ ) of the nset can be expressed as:

$$\vec{U}_i = \vec{U}_m + O_i \vec{O}_m \wedge \vec{\theta}$$

where  $\vec{U}_m$  and  $\vec{\theta}$  are functions of the 6 master dofs.

## Syntax:

\*\*mpc\_rb nset

\*\*\*feti

### **Description**:

This command allows input of parameters used by the FETI iterative parallel solver, and therefore may only be included in parallel mode (execution with switch -PP as described in the chapter about *Parallel computing* at page ??).

As the FETI solver is based upon a Conjugate Gradient (CG) method to solve the interface problem arising from sub-domain decomposition, those parameters are mainly conventional input parameters associated with an iterative CG method.

## Syntax:

	***feti		
	**projector	proj	
	**precision	eps	
[	**max_iteration	iter	]
[	**max_standing	max	]
Ε	**precond	type	]
[	**keep_direction	dir	]
[	**reprojection]		
Ε	**init_lp]		

### \*\*projector proj

After subdomain decomposition the local problem defined on each subdomain may be singular. To avoid rigid-body motions the FETI solver uses a *projected* CG algorithm (see page ?? for reference). Projection of the gradient requires solving a linear system of equation whose dimension is the total number of rigid-body motions in the subdomains. The string argument *proj* specifies the method used to solve this system of equation, and can be one of the following:

CODE	DESCRIPTION
direct	Direct Gauss solver
iterative	Iterative Conjugate Gradient solver

Direct solver is faster, and this option is strongly advised unless small pivots problems arise during resolution in which case the *iterative* option may be more efficient.

**\*\*precision** eps, where eps is a real value defining the relative precision required for convergence when solving the interface problem with the CG method. For a system of equation:

 $\mathbf{K}\mathbf{q} = \mathbf{F}$ 

this relative ratio is defined by:

$$ratio = \frac{||\mathbf{F} - \mathbf{K}\mathbf{q}||}{||\mathbf{F}||}$$

and convergence occurs when:

ratio < eps

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- **\*\*max\_iteration** *iter*, where *iter* is the maximum iterations when solving the interface problem (default is 100).
- \*\*max\_standing max, where max is an integer specifying the maximum number of CG iterations allowed without any significant decrease of the convergence ratio.
- \*\*precond type This optional keyword is used to specify the type of pre-conditioning used to accelerate CG convergence. Until now, the only pre-conditioner available is lumped. Pre-conditioning is strongly advised and can significantly reduce the number of FETI iterations.
- **\*\*keep\_direction** *dir*, where *dir* is the the integer value of the number of orthogonal descent directions retained during the CG iterations. Increasing *dir* leads to faster convergence but is more memory consuming. Default value is *iter*+1.
- **\*\*reprojection** This subcommand can significantly reduce the number of FETI iterations, when used in conjunction with quasi-Newton schemes of tangent matrix update (such as eeeeee or p1p1p1, see the **\*\*algorithm** command). With this option the descent directions calculated during previous load increments are reused, leading to convergence in just a few iterations when the tangent matrix and the load increment stay constant over several Newton increments.
- **\*\*init\_lp** When a new resolution is asked for (new iteration, or new increment), this option allows to reuse the previous interface force field, instead of starting from a zero force field (which is the default option). This option has to be used together with the **\*\*reprojection** option.

# Example:

```
***feti
    **projector direct
    **keep_direction 150
    **precision 0.00000001
    **max_iteration 1000
    **precond lumped
```

# \*\*\*file\_management

# **Description**:

This command allows the user to adjust Zebulon's use of external temporary files.

# Syntax:

\*\*\*file\_management
[ \*\*keep\_temporary\_files ]
[ \*\*max\_nb\_dof num ]
[ \*\*z7\_tmp\_dir path ]

# Example:

The following is an example of user controlled file management.

```
***file_management
 **keep_temporary_files
 **max_nb_dof 1
 **z7_tmp_dir ./
```

On a Linux OS, some graphical monitoring of the in-core memory and temporary disk storage usage is available.

Zrun -monitor myprob >& /dev/null & Zplot\_stats STAT.4390

# \*\*\*fluid\_structure\_interface

### **Description**:

The fluid structure interface option is for eigen value extraction problems to model fluid filled cavities and their effect on dynamic response.

# Syntax:

```
***fluid_structure_interface
**interface
*surface_name surf-name
*fluid_mass_vol value
*free_surface nset-name
```

# Example:

```
***fluid_structure_interface
**interface
*surface_name interf1
*fluid_mass_vol 1.0e-20
```

# \*\*\*impose\_kinematic

### **Description**:

This command imposes a kinematic configuration (displacements) for a problem which does not have any kinematic variables (e.g. thermal or diffusion problems). It is commonly used for coupled problems.

# Syntax:

The syntax is:

\*\*\*impose\_kinematic type
 \*file fname

## Example:

```
***impose_kinematic from_transfer
    *file MechTherm/kine_out
```

# \*\*\*init\_dof\_value

# **Description**:

This procedure imposes the initial values of specified degrees of liberty of the problem (at t = 0). The values may be either uniform either defined by nset or elset or read from an initializing binary file. This file may be for example the output from a previous calculation (see the example).

## Syntax:

The syntax used to initialize the problem DOFs is:

init_dof_value	
<pre>dof_name elset elset_name value ]</pre>	
dof_name nset nset_name value ]	
dof_name uniform value ]	
dof_name file file_name position	]
	<pre>dof_name elset elset_name value ] dof_name nset nset_name value ] dof_name uniform value ]</pre>

where dof\_name indicates replacement with the character name of the desired DOFs (see appendix). The keywords elset and nset indicate the type of set. These keywords require a set's name elset\_name or nset\_name followed by a real value The keywords uniform and file indicate the method upon which the values will be loaded. Specifying uniform requires a real value for value which is the absolute value of the DOF. File storage set by the file keyword requires a character name for the file, file\_name, and an integer value for the record position, position. The DOF values will be taken from this file position in sequence using single precision 4 byte floating point format.

### Example:

This example shows how to load a data file generated from a small C++ program. This source is as follows:

```
#include <fstream.h>
main()
{ fstream out("U2.dat",ios::out);
  float x=1.;
  for(int i=0;i<8;i++) out.write(&x,sizeof(float));
  out.close();
}</pre>
```

which sets a uniform value of 1 (for example).

The input for this example could be:

```
***init_dof_value
U1 uniform 2.
U2 file U2.dat 568
```

which will set all the  $u_1$  nodal displacements to 2, and all the  $u_2$  displacements to 1.

# $***initialize\_with\_transfer$

### **Description**:

The **\*\*\*initialize\_with\_transfer** section allows to initialize a computation from the results of a previous one which have the same material variables. This feature is mainly used when one needs to modify the geometry of the problem and continue the computation.

### Syntax:

**\*\*\*initialize\_with\_transfer** takes a number of main control commands, and sub-blocks defining the results fields transfer method.

```
***initialize_with_transfer
[**format result_format ]
**old_problem result_file_name
**map result_card_num
[**use_deformed_mesh]
[**dont_use_deformed_mesh]
[**put_nodes_back]
[**quiet]
[**initial_time lime ]
[**auto_resume]
[**reequilibrium]
 *algo algorithm
 *ratio convergence
 *iter max_iterations
[**skip_nodal_transfer]
[**skip_integ_transfer]
[**nodal_var_transfer]
  [*mapping mapping_method ]
[**integ_var_transfer]
  [*integ_transfer transfer_method ]
```

- **\*\*format** defines the imported results format. If this option is omitted, the default Z7 format is assumed.
- **\*\*old\_problem** defines the name of the results database to transfer to current computation.
- **\*\*map** defines the results map to transfer.
- **\*\*use\_deformed\_mesh** use deformed configuration of the old mesh to locate the nodes/IP of current mesh (assumed TRUE if not specified)
- **\*\*dont\_use\_deformed\_mesh** use initial configuration of the old mesh to locate the nodes/IP of current mesh (use this to disable previous)
- **\*\*put\_nodes\_back** when a deformed mesh is used (use\_deformed\_mesh) this option allows to retrieve the initial non deformed configuration on the current mesh by subtracting the transferred displacement
- **\*\*quiet** run silently without extra messages (default is a more verbose mode).

- **\*\*initial\_time** set the initial time for the current computation. If this option is omitted, the default initial time is 0.
- **\*\*auto\_resume** set the initial time for the current computation to be the last time step available in loaded results
- **\*\*reequilibrium** run a "zero delta time" increment of computation at the end of transfer to retrieve equilibrium. This command has the same syntax as **\*\*sequence** (see page 3.190)
- **\*\*skip\_nodal\_transfer** disable the transfer of nodal variables (assumed FALSE if not specified)
- **\*\*skip\_integ\_transfer** disable the transfer of IP variables (assumed FALSE if not specified)
- **\*\*nodal\_var\_transfer** choose the method and its parameters for transferring nodal variables (use default nodal transfer options if not specified)
- \*mapping allow to transfer nodal variables between meshes of different dimensions (no mapping if not specified). The possible mapping methods are summarized:

CODE	DESCRIPTION
2d_3d	the results from 2D computation are mapped to 3D mesh assuming extrusion along z direction
axi_2d	the results from 1D axisymmetric computation are mapped to 2D mesh assuming revolution around y direction
axi_3d	the results from 2D axisymmetric computation are mapped to 3D mesh assuming revolution around y direction

**\*\*integ\_var\_transfer** choose the method and its parameters for transferring IP variables (if not specified, IP variables are extrapolated to the nodes then transferred as nodal variables by nodal interpolation, finally they are interpolated back to integration points)

\*integ\_transfer The possible transfer methods are summarized:

CODE	DESCRIPTION
nearest_gp	for each integration point in the current mesh, locate the nearest one in the loaded initialization mesh and simply copy the value to transfer
nearest_gp_corrected	in this mode, we add a correction step after nearest_gp, where the correction try to enforce the compatibility between trans- ferred DOF and the IP variables. To achieve that, a local integration of the behavior is done using the difference be- tween the transferred gradient and the one recomputed from transferred DOF
moving_least_square	two types of moving least square interpolations are provided: linear and quadratic (which is the default one). To choose an interpolation type add : linear_2D, linear_3D, default_2D, default_3D

## Example:

This example can be found in Transfer\_test/INP/TransferQuadra.2.inp

```
***initialize_with_transfer
    **old_problem TransferQuadra.1.ut
    **quiet
    **use_deformed_mesh
    **format Z7
    **put_nodes_back
        *to_file xx.geof
    **auto_resume
```

The same example modified to use moving least square

```
***initialize_with_transfer
    **old_problem TransferQuadra.1.ut
    **quiet
    **use_deformed_mesh
    **format Z7
    **put_nodes_back
        *to_file xx.geof
    **auto_resume
    **integ_var_transfer default
        *integ_transfer moving_least_square default_2D
```

# \*\*\*make\_restart\_file

### **Description**:

Allows "restart" files to be created at certain pre-defined moments in addition to the default backup file (.rst).

# Syntax:

***make_restart_file when		
CODE	DESCRIPTION	Ň
always	save at each increment	nent
end_of_seq	guence save at the end of	each sequence
end_of_cyc	save at the end of	cycles

There is no default for *when*. The restart files will have the name *pb\_name.rst#* where *#* is the incremental number of the restart file.

# \*\*\*matrix\_storage

### **Description**:

This command specifies which global matrix solver to use.

### Syntax:

This command is a one-liner with no sub-commands. The command takes a parameter which must be a keyword for one of the available solvers.

#### \*\*\*matrix\_storage type

The possible solver keywords are:

CODE	DESCRIPTION
frontal	Standard frontal method where the solution is made by marching across the element "front," adding and eliminat- ing degrees of freedom; This is the default
skyline	Traditional skyline method
$sparse_direct$	Direct (inversion) of a sparse storage matrix $^{6}$
$sparse_iterative$	Iterative sparse storage matrix

\*\*\*\*calcul \*\*\*material \*\*elset \*material

### \*\*\*material

### **Description**:

This command marks the definition of the materials in a structure to be studied. The behavior of each material is defined in a file with special syntax (see the chapter *Material Behavior*). The purpose of this command is therefore to define the material file names, associate these files to different element sets, and specify other global applications on top of a material model such as rotation of material coordinates or give local integration methods.

### Syntax:

```
***material
[ **elset nom ]
    *file nom [ num ]
[ *integration ]
[ *rotation ]
[ *var_mat_ini ]
```

The sub-procedure **\*\*elset** is used when there is more than one material. It is necessary then to give the name of the element set considered (*elset*). In the absence of the **\*\*elset** command, the following options with one asterisk will be applied automatically to the ensemble of elements.

The sub-procedure **\*file** is used to specify the file name for a material file in absolute or relative pathnames. An optional integer parameter is now available after the file name, giving the number of material declaration in a particular file. The second example shows this type of syntax.

### Example:

a simple example of the **\*\*\*material** syntax is:

```
***material
*file mat
```

For a material with multiple materials in different element sets, and example is:

```
***material
**elset E1 *file E.inp 1
**elset E2 *file E.inp 2
```

The program will search the first instance of **\*\*\*behavior** in the input file E.inp and assign that material to element set E1, while element set E2 will have the material defined after the second **\*\*\*behavior** in the input file. Please use caution when playing around with instances like this - Always Verify Your Materials.

## \*material

### **Description**:

This option determines the local integration method for a material behavior.

### Syntax:

*integration	method params
he allowable methods are summarized in the table below:	
CODE	DESCRIPTION
runge_kutta	explicit integration with automated time steps based on integration error
theta_method_	implicit generalized midpoint integration; this method nor- mally supplies the best tangent matrix
theta_auto_a	automatic time stepping in the implicit $\theta\text{-method}$
theta_method_	implicit integration by trapezoidal rule

runge\_kutta The Runge-Kutta method implements a second order explicit integration with automatic time stepping. Variables are normalized to allow varied variable magnitudes in "stiff" sets of equations. The method takes two real parameters. These are the convergence criteria followed by a minimum value for normalization. Standard RK error calculation for each integrated variable will be normalized by either the increment of the variable or this second parameter, whichever is greater. the resulting error is compared with the first parameter.

The Runge-Kutta integration with the gen\_evp material behavior provides a tangent matrix in models with a single inelastic deformation. This matrix is however not consistent with the integration scheme, and thus yields less than optimal global convergence. The explicit integration also performs poorly in heavily time-dependent problems such as viscoplasticity. However, some complex models are only implemented with this method.

theta\_method\_a The  $\theta$ -A method is the standard integration for the majority of material laws requiring integration.

$$x(t + \Delta t) - x(t) = \dot{x} (t + \theta \Delta t) \Delta t$$

This method requires 3 parameters to describe the convergence. These are first the  $\theta$  value (real) followed by the residual required for convergence (real) and the maximum number of local iterations in the integration (integer).

The value for  $\theta$  must be greater than zero and less than one. It is **strongly** advised to use  $\theta$  values of 1 for time independent (plastic) materials, and 1/2 for time dependent (viscoplastic) problems. Time independent plasticity will normally show strong oscillations about the solution for values of  $\theta$  less than 1.

Reasonable values of convergence range from  $10^{-6}$  to  $10^{-10}$ . Values which are too large usually lead to poor global convergence. Too small values will not converge due to

```
****calcul
***material
**elset
*rotation
```

numerical roundoff  $(10^{-12} \text{ is about the limit})$ . Convergence will rarely take more than 25 iterations, and should not take more than 50. If this is the case, there may be some error in the integration (make a bug report), or the material parameters are excessive (damage laws may provoke this). If the local iterations are greater than 50 it is probably better to reduce the global iterations or use automatic time stepping (global or local).

The default integration is dependent on the material law used. Most behaviors modeling plastic or viscoplastic materials use a default of the  $\theta$ -method with theta = 0.5, eta = 1.e-9 and max\_iteration = 200.

#### Example:

% plasticity or large deformation \*integration theta\_method\_a 1.0 1.e-9 50 % difficult viscoplastic case \*integration theta\_method\_a 0.5 1.e-6 100 % complex law

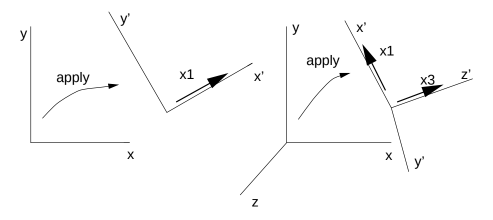
\*integration runge\_kutta 1.e-3 1.e-3

#### \*rotation

#### **Description**:

This material option is used to change a coordinate systems by rotation. It is used here to simplify specification of some materials (anisotropy, etc), but the syntax is general. Other applications using the rotation object include specification of grain orientations for polycrystals (see page 3.2), and mesh rotations (see page 2.100).

There are currently two methods for specifying a rotation. These are by vectors of the rotated coordinate axes in the global coordinate system, and by Euler angles (used for crystal orientation for example). The first case is displayed in the following figure:



\*\*\*\*calcul \*\*\*material \*\*elset \*rotation

For the material rotation of this section, the material gradient will be rotated (rotation is applied) before being integrated by the material behavior. For small deformation mechanics, this would be a rotation of the strain tensor.

$$\boldsymbol{\epsilon}_{tot}' = \mathbf{R}^T \boldsymbol{\epsilon}_{tot} \mathbf{R}$$

The material behavior then solves for the flux in terms of the new gradient, which is  $\epsilon'_{tot} \rightarrow \sigma'$  for the mechanical problem. Afterwards the flux is rotated to the global coordinates again:

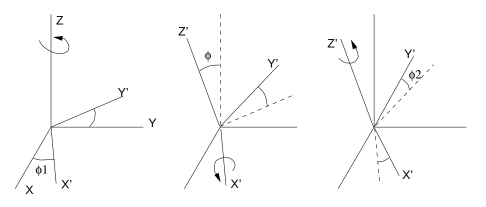
$$\boldsymbol{\sigma} = \mathbf{R} \boldsymbol{\sigma}' \mathbf{R}^T$$

Note that *only* the flux is rotated back to the global coordinates, but not the other quantities such as internal variables. For some material behaviors this can be modified, see for instance the **\*\*global\_output** option of the **\*\*\*gen\_evp** behavior in the Z-mat manual.



\*\*\*\*calcul \*\*\*material \*\*elset \*rotation

Rotation by giving Euler angles is similar. The significance of the three angles is given in the following figure:



#### Syntax:

For rotations specified using coordinate axes:

\*rotation
[ x1 x\* y\* [z\*] ]
[ x2 x\* y\* [z\*] ]
[ x3 x\* y\* [z\*] ]

The arguments x1, x2, x3 indicate the components of direction vectors for the transformed coordinate frame. Exactly one direction is required in 2D problems, and two directions are required in 3D. The order of definition is not important. The local coordinate system may be assembled with any of the geometrical axes. The input vectors will also be normalized by the program to automatically make unit vectors.

Using the notation here that t1 is the first direction vector defined, and t2 is the second (for 3D problems), direction vectors of the coordinate system are defined as follows: The first vector is collinear to t1. The second vector is a vector in the plane defined by t1,t2 and is perpendicular to t1. The third direction will always be calculated using the vector product of the first two vectors. The t vectors will replaced by those given by you using the x1, x2, x3 choices.

For rotations specified using Euler angles:

|--|

# \*\*var\_mat\_ini

### **Description**:

This option allows one to give the initial values for internal or auxiliary material variables. This option allows specifying any of the material variables, and thus depends on the individual material model. Common uses are to give the initial porosity or damage for initially porous or pre-damaged materials.

# Syntax:

The syntax for this option requires the name of an internal variable. The variable name is followed by a real number for the initial value to be applied, or by the keyword function and a function specification. The variable name can also be followed by either a character name for a binary data file, or the keyword ascii\_file followed by a character name for an ascii data file. Finally, it is also possible to initialize with a script, introduced by the z7p keyword.

Data files must furnish values for all the integration points stored as float variables in ascii or binary format. Ascii data files may also contain comments. Functions may only depend on coordinates x, y and z of the integration points. Any coordinate present in the function, but not at the integration point (typically a function depending on z for a two-dimensional mesh, as in the example below), will have a value of zero, so care should be taken in those cases.

The syntax is summarized below:

```
*var_mat_ini variable_name variable_value
*var_mat_ini variable_name function function;
*var_mat_ini variable_name [ ascii_file ] file_name
*var_mat_ini variable_name z7p script_name
```

### Example:

Some syntax examples follow:

```
*var_mat_ini epcum .015
  *var_mat_ini
    porosity
                0.1
                0.001
    grain_size
    epcum
                epcum.dat
  *var_mat_ini
                             x*x+(y-2.)-2000.*cos(z);
    porosity
                function
  *var_mat_ini
                ascii_file grain_size_distribution.dat
    grain_size
% from $Z7TEST/Program_test/Material_test/INP/bending.inp
```

```
*var_mat_ini eel11 z7p eel11-SC.z7p
```

# \*\*\*mesh

### **Description**:

Under the mesh description command we may specify the formulation used to satisfy the governing equations of the problem, including assumptions of a quasi-geometrical nature such as plane stress or plane strain conditions. The formulations possible in mechanical problems include classical small deformations, Total-Lagrangian, Updated-Lagrangian, or special cases such as the Cosserat continuum. Thermal problems do not have any alternate formulations.

The procedure also allows specifying a geometry file different than prob.geof.

## Syntax:

The syntax for this command is summarized below:

```
***mesh [ element_type ]
 **file filename
 **elset elset-name [ element_type ]
 *section type
 ...
 **local_frame type
 ...
 **import type fname
 **predefined type
```

The argument *element\_type* defines the type of element (its formulation) which will be used throughout the element set in question. The different types possible are described on the following pages. In order to simply apply the element formulation to the whole structure, fill in the *element\_type* on the same line as the **\*\*\*mesh** keyword. In order to define different element types for different regions of the mesh, put the *element\_type* keys after **\*\*elset** sub-commands.

Note that plane element geometries in 2D require a specification of element formulation in order to distinguish plane stress or plane strain. The default element type only applies to axisymmetric geometries in 2D. Element names with plane\_strain will enforce the  $\epsilon_{33} = 0$ condition, while elements with plane\_stress enforce  $\sigma_{33} = 0^7$ .

- **\*\*elset** elset-name sets the element formulation and possibly other characteristics for a given element set name. When used, no element formulation should be given after the **\*\*\*mesh** keyword.
  - \*section type this elset sub-command assigns section properties to a shell mesh. See the separate pages following which describe the particular sections available (starting at page 3.145).
- **\*\*file** Indicates that the mesh is in a file named other than *problem*.geof.
- **\*\*import** Indicates that the mesh file is in a separate file, and in a format other than native Z-set. It is normally recommended to use the batch mesher to translate the mesh before running the analysis (see page 2.7).

<sup>&</sup>lt;sup>7</sup>Plane stress is imposed in Z-set using additional degrees of freedom, and not at the material behavior level, although this exists too with some of the behaviors (notably gen\_evp; see page 3.2). For plane stress behaviors, one must choose ironically plane\_strain element formulations.

- **\*\*local\_frame** Assigns a local frame to the degrees of freedom attached to an elset (see page following at 3.146).
- \*\*predefined type Predefined elements may be used to quickly perform calculations
  on certain meshes. These are limited currently to single elements which can be used
  to quickly test and verify material behaviors. The predefined type type may be chosen from the following elements: cax8, cax8r, c2d8, c2d8r, cax4, cax4r, c2d4,
  c2d4r, rve1d, rve2d, rve3d, c3d20, and c3d20r. One can use Zrun -H to see the
  installed predefined elements.

The elements will be defined in the positive quadrant in a rectangle or cube form with its limits at 1. Two dimensional elements have liset and nsets defined as bottom, right, top, and left.

### Caution

In each calculation there may exist several different element types, and several different materials. Unfortunately, the dynamic structure of the program limits the ability to do exhaustive compatibility verification. Caution must therefore be given to the consistency of options within a calculation. The verification procedure will be improved in the next version of the code.

## Example:

The following is an example of the "simple" use of **\*\*\*mesh** to assign the whole problem to be a plane stress case, with a single predefined mesh as a 8 noded 2D element.

```
***mesh plane_stress
    **predefined c2d8
```

The second example is of a mesh stored in a fully qualified path name file, and two element sets with different element formulations.

\*\*\*mesh

```
**file /home/user/meshes/big_mesh.geof
**elset rubber_part total_lagrangian
**elset stiff small_deformation
```

## Mechanical calculations:

The following are brief descriptions of the mechanical element formulations available.

- spr Spring (or pinned truss) elements for one or 2 node geometries (12d1 12d2 13d1 13d2). This element requires that a valid spring material behavior be assigned to all such elements (c.f. 3.2). The two node element realigns with the line between those two nodes, and is therefore non-linear.
- linear\_spring A linearized version of the spring element, so no non-linearities are introduced.
- small\_deformation plane\_strain small\_deformation\_plane\_strain Mechanical elements in small deformation formulations. The integration volume is that of the initial structure, and significant rotations may cause erroneous calculation of the strain. The default 2D geometry is axisymmetric. Choosing the plane\_strain option will enforce the  $\epsilon_{33} = 0$  condition.
- plane\_stress Plane stress 2D element formulations. In order to be compatible with all material behaviors, this element adds degrees of freedom to each Gauss integration point which represent the strain  $\epsilon_{33}$ . These DOFs are used to enforce zero surface pressure and therefore  $\sigma_{33} = 0$ . Relationships may however be defined using the MPC command in order to specify generalized plane strain conditions. Setting the EZ ( $\epsilon_{33}$ ) DOF variable to be uniform over an elset yields zero overall applied pressure, but allows variation in the  $\sigma_{33} = 0$  field. The 3-direction strain will of course be uniform.
- small\_deformation\_updated plane\_strain\_updated plane\_stress\_updated Identical to the non \_updated formulations but using the geometry at the end of an increment as the integration volume. This produces non-linear structural behavior even with linear behavior models.
- small\_deformation\_select\_int small\_deformation\_select\_int\_updated Special version of small deformation elements for linear interpolation to solve the problem of strong local variations or oscillations in the stress fields (esp. pressure terms).
- cb\_shell cb\_shell\_updated\_lagrangian Shell elements formulated using the so-called "continuum based shell" assumption: the mechanical response of the shell element is computed using an underlying 3D volumic element whose geometrical configuration is dynamically constructed at runtime by the mother shell element. See for instance the corresponding chapter in the book: *Nonlinear Finite Elements for Continua and Structures*, T. Belytschko, W.-K. Liu and B. Moran, Wiley.

This element type uses 5 degrees of freedom per node: three displacements U1, U2, U3, and two rotations W1 and W2. Thus the rotation is assumed to be continuous along the shell.

Note that you must also give the initial thickness of the shell for this elset, using the following syntax:

\*\*elset shell\_part cb\_shell
 \*thickness 0.1

- total\_lagrangian total\_lagrangian\_plane\_strain Elements formulated for large displacement using Total-Lagrangian assumptions. These elements do not have any straining under rigid body motions.
- total\_lagrangian\_mixte\_u\_p total\_lagrangian\_plane\_strain\_mixte\_u\_p Incompressible elements in large displacements. The formulation is a mixed pressuredisplacement Total-Lagrangian method with degrees of freedom for the pressure at certain nodes.
- total\_lagrangian\_mixte\_u\_ps total\_lagrangian\_plane\_strain\_mixte\_u\_ps Incompressible elements in large displacements. The formulation is a mixed pressuredisplacement Total-Lagrangian method with a single pressure degree of freedom per element.
- updated\_lagrangian\_plane\_stress Updated Lagrangian\_plane\_strain updated\_lagrangian\_plane\_stress Updated Lagrangian element formulation for finite strain calculations. Must be used in the finite strain case for materials with internal variables such as metal plasticity or porous plastic materials. This element type *must* be used in conjunction with behaviors modified for updated Lagrangian methods (see the command \*\*\*behavior).
- periodic periodic\_plane\_strain periodic elements in axisymmetric, 3D, or plane strain. These elements allow one to impose the mean stress values for a periodic cell. Note that the displacement field that is saved by default is the *to-tal* displacement field and *not* only its the periodic part. You can override this choice and save only the periodic part by appending after the periodic keyword: periodic\_info \*\*periodic\_displacement\_field.
- smallw smallw\_updated Continuous mechanical element small displacement, small rotation - 3D only. This element combined with an appropriate behavior such as a smallw crystal can be used to model texture evolution in a crystal for moderate deformations.
- 2\_5D 2\_5D\_updated two and one half dimension elements. These use a 3D material law with a 2D geometry. There are six degrees of freedom per gauss point (element DOFs)  $[t_1 \ t_2 \ t_3]$  and  $[w_1 \ w_2 \ w_3]$ . The structures displacement field is:

$$\vec{u}(x, y, z) = \vec{u}_0(x, y, z) + \vec{u}_1(x, y)$$

$$u^{x} = u_{1}^{x} + zt_{1} - w_{3} (y - Y_{0}) z$$
  

$$u^{y} = u_{1}^{y} + zt_{2} + w_{3} (x - X_{0}) z$$
  

$$u^{z} = u_{1}^{z} + zt_{3} + w_{1} (y - Y_{0}) z - w_{2} (x - X_{0}) z$$

With this element, one can use full 3D material behaviors, and fix out of plane degrees of freedom. The values of  $X_0$  and  $Y_0$  are modified through the **\*\*\*specials** command.

• pressure Fluid interface mesh pressure elements.

#### \*section uniform

• cosserat cosserat\_plane\_strain cosserat\_plane\_stress Mechanical elements modified for the Cosserat continuum. This method adds DOFs at each node for an independent micro-polar rotation named W3. This method allows limitation of localization under shear deformation in strain-softening materials. The method uses a characteristic material length which must be input in the material definition, and the behavior must be modified for the Cosserat formulation (see the **\*\*\*behavior** command).

### Thermal calculations:

For thermal calculations, the type of element is completely defined by the geometry given in the .geof file. The **\*\*\*mesh** command should thus only be used if an alternate geometry file name is to be used.

### \*section uniform

#### **Description**:

This command assigns a uniform section of 1D elements to integrate the shell thickness. This command is a shortcut for having a separate mesh file for the section, which is possible too.

#### Syntax:

\*\*elset elset-name shell-formulation
\*section uniform num-elem ele-type thickness

### Example:

Here's a short example of a shell structure with 2 different section properties.

```
***mesh
**elset rib mindlin_shell
*section uniform 1 c1d3 .4
**elset flange mindlin_shell
*section uniform 1 c1d3 .25
```

# \*\*local\_frame

## **Description**:

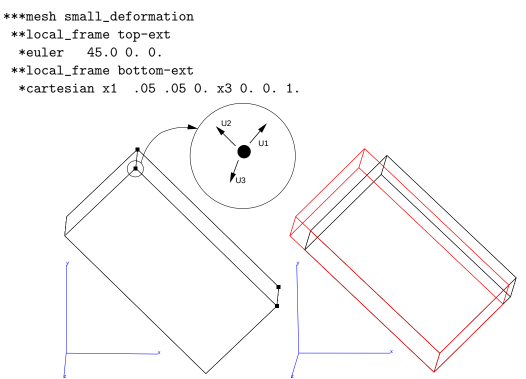
This command assigns a local coordinate system to the degrees of freedom attached to nodes of a given node set. The command is not compatible with every element formulation, and thus care should be used with this option. When in doubt, ask your supplier for additional advice.

# Syntax:

\*\*local\_frame nset
\*euler Euler rotation definition
\*cartesian Cartesian rotation definition
\*cylindrical
 center
[ axis ]

## Example:

The following example shows the use of this option to make a sliding boundary condition at an orientation away from the coordinate axes.



Another example is to impose a cylindrical coordinate system. This type linearizes the curvature, and is thus valid only for very small deformations.

```
***mesh plane_strain
**local_frame outer
*cylindrical
(2. 1.)
```

# \*\*\*sub\_problem

## **Description**:

This command is used to define a sub-problem in a weakly coupled analysis, and specify data to be transfered in order to interface with other sub-problems.

# Syntax:

\*\*\*sub\_problem type prob-name
\*\*transfer type
\*variable var-name
\*file out-file
....

The available transfer types are described below. In version 8.0 the transfers generate a separate set of files for each exported variable<sup>8</sup>, with names *out-file.catalog out-file.first out-file.initial* and *out-file.final*.

- integ\_integparam Make an integration variable (grad, flux, vint, vaux) and exported variable for use as an \*ip type external parameter in another sub-problem.
- integ\_nodeparam Make an integration variable (grad, flux, vint, vaux) and exported variable for use as an \*node type external parameter in another sub-problem. These variables are extrapolated, and will therefore have some error. The extrapolation is also averaged between elements, so it will not account for discontinuous materials well.
- **node\_nodeparam** Export one of the problems nodal variables (degree of freedom) for use as a node based external parameter in another problem (e.g. temperature).
- node\_kinematic Export nodal displacements in a way which is suitable for the
  \*\*\*impose\_kinematic option (see page 3.128).

 $<sup>^{8}\</sup>mathrm{these}$  will be combined into a "transfer" database in the next release

# \*\*\*output

### **Description**:

This command allows management of the output data files. By default, the code will only store the nodal DOF and associated reaction values in the file *problem.node*. In order to have the values of the material behavior variables the user is required to specify some of the sub-procedures below.

The version 7 of Z-set allows multiple output specifications, so the user may optimize disk usage for a problem. Normally this is very useful when the structure is to be output only at certain critical points during the calculation, while data plots may be output more frequently using the **\*\*curve** sub-option. Many of these curves may be specified with different output files, thus automating the post-processing stage of the analysis. Older versions of the code required saving the *entire* calculation with the frequency desired for the curves even if their information was rather specific. Use of output data files for the curves also allows automation of their generation using standard scripts with plotting software.

Syntax:

```
***output [ name ]
   **linear_solution ]
   **contour ]
   **contour_by_element ]
   **value_at_integration ]
   **save_in_material_frame ]
   **component name1 name2 ]
   **test tname ]
      *plot ]
      *precision digit ]
      *small small ]
     *gauss_var elem_id pg_id var1 var2 ... ]
      *node_var node_id dof_name1 dof_name2 ... ]
     *nset_var nset dof_name1 dof_name2 ... ]
     *liset_var liset var1 var2 ... ]
     *node_extrapolated node_id var1 var2 ... ]
      *element_node_var node_id var1 var2 ... ]
   **curve tname ]
   **frequency ]
      *dtime dtime ]
     *at time1 time2 ... ]
     *increment nb_increment ]
      *cycle cycle1 cycle2 ... ]
   **store_global_matrix matrix_type when filename ]
   **verbose ]
   **output_first ]
    **save_parameter [list of parameters' names] ]
[
[
    **reaction | **no_reaction ]
Γ
    **ele
                | **no_ele ]
```

[ \*\*node | \*\*no\_node ]

The optional character name may be used to specify the output file names. In the event of more than one **\*\*\*output** specification, the use of names will prevent overwriting of the same files for each output. This is essential to avoid, as the storage records are referenced using the .ut directory which would be overwritten for two like-named outputs. The default name will be the problem name *problem* which was given when running the Z-set program.

Names of the integration point variables are furnished for each material behavior. The names of the DOFs are given in the chapter *DOF*.

The sub-procedures are:

- **\*\*linear\_solution** This option indicates that the solution should be post processed with the intention to view linear summations of different load cases. The time steps are thus considered to be "load cases." In fact, this command only adds the keyword **\*\*linear\_solution** to the *problem*.ut file header, which can be edited by hand as well.
- **\*\*value\_at\_integration** This option indicates that default variables held at the integration points are to be stored in the file *problem.integ*. In each material behavior there will be a special set of variables which are stored by default (documented in the sections for each behavior). When the problem has more than one material specified, the union of all variable groups for all materials is stored at each integration point. If the value is not actually defined for a material at a certain integration point a null value will be stored<sup>9</sup>.
- **\*\*save\_in\_material\_frame** This option indicates that default variables held at the integration points are to be stored in the file *problem.integ*. By default they are stored in their local material coordinate system (if a **\*rotation** or a **\*pre\_problem\_treatment** is defined), but not the stress and strain fields **sig** and **eto**. With this option the latter are not only saved in the global coordinate frame, but also in the material coordinate system, in additional variables **sig\_lf** and **eto\_lf**. This option is very useful for composite materials.
- **\*\*contour** The contour option indicates that the integration point (material) variables will be stored at the nodal positions in the file *problem.ctnod*. Material variables are extrapolated to the nodal positions for all the elements. Each node has therefore a number of extrapolated values for each material variable to be stored, equal to the number of elements connected to that node. The recorded output value at the node will be the arithmetic mean of the individual element extrapolations. Data stored in this way will incorrectly smooth stress discontinuities across material boundaries. Such output is however better in the single material case because it assures continuity of variable fields from one element to another.
- **\*\*contour\_by\_element** This output option causes the material variables to be saved at nodal locations by extrapolation, element by element. The data is stored in a binary file named *problem.ctele*. Element extrapolation leaves the discontinuities at boundaries, and will thus represent the multi-material interfaces correctly. Each node will have a number of values stored in the output file corresponding to each attached element.

<sup>&</sup>lt;sup>9</sup>More intelligent strategies for outputting variable values will be implemented in future versions of the code

The magnitude of discontinuity as observed using this method of extrapolation may often be used as a measure of the mesh quality. Better meshes should show small differences element to element in the material variable values.

**\*\*component** This option allows direct specification of the material variables to be stored. This allows a reduction of the output file sizes by limiting the material variables output, or outputting of secondary variables not normally stored by the default output.

The code does not verify that the variables exist, as that would pose problems for multimaterial problems having a defined variable in one element set only. The stored value for undefined variables will always be zero. Note that if a variable is forgotten with this option it will be required to re-run the entire problem to obtain its value. It is also essential to leave this option untouched when using the **\*\*\*restart** options.

Syntax is the following:

\*\*component name1 name2 ...

where name# are the variable names to be output. One may see all the variable names defined by each material law by running Z-set with the verbose -v command line switch, or using the **\*\*verbose** option (see below).

- **\*\*save\_parameter** This option indicates that we wish to store the nodal parameters defined in the calculation. These parameters are stored in the file *problem.node* which contains the DOF values and associated reactions. These parameters are then able to be examined together with the calculation results in the post-processor.
- **\*\*frequency** This option adjusts the frequency of output storage for all the options within an **\*\*\*output** block. The option allows a variety of definitions in problem time or in increments. If a given output time does not correspond directly to a discrete calculation time, the next solution after the requested time will be output. The problem is always stored at the end of the calculation.

The sub-options are described below:

- \*dtime sets the output in by increment of time. This option takes a single real value for the increment of time between each output.
- **\*at** specifies the exact times for output. This option takes a list of real values for the output times.
- \*increment outputs at every specified number of increments. The option takes
   a single integer which defines the increments between outputs. For example
   \*increment 3 will store increments 3,6,9, etc.
- \*cycle outputs only during the specified cycles numbers. Any number of integer cycle numbers may be entered after the keyword.
- \*\*test is used to output specific data in an ASCII text file. Giving an optional character name after the \*\*test option will define the name of the output file for all the variables specified after this option. By default, the filename will be the output name (as given after \*\*\*output or the problem name in the absence of an output name), suffixed by .test.

The output data will be controlled by a variety of options:

- \*plot specifies that the output file is to be stored in column format (recommended). This allows direct plotting of the output data. The variable names will be listed at the head of the file with the **#** character before (a standard comment character).
- \*precision gives the precision for the output variables (number of digits after the decimal). All variables are output in exponential form. The default precision is 6 decimal places.
- \*small gives the value below which the result must be considered zero. Default value is  $10^{-9}$ .
- \*gauss\_var Specifies output of Gauss point data. The syntax requires the element number (integer) followed by the local Gauss point number (integer - numbering is described in the appendix). After the Gauss point specification an unlimited list of character names for the material variables to be output is given. These names may be any of the valid material names as viewed while running with the -v command line switch.
- \*node\_var This option specifies that nodal quantities are to be output. The syntax requires a node number (integer, or alternatively the name of a nset containing a single node) followed by a list of character names for the nodal variables to be output. The variables may be DOF names (such as U1, U2, etc) or the DOF reactions prefixed with an R ((such as RU1, RU2, etc).
- \*nset\_var Stores nodal reactions as summed over a node set. The syntax requires a node set character name to be given followed by a list of the nodal DOFs from which the reaction is to be taken. Note that the R is not given here. An example is \*nset\_var top U2 which will output the applied load  $(F_2)$  over the surface defined by the node set top.
- \*liset\_var Output data along a line set. Data is output in separate files for each output time, so it is strongly suggested to use the \*\*frequency option. The syntax takes a liset name, and a list of variables to be output on the liset. The standard definitions X, Y and Z may be used to output the coordinates. Any other nodal variable is possible to output on the liset. If material variables are demanded, they will be output using an averaged extrapolation of the adjacent elements. Example: \*liset\_var outside Y U1 U2 RU1
- **\*node\_extrapolated** Material variables will be extrapolated to the nodal coordinates. The syntax takes the node number, followed by a list of integration variables.

\*node\_extrapolated 22 sig22

Note that mixing of the **\*-\_var** options may be given in order to force sequential output of all the variables.

- **\*\*curve** This option is identical to **\*\*test** with the **\*plot** sub-option automatically set.
- **\*\*store\_global\_matrix** allows storage of the global matrix associated to the structure's mesh in a file. The matrix may thus be saved at different points in the calculation

and re-used for the calculation of resonant eigen frequency analysis of the pre-loaded structure (see **\*\*\*preload**). The syntax is the following:

\*\*store\_global\_matrix

matrix\_name when filename

[...]

where *matrix\_name* is the name of the matrix to be stored, *when* is the time at which the matrix is to be saved, and *filename* is the filename to be used for the matrix. Currently the only value for *matrix\_name* is **stiffness\_matrix** which stores the global problem stiffness, or **conduction\_matrix** for thermal problems. Such triplets may be repeated, in order to write the matrix at different times.

\*\*verbose Specifies verbose screen output during the calculation. As the screen output is stored in the file *problem.msg* this option may create a very large file for long (many increments) problems. It has been found that the .msg file may rapidly become larger than the primary output files so verbose output is not normally set. This option is equivalent to using the -v command line switch.

The verbose option is especially useful in problem setup, however, as it causes output of the local material variables to the screen.

**\*\*reaction, ele, node** Enables (resp. disables for the **no\_** prefixed options) the storage of these values in the results files.

# Example:

```
***output
     **test
      *precision 4
      *small 1.e-6
       *gauss_var 1 1 evi11 sig22
       *node_var 53 U1
     **frequency
       *dtime 1.5
       *increment 10
     **store_global_matrix
       stiffness_matrix 10.0 disk.stif
***output
   **frequency
     *at_time
               30.
                     60.
               270.
                     300.
               1770. 1800.
               3570. 3600.
               7170. 7200.
   **contour
   **value_at_integration
   **save_parameter
 ***output none
    **curve q1.charge
       *nset_var top U2
    **curve q1.dv9
       *node_var 2 U1
       *node_var 25 U1
    **curve q1.stress_strain
      *gauss_var 1 1 eto22 sig22 epcum
```

\*\*\*\*calcul \*\*\*output \*\*test \*J

\*\*test

#### \*J

#### **Description**:

This command is used for calculating the J integral or  $\Delta J$  using a line set only. The command will work in 2D plane stress and plane strain only. Any material may be used, and any standard mechanical element formulation.

The integral is evaluated by performing direct integration of the contour integral

$$J = \int_{\Gamma} \left[ W n_x - \sigma_{ij} n_j \frac{\partial \mathbf{u}}{\partial x} \right] d\Gamma$$

The calculation will be negative if the path is oriented clockwise. Also, remember that if there is a plane of symmetry about the crack plane, and you only model this one half, the associated J integral will be one half of its real value.

#### Syntax:

. . .

The syntax for this output command is the following:

\*J liset\_name [ time-ini time-end ]

Each contour desired should be entered as a separate \*J command.

Adding the two optional real values time-ini and time-end will cause the J integral to be evaluated as the  $\Delta J$  integral. Note that this is not an incremental version of J as the LEFM  $\Delta K$  is used, but rather an evaluation of J over a path with residual stresses. Caution

Because the J integral is computed along a boundary, it is using extrapolated Gauss point data to the nodes, which is then again re-interpolated to the intermediate points on the line. This can cumulate some error, especially if the mesh is too coarse. One should verify the stability of J calculations with respect to path (path independence), and the mesh size.

Also, because this is a **\*\*\*output** option, it is subject to the **\*\*frequency** command. As the J integral is integrated over time, for non-linear problems it's calculation requires a sufficient number of time steps.

#### Example:

\*\*\*output
 \*\*contour
 \*\*test
 \*J path1
 \*J path2
\*\*\*output
 \*\*contour
 \*\*test
 \*J path1 0. 10.
 \*J path1 20. 30.

# \*\*\*parameter

#### **Description**:

Using the coefficient mechanism, material behavior may depend on both internal variables calculated by the material law, and external user specified parameters which are essentially problem loading. The coefficient syntax is discussed in the **\*\*\*behavior** chapter in the Z-mat manual.

Use of the parameter mechanism may also intervene directly in the material law such as thermal deformations which depend on a temperature parameter. This temperature field may be the output of previous calculations to provide immediate capability of un-coupled or weakly coupled multi-domain solutions. Other non-standard parameters are also permissible allowing user data to be used in the calculation. One example is specifying a distribution of the density, and then making the elastic modulus depend on this density.

There are several methods of specifying the parameters within a geometry.

#### Syntax:

The baseline syntax for external parameters follows.

***parameter	type parameter_name
[ *node	*ip ]

- type a \*\*-level sub-command for the type of external parameter input which is desired. The different types are described in the pages which follow, including their specific values. In the absence of a type (input starts directly with \*-level commands, or data entry fields, the type \*\*file is taken as the default.
- parameter\_name Character name of the parameter. The naming of parameters is inherently open in the code. However, the use of certain names have particular signification for different options in the material behavior or global loading. An important example is the name temperature which is searched by the thermal strain material objects.
- **\*node** indicates that the parameter exists at nodal locations, and will be interpolated to integration points. Record sizes are the number of nodes. This is the default.
- \*ip indicates that the parameter exists at integration points. The record size will be the number of integration points in the mesh.

It is imperative that the parameters be defined at all times from t = 0. to the end time of the problem. Between the times for which the parameter is defined through *file*, *function* or *uniform* the value will be linearly interpolated. It is therefore desirable that the times of parameter definition correspond to the end sequence times. Uniform values may be applied in an analogous manner as the boundary conditions with the use of tables. ... continued

### \*\*\*\*calcul \*\*\*parameter

# Examples:

Two simple examples are shown below.

#### \*\*\*parameter temperature

- 0. uniform 125.
- 1. uniform 175.
- 2. uniform 175.
- 3. uniform 125.

\*\*\*parameter humidity

- \*ip
- 0. uniform 0.25
- 1. uniform 0.25
- 2. uniform 0.75
- 3. uniform 0.25

\*\*file

# **Description**:

This is the default parameter type. It allows mixed use of uniform parameter values, parameter values specified by a function and binary file records containing the values of the parameter field. This parameter type is very similar to the ascii\_file type (see page 3.160), but binary files generally load faster, which is important for very large parameter files. Note: some command options are available only in Z-set version 8.4 or newer.

# Syntax:

```
**file param-name
[ *node ]
[ *ip ]
[ *dtime ]
[ *table_file size ]
[ *table_file tablefile name ]
[ *cycle_conversion start end period ]
   time uniform value
   time function func;
   time file filename record
```

- **\*\*file** takes *param-name* (a character string) as argument for specifying the name of the parameter. This name will be used to reference the parameter values elsewhere in the problem.
- **\*node** specifies that the parameter is applied at nodes. This is the default.
- \*ip specifies that the parameter is applied at integration points.
- \*dtime converts the absolute time values given in the table below to time increments.
- \*table\_file takes as argument the character string table\_file name as the name of the file from which the table containing the list of times or time increments with their associated parameter types (uniform, function or file) can be read. This is an alternative to directly specifying the table here (see below), and can be useful for automatic table generation by scripts.
- **\*rec\_size** enter an integer defining the record size. This value should be the number of nodes for **\*node** type parameters, or the total number of integration points for **\*ip** type parameters.
- \*cycle\_conversion real values for *start* time of cycles, *end* time, and cycle *period* are to be input. This command generates a variable time which can be used in function records to define the time.
- $\dots$  continued

The table specifying parameter values contain the following elements:

time a time value for the given parameter data. Parameter values during the calculation will be found using linear interpolation between given times. Do not forget to specify a value for time = 0.0. If the \*dtime command is given, the time value becomes a time increment value. The time values may also be specified as a FUNCTION. This is especially useful in conjunction with the \*cycle\_conversion command (see example below).

value a uniform parameter value.

func a function specified in the standard Z-set manner. Do not forget the ; at the end.

filename name of the binary file to use. Different files can be given for different time values.

record record number in the file. These are numbered from zero (0 is the first record).

## File format:

The format for the parameter files consists of an ordered list of float (single precision floating point) values in big endian format. The ordering corresponds to the nodal numbering in the *problem*.geof file. Each record therefore comprises only this list of float values for each node. The record numbers are noted to be the record number (number of the block storing the entire nodal data set) and not the file position as would be read in C or Fortran programming.

# Examples:

The example below shows the use of mixed parameter types in order to impose a variable temperature field.

```
***parameter
**file temperature
*node
*rec_size 8
0. uniform 40.
1. file ../DATA/thermal_field.node 1
2. file ../DATA/thermal_field.node 2
3. uniform 120.
```

 $\dots$  continued

Here is a relatively complex example using a function to repeat values for many cycles. The *time* values are now specified using a function.

```
***parameter
 **file temperature
  *node
  *rec_size 757
  *cycle_conversion 1. 1.e20 60.0
   0.
                                    uniform 23.0
                       cycle*60.0; file mechanical.data 20
cycle*60.0; file mechanical.data 5
   function 1.0+
   function 1.0+ 10.0+cycle*60.0; file
   function 1.0+ 14.0+cycle*60.0; file
                                             mechanical.data 6
   function 1.0+ 18.0+cycle*60.0; file
                                             mechanical.data 7
   . . .
   function 1.0+ 56.0+cycle*60.0; file
                                             mechanical.data 19
   function 1.0+ 60.0+cycle*60.0; file
                                              mechanical.data 20
```

# \*\*ascii\_file

## **Description**:

This parameter type allows an easy input of parameter data with an ASCII file. The use of ASCII allows simple generation of the table values, for example by using scripts. Just as for the binary file parameter type (see page 3.157), it allows mixed use of uniform parameter values, parameter values specified by a function and binary file records containing the values of the parameter field. Note: some command options are available only in Z-set version 8.4 or newer.

# Syntax:

```
**ascii_file param-name
[ *node ]
[ *ip ]
[ *dtime ]
[ *table_file tablefile name ]
[ *cycle_conversion start end period ]
[ *rec_size size ]
   time uniform value
   time function func;
   time file filename record column
```

- **\*\*ascii\_file** takes *param-name* (a character string) as argument for specifying the name of the parameter. This name will be used to reference the parameter values elsewhere in the problem.
- \*node specifies that the parameter is applied at nodes. This is the default.
- \*ip specifies that the parameter is applied at integration points.
- \*dtime converts the absolute time values given in the table below to time increments.
- \*table\_file takes as argument the character string table\_file name as the name of the file from which the table containing the list of times or time increments with their associated parameter types (uniform, function or file) can be read. This is an alternative to directly specifying the table here (see below), and can be useful for automatic table generation by scripts.
- \*rec\_size specifies the size of each data entry record. For FEA problems and \*node type
  parameters, this is the number of nodes (can be found in the problem.geof file, or in
  Zmaster). For \*ip type parameters, this is the number of integration points.
- \*cycle\_conversion real values for *start* time of cycles, *end* time, and cycle *period* are to be input. This command generates a variable time which can be used in function records to define the time.

 $\dots$  continued

The table specifying parameter values contain the following elements:

time a time value for the given parameter data. parameter values during the calculation will be found using linear interpolation between given times. Do not forget to specify a value for time = 0.0. If the \*dtime command is given, the time value becomes a time increment value. The time values may also be specified as a function. This is especially useful in conjunction with the \*cycle\_conversion command (see the example given in the \*\*\*parameter \*\*file section on page 3.157).

value a uniform parameter value.

func a function specified in the standard Z-set manner. Do not forget the ; at the end.

filename name of the ASCII file to use. Different files can be given for different time values.

record record number in the file. These are numbered from zero (0 is the first record).

column column number for the data. Column number starts with 1.

## Example:

The following is an example of a variable Young's modulus imposed with an ascii\_file parameter.

```
***parameter
**ascii_file yng
*rec_size 20
*node
0.0 file param_as_young.inp 0 2
1.e20 file param_as_young.inp 1 2
```

The file param\_as\_young.inp of the example above might look like this :

```
1. 200000.0 % first line in file (rec 0)
2. 195000.
3. 198000.
...
20. 199000. % end of 0th rec
1. 220000.0 % start 1st record
...
```

The parameter is then applied in the constitutive law:

```
***behavior gen_evp
**elasticity isotropic
young = yng % young assigned to field parameter
poisson 0.25
**save_coefficients
young
```

If the parameter is to be applied at integration points, the data file must look like this :

\*\*\*\*calcul
 \*\*\*parameter
 \*\*ascii\_file

- 1. 200000.
- 1. 195000.
- 1. 198000.
- 2. 211000.
- 2. 205000.
- 2. 203000.

. . .

where the fist colomns indicates the element number. In this exemple, the elements have 3 integration points. rec\_size hes to be set to the total number of integration points.

# \*\*from\_results

### **Description**:

The **\*\*from\_results** command indicates that the parameter will be imported from a previous computation. It can of course be imported from a Z-set computation<sup>10</sup> but can also be imported from other software (Abaqus odb for example). Z-set then does the necessary time interpolations. In case of non-conforming meshes, where space interpolation is also necessary, use the **\*\*from\_results\_with\_transfer** syntax instead (see next page).

# Syntax:

\*\*from\_results param-name
 \*database format file\_base
[ \*initial\_value initial\_value ]
[ \*remap old\_name new\_name ]
[ \*ip ]

- param-name a character string name for the parameter. This name will be used to reference the parameter values elsewhere in the problem.
- format is the format of the imported file. For example, use Z7 to import Z-set results or odb for Abaqus. See page 2.8 for a complete list of known formats.

file\_base is the filename prefix of the results files.

initial\_value is a constant value to be used at t = 0.

# Example:

The following is an example of temperature imported from a previous Z-set computation:

```
***parameter
**from_results temperature
*database Z7 specimen_thermal
*remap TP temperature
```

 $<sup>^{10}</sup>$  and thus gives an alternate and simpler syntax than \*\*results, page 3.166

# \*\*from\_results\_with\_transfer

### **Description**:

The **\*\*from\_results\_with\_transfer** command indicates that the parameter will be imported from a previous computation. It is similar to **\*\*from\_results** (see previous page) and additionally allows for non-conforming meshes by doing the proper space interpolations.

## Syntax:

**\*\*from\_results\_with\_transfer** has the same syntax as **\*\*from\_results**. Refer to the previous page.

## Example:

The following is an example of temperature imported from a previous Abaqus computation:

#### \*\*\*parameter

\*\*from\_results\_with\_transfer temperature
 \*database odb thermal\_model

# \*\*function

### **Description**:

This parameter type is specified as a (t, x, y, z) function.

## Note:

If the function only depends on time, you may also use the **\*\*table** syntax.

## Syntax:

\*\*function param-name
 \*function function(t,x,y,z);
 \*tables table-names
[ \*time\_independant ]
[ \*node ]
[ \*ip ]

- *param-name* a character name for the parameter. This name will be used to reference the parameter values elsewhere in the problem.
- function a (t, x, y, z) function (do not forget the terminating;)
- table-names a (list of) table name entered in a **\*\*\*table** section. If more than one name is given, the first valid one is used (this validity being decided at each increment).
- \*time\_independant This switch allows the parameter to be time independent. The parameter is then be computed once at the beginning of the computation. This can be useful when random numbers are used in the function.

The parameter's value will be computed as the product between function and table.

#### Example:

The following is an example of a temperature gradient along x:

```
***parameter
 **function temperature
 *function 300. + 600. * x ; % gradient between 300 and 900 degrees
 *node
 *tables plateau % time dependence is given here
```

# \*\*results

# **Description**:

The **\*\*results** command indicates that the parameter will be read in from a Zebulon FEA results file (**\*\*changing** is a deprecated alias for the **\*\*results** parameter).

# Syntax:

The syntax required is:

\*\*results param-name
[ \*node ]
[ \*ip ]
 \*file\_base fname

- *param-name* a character string name for the parameter. This name will be used to reference the parameter values elsewhere in the problem.
- \*file\_base enter fname a file name prefix for the results files. The first file of concern is fileprefix.catalog, which lists the initial and final times for the file inputs (presumably from another time step of a mechanical problem, see below). Binary files for the beginning of the problem, start of an increment and end of an increment are: file-prefix.first file-prefix.initial file-prefix.final The current increment should be within the time bounds given in the catalog file.

# Example:

The following is an example of temperature values imposed with a table parameter.

```
***parameter
 **results temperature
 *node
 *file_base MechTherm/temp_out
```

# \*\*table

# **Description**:

This type of external parameter is used for uniform values which use entries from **\*\*\*table** (see page 3.199) to determine the parameter value through time. The input is rather like a boundary condition.

# $\underline{Syntax}$ :

The syntax required is:

\*\*table param-name
[ \*node ]
[ \*ip ]
value table

*param-name* a character string name for the parameter. This name will be used to reference the parameter values elsewhere in the problem.

value a scaling value (base value - decimal input).

table a table name entered in a **\*\*\*table** section.

# Example:

The following is an example of temperature values imposed with a table parameter.

```
***parameter
**table temperature
1.0 temp_tab
```

# \*\*\*post\_increment

#### **Description**:

This command starts reading of any number of post-increment calculations which are to be processed as additional output, or modifications based on the converged solution.

### Syntax:

***post_in	crement				
• • •					

The post-increment sub-options depend on the installed modules, and so may be expanded by using "plug-ins." As of Z8.0, the following models are included.

CODE	DESCRIPTION
parks	Parks method for Stress Intensity Factor
j_integral_lorenzi	a method implementing the $J$ -integral with virtual crack extension
non_local	A "weak" non–local model which can be applied to any state variable

## Example:

```
***post_increment
    **parks
    perturb elset rg1
    tip_radius 55.
    da (1.e-3 0.)
    da (1.e-5 0.)
    **j_integral_lorenzi
    perturb elset rg1
    tip_radius 55.
    da (1.e-3 0.)
    **j_integral_lorenzi
    perturb elset next
    tip_radius 55.
    da (1.e-3 0.)
```

\*\*\*calcul
\*\*\*post\_increment
\*\*i\_integral

# $**i_integral$

### **Description**:

This option calculates the T stress using the method of Chen etal. [Chen01]. and DeLorenzi [Hors85]. The implementation is similar to the **\*\*j\_integral\_lorenzi** in its use of a virtual crack extension.

## Syntax:

**i_integral	
perturb (elset   1	tip)(name1   node_num)
da (dax day)	to indicate the crack direction
	(and not the crack growth direction !).
tip nset	indicates the node corresponding to the tip
young E	gives the Young's modulus
poisson nu	gives the Poisson's ratio
[factor f]	indicates a multiplicative factor to be used
	in the case of symmetries. Default value is 1.

The T—stress calculations works with small deformation plane stress and plane strain elements. Like for the Delorenzi or Parks methods, the syntax perturb elset next will search the next elset surrounding the crack tip. This syntax can be repeated.

# \*\*j\_integral\_lorenzi

## **Description**:

This option calculates the J integral using the method of Horst and DeLorenzi [Hors85]. The implementation is analogous to the **\*\*parks** (page 3.173) in its use of a virtual crack extension. This method is more reliable (accurate, insensitive to mesh density, and path independent) for small deformation analysis than the output method **\*J** described on page 3.154. It is also valid for axisymmetric or 3D analysis where **\*J** is not. However, unlike that method the **\*\*j\_integral\_lorenzi** is limited to small deformation, and can not calculate  $\Delta J$ .

Remember that is the case of linear elasticity:

$$J = \frac{\frac{K^2}{E}(1-\nu^2)}{\frac{K^2}{E}} \quad \text{plane strain}$$

## Syntax:

\*\*j\_integral\_lorenzi
perturb (elset | tip) (name1 | node\_num)
continuing with as many lines as G calculations
[ tip\_radius val ]
da (dax day)

Refer to the **\*\*parks** option for a complete description of these options.

### \*\*non\_local

#### **Description**:

This post increment method provides a "smoothing" of specified material variables in order to constrain the gradient in those variables to be below a given limit. This is an important method in eliminating localization when stiffness drops according to those parameters (i.e. damage or strain softening). This is a "weak" method in that it allows localization during the solution convergence of a given increment.

Let d be the non–local variable ;  $\Delta d$  the local time increment ;  $\Delta d_{nl}$  the non–local time increment.

$$\Delta d_{\rm nl}\left(\vec{X}\right) = \int_{V} \Delta d(\vec{x}) \psi(\vec{x} - \vec{X}) \, dx$$

$$\psi(\vec{x}) = \frac{1}{\lambda^N \pi^{N/2}} \exp\left(-||x||^2 / \lambda^2\right)$$

where N = 1, 2, 3 is the space dimension.

$$\int_{-\infty}^{+\infty} \dots \int_{-\infty}^{+\infty} \frac{1}{\lambda^N \pi^{N/2}} \exp\left(-(x_1^2 + \dots + x_N^2)/\lambda^2\right) \, dx_1 \dots dx_N = 1$$

Using the FE method integrals are discretized using Gauss quadrature. So that

$$\Delta d_{\rm nl}(\vec{x}_i) = \sum_j \Delta d(\vec{x}_j) \psi(\vec{x}_j - \vec{x}_i) V_j = M_{ij} \Delta d(\vec{x}_j)$$

and  $V_j$  the volume associated to Gauss point j. where  $x_i, x_j$  are the positions of the Gauss points (G number of Gauss points).

Due to numerical integration and other limitations (optimizing memory) it is necessary to correct the M matrix which should fulfill the conditions

[c<sub>1</sub>] 
$$\sum_{j} M_{ij} = 1$$
 and [c<sub>2</sub>]  $\sum_{i} M_{ij} = 1$ 

Several correction methods can be proposed  $(M \to M')$ :

average\_line  $M'_{ij} = M_{ij} / \sum_i M_{ij}$ . [c<sub>2</sub>] is not fulfilled.

average\_column  $M'_{ij} = M_{ij} / \sum_{j} M_{ij}$ . [c1] is not fulfilled.

diagonal\_line  $M'_{ii} = M_{ii} + (1 - \sum_i M_{ij})$  and  $M'_{ij} = M_{ij}$  for  $i \neq j$ . [c<sub>2</sub>] is not fulfilled.

diagonal\_column  $M'_{ii} = M_{ii} + (1 - \sum_{j} M_{ij})$  and  $M'_{ij} = M_{ij}$  for  $i \neq j$ . [c<sub>1</sub>] is not fulfilled.

iterative Line *i* is multiply by a scalar  $l_i$  and column *j* is multiply by a scalar  $c_j$ . Conditions  $[c_1]$  and  $c[c_2]$  can be fulfilled simultaneously. There are  $2 \times G$  equations

$$\sum_{i} l_i c_j M_{ij} = c_j \sum_{i} l_i M_{ij} = 1$$

Z-set — Non-linear material & structure analysis suite

\*\*\*calcul
\*\*\*post\_increment
\*\*non\_local

$$\sum_{j} l_i c_j M_{ij} = l_i \sum_{j} c_j M_{ij} = 1$$

Coefficient can be found iteratively:

$$c_{j}^{k+1} = \frac{1}{\sum_{i} l_{i}^{k} M_{ij}} = \frac{1}{M^{T} l^{k} | j}$$
$$l_{i}^{k+1} = \frac{1}{\sum_{j} c_{j}^{k} M_{ij}} = \frac{1}{M c^{k} | i}$$
with  $l_{i}^{0} = 1$  and  $c_{j}^{0} = 1$ .

Syntax:

The command syntax is described below, with a few sub-commands.

```
**non_local
*lambda lam
*variable var1 ... varN
*elset elset-name
*cut_off val
*normalize type
```

\*lambda takes a real value for  $\lambda$ 

\*variable list the material variables which are to be the subject of non-local.

**\*elset** apply the method to the given elset name.

\*cut\_off specify a real value for the cut-off.

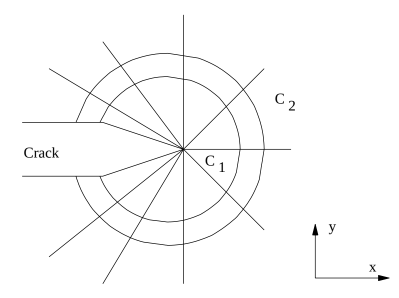
**\*normalize** replace *type* with one of the keywords described above (average\_line, etc).

### \*\*parks

#### **Description**:

The **\*\*parks** option indicates that the crack tip energy release rate G is to be calculated by the perturbation method (c.f. Parks [Park74]). The method is only available in 2D calculations.

Parks' method advances the crack length a small amount in order to evaluate the energy release per crack length unit. The obtained stress intensity factor taken from G is thus a "global" measure of the mixed mode I and II cracking.



It is necessary to define a translated internal zone (in the interior of the contour  $C_1$ ). The perturbed zone by the translation is that constituted by the first arrangement of elements about the translated contour (contour between  $C_1$  and  $C_2$ ). Beyond the contour  $C_2$  nothing will be modified.

The internal translated contour is fabricated from either an element set or a single node defining the crack tip. These geometrical groups must be defined in the *problem.geof* file with the meshing commands described in the leading chapters of this manual. For the case of axisymmetric calculations, it is necessary to define the radius of the crack tip point.

#### <u>Note</u>:

The results are stored in the sequential formatted file *problem*.parks with one line per increment calculated. Be careful that for for sructures exploiting symmetry (one half the crack is modeled) the value of G must be multiplied by a factor of 2, while K must be multiplied by the square root of two.

## Syntax:

name1 the character name of an elset composed of the translation elements.

```
node_num integer corresponding to the crack tip.
```

- *val* real value designating the radius of the crack tip. This option only applies to axisymmetric geometries.
- da vector describing the virtual advancement of the crack tip.
- dax real value for the advancement of the crack along the x axis only. This value may be negative.
- $day\;$  real value for the advancement of the crack along the y axis only. This value may be negative.

# Example:

```
***post_increment
    **parks
    perturb elset inter
    da (1.e-2 0.)
    da (1.e-4 0.)
    da (1.e-6 0.)
    da (1.e-8 0.)
    da (1.e-10 0.)
    da (1.e-12 0.)
    da (1.e-16 0.)
    da (1.e-20 0.)
    da (1.e-30 0.)
    da (1.e-100 0.)
```

# \*\*\*pre\_problem

## **Description**:

This procedure and its options are executed at the beginning of the problem, before doing anything.

# Syntax:

\*\*\*pre\_problem
\*\*first-pre-problem-type
first-pre-problem-specific options
\*\*another-first-pre-problem-type

There may be any number of sub-options defining the different pre\_problem to treat, and also any number of **\*\*\*pre\_problem** instances.

The different pre\_problems commands are the subject of the following pages. The following tables are included as a quick-directory to the pre\_problems.

# <u>General purpose pre\_problems</u>:

CODE	DESCRIPTION
**init_z7p_rotations	used to impose a material rotation to a partic- ular elset using a z7p program
$**layer_orientation$	used to define the stack sequence when a layer element (c3d16l for instance) is used.

\*\*\*\*calcul
\*\*\*pre\_problem
\*\*init\_z7p\_rotations

## \*\*init\_z7p\_rotations

#### **Description**:

The goal is there to compute a material frame (material rotation) for the given elset. For all the integrations points of this elset, the positions if each points is given to a t should then return a rotation matrix which will be used as the material for that integration point. This is very useful for composite materials.

#### Syntax:

\*\*init\_z7p\_rotations
 \*elset\_name name\_elset
 \*script name\_script

- name\_elset character name of the element set. This must be the name of a valid elset defined in the geometry file. The rotation will be imposed at every Gauss point in this element set.
- *name\_script* The character name of the zLangage script which defined the material frame as a function of the gauss point positions.

#### Example:

The following example apply a rotation to all gauss points of the elset called elbow. The rotation is around the z axis, such that the center is cx=27. and cy=10..

```
***pre_problem
    **init_z7p_rotations
    *script rotation.z7p
    *elset_name elbow
```

The script called rotation.z7p is given as follows :

\*\*\*calcul
\*\*\*pre\_problem
\*\*layer\_orientation

# \*\*layer\_orientation

#### **Description**:

The goal is there to define the stack sequence when a layer element (c3d16l for instance) is used.

## Syntax:

\*\*layer\_orientation
 \*elset name\_elset
 \*stack\_sequence number\_of\_layer
 frame\_1
 frame\_2
 ...
[\*elset\_rotation frame]
[\*element\_rotation\_script/ name\_script]/

- name\_elset character name of the element set. This must be the name of a valid elset defined in the geometry file.
- number\_of\_layer (integer) gives the number of layers of the element definef in name\_elset. This number must be consistent with the one defined in **\*\*\*mesh**.
- stack\_sequence used to indicate the type of stack, Define the rotation from the laboratory coordinate system (in which the material behavior is defined) to the coordinate system of the ply. Use Euler angle (see page 3.135) or a around the y axis.
- \*elset\_rotation frame These rotations are defined using Euler angles. It gives the rotations to chage the coodinate system from the glabal coordinates to the element coordinates (see example)
- \*element\_rotation\_script name\_script/ gives the rotations to chage the coodinate system from the glabal coordinates to the element coordinates using a zLangage script (see page 3.176).

# Example:

First example :

The following example define a laminate composite (0,+45). The material is defined in the global coordinate system shuch as the fiber direction is z.

```
***pre_problem
**layer_orientation
*elset ALL_ELEMENT
*stack_sequence y_axis 2
0. 45.
*elset_rotation
0. 0. 0.
```

For a material define such as 1 is the direction of fibres, 2 and 3 are equivalent

Second example :

The next figure shows an example such as the laminate composite is defined by a the (0,+45) stack sequence, and the elset is rotated at 90 around the z axis.

```
***pre_problem
 **layer_orientation
    *elset ALL_ELEMENT
  *stack_sequence y_axis 2
    0. 45.
    *elset_rotation
    90. 0. 0.
                                                                45
                                                                       х
                y
                                                                                        z
                             \mathbf{x} from the global coordinate system
                               to the element coordinate system
                                                                  y
           z
                                                                 from the element oordinate system
                                                                     to the ply coordinate system
                                                                       х
                                                                                z
                                                                                45°
                                                                             y
```

# \*\*\*random\_distribution

#### **Description**:

This command creates a random distribution in the structure, which can be used to create spatially randomized parameters such as material coefficients.

## Syntax:

\*\*\*random\_distribution
\*\*name name
\*\*type dimen type
\*\*law law
\*\*load load-file

- **\*\*name** gives a character name to the distribution. This user assigned name will be used to reference the random value at a particular point in space.
- **\*\*type** specify *dimen* (real) for the space dimension and a distribution *type*. The later is currently restricted to be **cellular**.
- **\*\*law** *law* define the type of random law to use. *law* should be a keyword defining the type. In the standard distribution, there are the following types available.

CODE	DESCRIPTION
uniform	Uniform law $F(x) = x$
weibull	Weibull type statistical distribution $F(x) = 1 - \exp\left[-((x-A)/b)^C\right]$
discrete	User gives $F(x)$
normal	F(x) = erf(x)

These laws are described in more detail in the following pages.

**\*\*load** is used to load the random field from a pre-existing file. This is used for example for verification tests where it is desired that the distribution be the same every time the case is run. One can use this to preserve interesting random distribution cases for more than one run.

#### ... continued

#### Example:

An example of random distribution being used for radomized coefficients follows.

```
***random_distribution
    **name Erand
    **type 2 cellular
    *start (0. 0.)
    *end (1. 1.)
```

```
*cell (.2 .2)
**load random.data
....
***behavior linear_elastic
**elasticity
young random Erand
poisson 0.3
***return
```

The random distribution can also be used to initialize material state variables.

```
***random_distribution
 **name eelrand
 **type 2 cellular
 *start (0. 0.)
 *end (1. 1.)
 *cell (.2 .2)
 **load random_iv.data
***material
 *file ../MAT/random_iv.mat
 *var_mat_ini eel11 random eelrand
****return
```

# \*\*\*resolution

### **Description**:

This procedure is used to fix the calculation parameters and method of resolution for the global matrix problem  $\mathbf{Ku} = \mathbf{R}$ . Except for the simplest linear problems, at least some of the procedures sub-options will be required. Non-linear problems are best solved with some substantial customization of these parameters.

#### Syntax:

Specification of the method of resolution requires a definition of the form:

```
***resolution type
```

which will be followed by a number of options of the **\*\*** command level. The possible types of resolution are summarized in the following table:

CODE	DESCRIPTION		
newton	resolution of type Newton-Raphson, or modified		
	Newton		
bfgs	resolution of type BFGS quasi-Newton		
riks	a Riks solution to pass by local instabilities or snap- throughs		

In the absence of type the default **newton** will be substituted.

This **\*\*\*resolution** command has a variety of sub-procedures to define load sequencing:

**\*\*sequence** Define loading sequences.

**\*\*cycles** Define cycles of loading sequences.

**\*\*automatic\_time** Define automatic time stepping.

**\*\*init\_d\_dof** Accelerate convergence by pre-initializing the DOF increment.

**\*\*no\_symmetrize** Use the non-symmetric solver.

**\*\*max\_divergence** Specify the maximum number of divergences allowable before the problem stops.

**\*\*skip\_cycles** Use the cycle skip algorithm.

**\*\*use\_lumped\_mass** Use the lumped mass matrix (only available for **\*\*\*\***calcul dynamics)

### \*\*\*resolution newton

#### **Description**:

The default algorithm is the **newton** method, which can be used as a full updated Newton-Raphson algorithm, or as a modified Newton-Raphson method depending on the **\*algorithm** choice in the sequence definitions.

Note that the finite element method calculates the following residual:

$$\mathbf{R}^i = \mathbf{F}_{ext} - \mathbf{F}^i_{int}$$

where  $\mathbf{F}_{ext}$  are terms of the discretized weak form of the problem variational statement due to externally applied forces,  $\mathbf{F}_{int}^i$  are the terms due to "internal forces" (e.g. the  $\int_V \delta \boldsymbol{\epsilon} : \boldsymbol{\sigma} dV$ virtual work term in mechanics), and  $\mathbf{R}^i$  is the residual imbalance at an iteration *i* due to the non-linearity of the problem. Convergence of a loading step is achieved when a measure of this residual falls below a desired magnitude.

A truncated Taylor series is used to find the adjustment to the problem variables (degrees of freedom):

$$\begin{split} \mathbf{R}^{i+1} \left( \mathbf{q}^{i} + \Delta \mathbf{q} \right) &= \mathbf{R}^{i} \left( \mathbf{q}^{i} \right) + \mathbf{K} \left( \mathbf{q}^{i} \right) \Delta \mathbf{q} \\ \Delta \mathbf{q}^{i+1} &= -\mathbf{K}^{-1} \mathbf{R}^{i} \qquad \mathbf{q}^{i+1} = \mathbf{q}^{i} + \Delta \mathbf{q}^{i+1} \end{split}$$

Where  $\mathbf{K}$  is a stiffness matrix. Note that The stiffness can be an approximate measure of the real stiffness at a give time, and the solution can still be found. Typically however, any approximations to the stiffness other than the real algorithmic consistent tangent will result in increased iterations, and a reduction in the convergence "radius."

#### Syntax:

The Newton-Rapshon method provides a quadratic convergence: the residual of the problem to be solved can not *theoryticaly* increase. Because real problems usually do not fall into the Newton-Raphson assumptions, one may observe that the redisual increases during iterations. In this case, use the **\*\*line\_search** option to activate the line search procedure. The idea behind it is that if the residual increases, it means that the magnitude of the current Newton-Raphson correction is too large: a search is done along the descent direction to find the largest magnitude so that the residual decreases (one show that this optimum value always exists).

\*\*\*resolution newton
 \*\*line\_search

## \*\*\*resolution bfgs

#### **Description**:

The BFGS algorithm is a quasi-Newton method of matrix updating which can be used to efficiently solve large non-linear problems. The algorithm consists of two methods of convergence acceleration [Matt79]. The first is the updating process which corrects an approximation of the inverted non-linear tangent with the history of trail solutions and their errors. The second is a relaxation of the displacement increment to be compatible with the "direction" of the solution. The algorithm requires an initial calculation of the matrix inverse. Further iterations with the BFGS algorithm do *not* however require additional inversions of the rigidity matrix, making the method very efficient for large non-linear structures.

Given an inverted matrix  $\mathbf{K}_{ini}^{-1}$  which could have been calculated at the beginning of an increment, or just one time at the start of the problem (inverted elastic behavior stiffness), one updates the inverse through the course of iterations as follows:

$$\mathbf{K}_{N}^{-1} = \left[\prod_{j=1}^{m} \mathbf{A}_{j}\right] \mathbf{K}_{ini}^{-1} \left[\prod_{j=1}^{m} \mathbf{A}_{j}^{T}\right]$$

$$\mathbf{A}_i = \mathbf{1} + \mathbf{w} \mathbf{v}^T$$

which shows that the storage requirements are simply that of two vectors of size N (the number of degrees of freedom in the problem). The vectors  $\mathbf{v}$  and  $\mathbf{w}$  are calculated using the changes in displacement and changes in residual between updates.

#### Syntax:

The BFGS algorithm allows the following syntax to adjust the solution process:

```
***resolution bfgs [ iter ][ no_optimize ]
```

The parameter *iter* is an integer specifying the number of iterations which will performed before the BFGS updates begin. The optional token no\_optimize indicates that the optimization (line search) procedure is not desired. The line search adds evaluations of the material law per iteration which may be very expensive for complex material behaviors. *Caution* 

The contact solutions are not compatible with the BFGS algorithm. Also, the use of BFGS requires matrix updating of the types eeeee, p1p1p1 or p1p2p2. Use of p1p1p1 requires the number of iterations *iter* to be greater than one, and p1p2p2 requires *iter* to be greater than two.

# \*\*\*resolution riks

### **Description**:

The **riks** algorithm is a method for which local instabilities can be passed in a quasi-static analysis, such as a bucking situation of a roof collapse. Note that these "snap-though" type problems are in reality dynamic events which can also, perhaps more appropriately, be solved using implicit dynamics.

The method assumes proportional loading as well. So if the load at a point is not merely scaled through time (without change in direction) the algorithm can be used succesfully.

## Syntax:

The syntax for Riks allows a number of keywords and keyword parameter pairs to be entered after the **\*\*\*resolution** command.

```
***resolution riks option-keys
```

Example:

```
***resolution riks
   opt 4 % optimal number of iteration
   max 30 % maximum number of iteration
   div 2. % divergence division factor
   maxDs 0.3 %
   Plus % Keep going positive..
```

#### \*\*automatic\_time

#### **Description**:

This procedure sets automatic calculation of the time stepping (increments) within a sequence. The method will attempt to optimize time steps based on user adjustable convergence and accuracy parameters, and also provide divergence control. The optimization parameters are currently limited to the following:

- optimal number of iterations per increment.
- maximum variation in material variables (e.g. max allowable plastic flow).

The step calculation is made by first examining the integrated material variables on which control has been set. The estimation of the next step size will then be:

$$\Delta t_{i+1} = \Delta t_i \frac{\Delta v}{\Delta v_i}$$

with  $\Delta t_i$  the last increment of time (just solved),  $\Delta t_{i+1}$  the new estimated time step,  $\Delta v_i$  the internal variable increment just achieved, and  $\Delta v$  the desired internal variable increment. This value is calculated for each of the controlled internal variables. The new time increment to be used for the next increment will be the minimum of these  $\Delta t_{i+1}$  values. It is remarked that control may be placed on any of the internal variables existing in the problem. This includes variables defined for only a portion of the structure in a multi-material problem. The problem variables may be confirmed at run-time with the -v command line switch or the **\*\*verbose** output command.

If the time stepping is also controlled by the number of desired iterations, an additional estimation of time step is made as follows:

$$\Delta t_{i+1} = \Delta t_i \sqrt{\frac{nc}{ni}}$$

with ni the number of iterations for the last convergence, and nc the desired number of iterations. All other variables are as described above. The next time increment will be taken as the minimum value calculated from the internal variable control and the iteration control.

As the time step is determined from the previous time step, it remains to define the initial time step in a sequence. This initial step may be given using two different methods. 1) The default method uses the number of increments in the sequence as defined under the **\*\*sequence** command (the **\*increment** sub-command). This method gives the standard  $\Delta t_0 = \Delta t_{sequence}/\#$  increments. 2) Explicitly enter the first time step with the **first\_dtime** sub-command described below.

Divergence control is available when the maximum number of allowable iterations is passed within any of the defined increments. The sub-command **\*divergence** will allow specifying a dividing factor used to reduce the time step. The maximum number of successive divergences is also adjustable.

If the increment of all controlled internal variables is less than the desired amount, or if the convergence is achieved in less than the desired number of iterations, the time step will be increased by a user defined factor (\*security) in order to accelerate the solution.

#### Syntax:

The following syntax is used to define the automatic time-stepping in a calculation:

```
**automatic_time [ type ] ( var1 val1 [ var2 val2... ] | global iter | mandatory)
[ *divergence div [ times ] ]
[ *security ratio ]
[ *max_dtime max_dtime [ max_dtime2... ] ]
[ *min_dtime min_dtime [ min_dtime2... ] ]
[ *first_dtime first_dtime [ first_dtime2... ] ]
```

CODE	DESCRIPTION
standard	Automatic time assuming that the loading path is basi- cally continuous. If mandatory is specified the indicated incremental value is considered to be mandatory: if the increment of the specified variable is too large, one consid- ers that a divergence occured, and the automatic time will reduce the time step.
by_sequence	The algorithm is initialized to the user's increment input for each sequence; recommended for cyclic loading for example
divergence_control	Automatic time only to control divergence; the input incre- ments will be used except when a divergence occurs

- \*divergence div [ times ] Divergence control taking a real value for the devising factor div and an integer for the number of allowable divergences times. The default value for times is one.
- \*first\_dtime first\_dtime Set the first time step in all sequences with the real value first\_dtime. This option is advised only for monotonic or other simple loading paths. A list of first\_dtimes may also be given, corresponding to the first dtime of each sequence. The last is implicitly repeated, if necessary.
- \*max\_dtime max\_dtime Set the maximum allowable time step to the real value max\_dtime. In the case of overestimating the number of iterations or change in the internal variables, this option will limit the acceleration of solution.
- \*min\_dtime min\_dtime Set the minimum time step to the real value min\_dtime. In the event where the number of iterations or the change in internal variables are grossly underestimated, this option will limit the time step to a minimum value. It prevents extreme reduction of the time step which will effectively limit the calculation from advancing further.
- **\*security** *ratio* Defines the multiplicative factor used to increase the time step in the event of good convergence or small changes of the internal variables. The parameter *ratio* is a real value specifying the factor to be used.

If automatic\_time exhibits strange behavior with small or large time steps, try to use \*\*\*dimension instructions 3.106.

#### Example:

A very frequent use of this command follows:

\$

\*\*automatic\_time global 3
\*divergence 2.0 10
\*security 1.2

Frequently it is a good idea to keep the allowable iterations small (\*\*sequence \*iteration command), and let the automatic time reduce the step size more readily.

Some other variations to control accuracy based on changes in material variables follow.

```
**automatic_time evcum 0.001 eel11 0.001 global 5
```

```
**automatic_time f 0.01 p 0.01
    *divergence 2.0 50
    *security 1.2
    *max_dtime 0.03
    *min_dtime 0.001
```

# \*\*init\_d\_dof

## **Description**:

This procedure sets the initial increment DOF values for iteration i + 1 as a function of the previous solution i:

$$\Delta \text{DOF}_{i+1} = \text{ratio} \times \frac{\Delta t_{i+1}}{\Delta t_i} \times \Delta \text{DOF}_i$$

The objective of this method is to accelerate convergence during relatively stable loading paths. The assumption is that the next solution will not vary greatly from the previous one. This method is therefore dangerous to convergence if the loading changes rapidly, such as a newly unloaded section of the structure.

# Syntax:

\*\*init\_d\_dof [ ratio ][ sequence ]

ratio is a real value acting as a scaling factor, as shown above. The default value is 1.0.

**sequence** indicates that the initialization is only applied within each sequence, based on the first solution of that sequence. In absence of this option the initialization is carried out for the entire calculation, even across sequence boundaries.

# Example:

\*\*init\_d\_dof 0.9 sequence

\*\*\*\*calcul
\*\*\*resolution
\*\*max\_divergence

# \*\*max\_divergence

## **Description**:

This procedure allows a given value to be set for the maximum allowable ratio of two successive convergence ratios. The calculation will be terminated if this critical ratio is exceeded.

## Syntax:

\*\*max\_divergence value % value is a double

## Example:

\*\*max\_divergence 1.5

## \*\*sequence

## **Description**:

This command writes the sequences or blocks of time steps for the problem<sup>11</sup>. Solution requires specification of discrete times for which the governing equations are to be evaluated. The program allows an unlimited number of **\*\*sequence** or **\*\*cycles** (see next command page) instances in order to create complex loading histories. There is really no distinction between the sequence and the cycle commands; the cycle command automatically generates a number of repeated sequence groups to form cycles.

As discussed in the introduction of **\*\*\*\*calcul** (page 3.7), the sequences are broken up into increments which are the most fundamental unit of time stepping.

### Syntax:

Sequences are created as necessary according to the sub-options of the **\*\*sequence** command. Normally, the sequences are defined by end times or incremental times of the sequences. The syntax for these definitions is the following:

	**sequence	[	N ]
[	*algorithm		algo1 [ algo2, algoN ] ]
[	*increment		inc1, inc2,, incN ]
[	*iteration		iter1, iter2,, iterN ]
[	*limit_dof		vari, vali1,valin [, varj, valj1,, valjN ] ]
[	*time		time1, time2,, timeN ]
[	*dtime		dtime1, dtime2,, dtimeN ]
Γ	*ratio		[ absolu ] ratio1, ratio2,, ratioN ]

The integer value N defines the number of sequences which will be run, even if there are more defined in the sub-commands. This allows the run-time of a problem to be limited quickly, without deleting or commenting information which may be useful later on. In the absence of this value, the number of sequences will be determined based on the number of time or dtime values input.

All the other options are used for specifying particular parameters which correspond to the sequences. If the number of values given after an option is less than the number of sequences of the current block (the last **\*\*sequence** command), the last value given will be repeated for all the subsequent sequences. If one or several of the sub commands are absent, the default value will apply to all the sequences defined.

\*algorithm String values substituted for algo1 to algoN are the algorithm solution methods for each segment of the loading. The values for these methods are summarized:

 $<sup>^{11}\</sup>mathrm{Sequences}$  are extensively discussed in the examples manual

CODE	DESCRIPTION
eeeeee	Newton "elastic" method. Matrix is not re-calculated during iterations
EEEEE	Linear method. No iterations, matrix calculated only once (once after time step or Newmark coefficients changes in dy- namics). The computation of internal forces is made through the K.U multiplication which is much faster than the classical element integration procedure. Only nodal fields are post- treated, no element fields are available. This can be useful to treat linear dynamics problems in a very fast way.
p1p1p1	Modified Newton algorithm with the matrix calculated and inverted the first and second iterations
p1p2p2	Modified Newton algorithm with the matrix calculated and inverted the first, second, and third iterations
p1p2p3	Newton-Raphson algorithm where the matrix is always up- dated during iterations

- \*increment The integer values *inc1*, *inc2*, ..., *incN* are used to define the number of increments in corresponding sequences. In the absence of this option, all sequences are calculated in a single loading increment.
- \*iteration *iter1*, ..., *iterN*. Integer values used to limit the maximum number of iterations in a given increment, valid during a sequence. The default number of iterations is 10. These values may be used to adjust the divergence control, or limit the CPU consumption of a poorly converging problem.
- \*limit\_dof vari, vali1,...valin, ... This option takes a DOF character name (U1, U2, etc) and a list of real values corresponding the given segments. Any number of DOF name / list of limiting values may be input sequentially to put limits on the different DOFs of the problem. Real numbers input are limiting values for any iteration of the degree of freedom. In the course of iterations, we will have available  $\Delta \mathbf{u}_i$  at iteration i + 1 from the solution of the linear system,  $d\mathbf{u}_i = (\mathbf{K}_i)^{-1} \mathbf{R}_i$ . In place of the standard updating of the DOF values,  $\Delta \mathbf{u}_{i+1} = \Delta \mathbf{u}_i + d\mathbf{u}_i$  we have  $\Delta \mathbf{u}_{i+1} = \Delta \mathbf{u}_i + \beta d\mathbf{u}_i$  where  $\beta$  is a limiting factor:

$$\beta = \min\left(\frac{\text{limitation value}}{\max_{DOF}(d\mathbf{u}_i)}, \beta\right)$$

with  $0 < \beta < 1$ .

This option allows the calculation to pass points where the algorithm would predict instability, at the cost of increased number of iterations. Excessive trial delta values in the DOFs may occur when the problem tangent matrix displays softening behavior.

\*time time1, time2,..., timeN specifies the absolute times at the end of the sequences. There are as many values as the number of sequences. The first sequence of the problem is at

t = 0, or the last time given in the previous **\*\*sequence** command. This initial value must be less than the first time *time1* given here. In the absence of a *\*time* option, the times will be assigned to the sequence numbers.

- \*dtime dtime1, dtime2,..., dtimeN specifies the increment of time taken over a sequence. This method of specifying the sequence times is useful if a "time shiftable" block of sequences is desired. An example is a block of loadings to be subjected to different pre-loadings, or loading after a series of cycles, where additional computations would be necessary to assure that the times are well-posed. Options \*dtime and \*time are mutually exclusive.
- **\*ratio** [absolu] ratio1, ratio2, ..., ratioN This option is for specifying the maximum global residual for convergence in a sequence. The real values ratio1, etc correspond to the sequences defined. A single given value will be repeated for all the sequences.

The residual calculation will be calculated according to the following formula:

$$\begin{aligned} \text{ratio} &= \frac{\sqrt{\sum_{\text{iddl}} \left(R_{\text{iddl}}^{\text{ext}} - R_{\text{iddl}}^{\text{int}}\right)^2}}{\sqrt{\sum_{\text{iddl}} R^{\text{ext}^2}}} & \times 0.01 \text{ ratio} \\ \text{ratio} &= \sqrt{\sum_{\text{iddl}} \left(R_{\text{iddl}}^{\text{ext}} - R_{\text{iddl}}^{\text{int}}\right)^2} & \text{ratio absolu} \end{aligned}$$

where the option absolu indicates non-normalized ratio calculation.

#### Example:

```
**sequence
                          5.0
  *time
                  1.0
                  10
  *increment
  *iteration
                  20
  *ratio absolu
                  1.0
  *algorithm
                  p1p2p3
                             U2 .02 .03
  *limit_dof
                  U1 .1 .1
%
% from ab429.inp making 3 sequences
%
**sequence
               3.
                     33.
                          50.
  *time
  *ratio
               1.
                      3.
                           3.
  *iteration
               50
                     30
                          30
                     15
  *increment
               10
                          34
  *algorithm
               p1p2p3
  *limit_dof
     U1
           0.2
                 0.5
                        0.5
     U2
           0.2
                 0.5
                        0.5
         10.0
                      10.0
     PR
                10.0
```

\*\*cycles

# **Description**:

The cycles option is essentially the same as the **\*\*sequence** option except that it repeats the block of sequences a given number of times. Normally, use of the **\*\*cycles** option will be employed after a series of pre-loading sequences, in order to attain the initial loading state of the cycles. Therefore, cyclic loadings will usually require a **\*\*sequence** definition for the pre-load, followed by a **\*\*cycles** definition for the cycles themselves. Unless the end segment of the cycles will return to zero loading, additional **\*\*segment** definitions may be given to relax the structure to an unloaded state.

An example loading is a pre-load in tension, followed by tension-torsion cycles, and finally a relaxation of the initial tensile load. Many more cases may be imagined.

As the calculation time scale is the principal measure of the solution, it is important to understand the method of time scale construction from a cycles definition. The times given for this option are always relative to the beginning of the cycle. This defines a local cycle-based time scale, which may be repeated.

# Syntax:

The syntax for cyclic loading is the following:

	**cycles [ N	1]
Ε	*algorithm	algo1 [ algo2, algoN] ]
Ε	*increment	inc1, inc2,, incN]
Ε	*iteration	iter1, iter2,, iterN]
Ε	*limit_dof	vari, vali1,valin [, varj, valj1,, valjN] ]
Ε	*time	time1, time2,, timeN]
Ε	*dtime	dtime1, dtime2,, dtimeN]
[	*ratio	[ absolu ] ratio1, ratio2,, ratioN]

the integer N is the number of cycles to be created.

Except for the time scale alteration for the cyclic sequences defined with the **\*time** or **\*dtime** options, all sub-commands for this command are equivalent to the **\*\*sequence** command.

# Example:

An example loading gives 5 cycles with the waveform 10s-300s-10s after a pre-load of 1s.

```
**sequence
 *time 1.
**cycles 5
 *dtime 10. 300. 10.
 *increment 5 10 1
 *algorithm p1p2p3
```

This example assumes the first one second loading segment will be a linear solution. The first segment of the first cycle (from 1 second to 11 seconds on the "global" clock) may be a loading with non-linear behavior, so 5 increments are given. The following rest time has 10 increments, also assuming non-linear behavior. The final segment from times 311 seconds to 321 seconds is assumed linear and will be solved in one increment. The same increments will

then be repeated 4 more times. A corresponding loading waveform is given in the **\*\*\*table** command.

\*\*\*\*calcul \*\*\*resolution \*\*skip\_cycles

# \*\*skip\_cycles

## **Description**:

The skip cycles command is used for cyclic calculations to extrapolate the problem variables and thereby reduce the number of cycles actually calculated. Because there can be many time steps per cycle, this command has the possibility of greatly reducing the calculation time.

# Syntax:

```
**skip_cycles [ type ]
*precision val
*during_cycles cyc-st cyc-end
*use_sequence seq-num
*extrapolate_from cyc1 cyc2 cyc3
*check_with_component var-name
```

# Example:

For example we give a viscoplastic problem with 300 cycles and several sequences per cycle. The option **\*use\_sequence** was used to specify the beginning of sequence 1 because the material response is pretty much elastic there throughout the cycling. The response changes rapidly before cycle 4, so **\*during\_cycles** was used to start the cycle skip after that point.

```
**cycles 300
 *time
            15.0 30. 40.0 50.0
 *increment 20
                 4
                     5
                           4
 *ratio absolu 1.e-3
 *algorithm p1p2p3
**skip_cycles polynomial_extrapolation
 *precision
                0.2
 *trim
                0.8
 *order
                2
 *error_skip
                2
                    290
 *during_cycles 4
 *use_sequence 1
 *extrapolate_from 1 2 3
 *check_with_component evcum
```

# $**use\_lumped\_mass$

# **Description**:

This procedure replaces the mass matrix by the lumped one currently used with explicit algorithm (see \*\*\*\*calcul mechanical\_explicit in 3.15). Only available for \*\*\*\*calcul dynamics.

# \*\*\*restart

# **Description**:

Manages the restart of an interrupted or unfinished calculation. The program automatically saves a restart file (filename prob.rst) at the end of each increment. If special restart saves are desired at different stages of the calculation, use the **\*\*\*make\_restart\_file** option. Saving a calculation's .rst file may be used to repeatedly calculate a second part of a calculation without repeating the initial portion. The option may also be used to change the parameters of the calculation at intermediate points in the time-scale, although with significant user work.

# Syntax:

\*\*\*restart
[ \*\*file name-of-restart-file ]

The default restart file name is *prob.rst*; you may specify a different name with the **\*\*file** option.

Drastically changing the boundary conditions or other parameters in the .inp file and re-starting the calculation may lead to poor results.

# \*\*\*auto\_restart

# **Description**:

This option allows automatic restart of a problem, if a restart file exists. The only difference with **\*\*\*restart** is that no error message is output if no restart file is available.

# \*\*\*table

## **Description**:

The procedure **\*\*\*table** indicates a loading table definition in time. The table will be available in the other commands of the .inp file such as the boundary conditions or the external parameter definitions. Loading tables are currently only defined as a series of time-magnitude points which are linearly interpolated to intermediate times.

The tables are named with arbitrary, user-defined character strings which are used in the syntax of other commands to indicate which table to use. There are no conditions on the number or size of the tables.

# Syntax:

Loading tables are assembled through the use of the following sub-commands:

```
***table

**name name

*time t_1 \dots t_n

*value v_1 \dots v_n

**cycle name t_{ini} t_{end}

*time t_1 \dots t_n

*value v_1 \dots v_n

**function name FUNCTION

**file name file_name ctime cvalue
```

- **\*\*name** Specifies that a simple table is to be created with name as given directly following the **\*\*name** keyword. A table defined as such is assembled with the **\*time** and **\*value** options.
  - \*time indicates that a list of real values will follow giving the individual time points of the table. These time values are in absolute time measured from the beginning of the problem. The current version of the code requires in most applications that the table be able to give a value for all valid times in the problem. This is noted to begin with the initial time t = 0. Most tables will therefore be defined starting with this initial point.
  - \*dtime This option serves the same function as the \*time command, but is defined in incremental form. The option is useful for complex loadings where it is easier to think in incremental form, or as the only way to currently model step loading (using segments of  $\Delta t = 0$ ).
  - \*value This option indicates that a series of real numbers will be given to define the table values. It is necessary that there be exactly one value for each time specified with the \*time command. As the time definitions are given from the beginning of the problem, there must be a corresponding initial value for time t = 0. This value will normally be zero in the case of boundary condition loading, as the problem may not begin in a pre-deformed state through standard boundary conditions (use a restart procedure with the pre-deformed structure stored in the restart file). If one table is succeeded by another, it is necessary to ensure the continuity from one table to another.

- **\*\*cycle** name  $t_{ini}$   $t_{end}$  In order to generate cyclic loadings it is convenient to simply specify a load cycle to be repeated a number of times. This option does exactly that with the initial time for the cycles given by  $t_{ini}$  (a real value). The cycles will continue repeated until the time  $t_{end}$  (real) regardless if that time is at the end point or an intermediate point in the cycle. The cycle is defined as a simple table above using the **\*time** and **\*value** options, only that the time will be in a local cycle scale. The start of the cycle (first at time  $t_{ini}$ ) is at the local time zero. The zero time point must therefore always be given for the cycle table format. In contrast with the standard table, the corresponding initial value is not usually zero, as some pre-loading may have been applied before the cycles begin. This command is examined more fully in the examples below.
- **\*\*function** This allows to generate tables defined by functions depending on time. name specifies the table name followed by a FUNCTION object. The variable time MUST be the only argument of the function.
- **\*\*file** This allows to generate tables defined by a file (column format). *name* specifies the table name. *file\_name* specifies the file name. *ctime* specifies the column of *file\_name* representing the time and *cvalue* the column of *file\_name* representing the table value (the first column is numbered 1).

# Example:

1. Suppose that a U2 displacement is to be applied to the top surface of a structure. This surface will have a node set defined by the user in the geometry description with the name top. The loading profile is to be:

at $t = 1$ ,	u = 10.
at $t = 5$ ,	u = -10.
at $t > 5$ ,	u = -10.

On a face set named inter a pressure is also to be applied. The pressure value will be zero until t = 1, after which there is a ramp loading in the pressure until t = 7 at which point the pressure is 100.

The loading sequences will correspond to the points of load change:

```
**sequence
```

\*time 1. 5. 7.

Declaration of the displacement and pressure boundary conditions are written:

\*\*\*bc

```
**impose_nodal_dof
    haut U2 1. tab1
**pressure inter 100. tab2
```

Note the two different table names tab1 and tab2 because the loading waveforms are different. The table definitions may be written as:

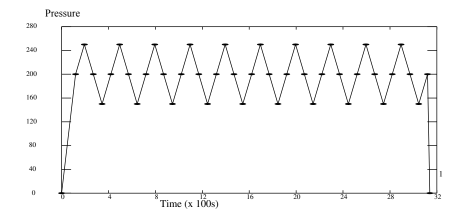
\*\*\*\*calcul \*\*\*table

```
***table
 **name tab1
 *time 0. 1. 5. 7.
 *value 0. 10. -10. -10.
 **name tab2
 *time 0. 1. 7.
 *value 0. 0. 1.
```

2. Cyclic loading is given as a second example. This case represents the calculation of a structure under a pre-loaded pressure of 200 MPa followed by a cycling of the pressure about the mean value of 200 MPa with a cyclic magnitude of 100 MPa. After 10 cycles the pressure is returned to zero to observe the residual stress field.

```
***resolution
   **sequence
                 120.
    *time
                 5
    *increment
   **cycles 10
    *dtime
                 75. 75. 75. 75.
                10
    *increment
    *algorithm
                p1p2p3
   **sequence
    *dtime
                 20.
    *increment
                 5
  ***bc
   **pressure
      top 1.0 tab1 cycs tab2
  ***table
   **name tab1
    *time
               0.0 120.
               0.0
                     200.
    *value
   **cycle cycs 120. 3120.
    *time
               0.0
                      75. 150. 225. 300.
    *value
             200.0
                     250. 200. 150. 200.
   **name tab2
    *time
            3120.0
                     3140.
    *value
             200.0
                       0.
```

\*\*\*\*calcul \*\*\*table



# \*\*\*function

# **Description**:

This command applies as **\*\*\*table** in the preceeding section, but defines the magnitude as a function of time (see page 5.2 for a discussion of functions).

The **\*\*\*function** command can only be defined in terms of the variable **time**.

# Syntax:

The syntax is the following:

**\*\*\*function** name func\_def;

Note that semi-colon is **required**. The name has the same significance as in the **\*\*\*table** command. Each function requires its own **\*\*\*function** declaration.

# Example:

\*\*\*function xtab cos(time); \*\*\*function ytab sin(time); \*\*\*function funny time + 3.0\*exp(-8.2\*time); \*\*\*function constant 1.0;

The last example will give a warning, but works.

# \*\*\*specials

# **Description**:

The command **\*\*\*specials** allows one to specify certain special boundary conditions. Currently, the command may only be used to declare mesh conditions which exist in the mesh, and is thus could be thought of as a "mesh modifier<sup>12</sup>."

# Syntax:

\*\*\*specials
[ \*\*mesh ]
[ \*specify ]

**\*\*mesh** is used to specify special conditions which exist in the mesh.

\*specify Introduces the specified parameters to the mesh. The only option currently available for this command is X0\_Y0\_for\_E2\_5 which fixes the origin of moments applied to 2.5D elements (generalized plane strain). The moments are applied throughout the thickness of an elset. The syntax requires an element set name follow the keyword, and also real values for the center on the axes 1 and 2. This option is associated to the BC command \*\*impose\_elset\_dof.

# Example:

```
***specials
**mesh
*specify X0_Y0_for_E2_5 eprouvette 1. 1.
```

 $<sup>^{12}\</sup>mathrm{I}$  really don't understand what this is - rf

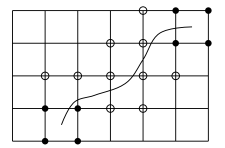
# \*\*\*xfem\_crack\_mode

# **Description**:

This command allows the use of XFEM enriched elements, in order to model the influence of a discontinuity (crack) on mechanical fields, without the need to explicitly introduce this crack in the FE mesh. Note that the graphical interactive **Zxfem** script can be used to handle Xfem models, and automatically generate input data for this command.

In this context the crack is defined by 2 levelsets,  $\phi$  (signed distance to the crack plane) and  $\psi$  (signed distance to the front of the projection on the crack plane). The **\*\*compute\_predefined\_levelset** mesher command can be used to generate such levelsets for basic crack geometries (see page 2.31. Let *I* be a set containing all nodes in the FE mesh, we then define 2 subsets *J* and *K* of *I* (see figure below):

- $J \subset I$  correspond to nodes in elements E completely cut by by the crack ( $\phi$  is changing sign on E, but sign of  $\phi$  is constant),
- $K \subset I$  correspond to nodes in elements that include the crack front (both  $\phi$  and *psi* are changing sign on *E*).
  - *J* set: nodes  $\in$  elements cut by the crack
  - *K* set: nodes  $\in$  elements containing the crack front



Enriched shape functions are then defined in the following way:

$$u^{E}(x) = \sum_{i \in I \cap E} u_{i} \phi_{i}(x) + \sum_{i \in L \cap E} a_{i} \phi_{i}(x) H_{i}(x) + \sum_{i \in K \cap E} \phi_{i}(x) \left[ \sum_{j=1}^{4} b_{i}^{j} F_{i}^{j}(x) \right]$$

where:

- $\phi_i(x)$  denote the shape functions of classical (ie. not enriched) finite elements and  $u_i$  the conventional displacement degrees of freedom ( $i = 1, N_E$ , where  $N_E$  is the number of nodes of element E)
- $H_i(x)$  are shape functions taking a value of 1 on one side of the crack, and -1 on the other. They are used to model the discontinuity of displacement fields across the crack. The corresponding  $a_i$  degrees of freedom are called Xh in results files.
- the  $F_i^j(x)$  (j = 1, ..., 4) shape functions account for classical elastic analytical solutions around a crack tip:

$$(F^1, F^2, F^3, F^4) =$$

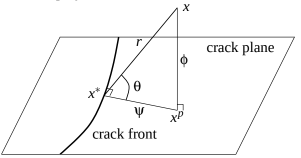
Z-set — Non-linear material & structure analysis suite

$$\left(\sqrt{(r)}\,\sin\left(\frac{\theta}{2}\right)\;,\;\sqrt{(r)}\,\cos\left(\frac{\theta}{2}\right)\;,\;\sqrt{(r)}\,\sin\left(\frac{\theta}{2}\right)\;\sin(\theta)\;,\;\sqrt{(r)}\,\cos\left(\frac{\theta}{2}\right)\;\sin(\theta)\right)$$

where  $(r, \theta)$  are polar coordinates defined around the crack front as defined on the next figure:

 $x^p$ : projection of x on the crack plane

 $x^*$ : projection of  $x^p$  on the crack front



• the  $b_i^j(j=1;4)$  degrees of freedom associated to the  $F_i^j$  shape functions will be called Xa, Xb, Xc, Xd in results files.

Note that the basic enrichment scheme described above is the so-called *topological enrichment* mode. A major problem associated with this mode, is that the influence of the refined shape functions used for nodes in the K set ( $\sqrt{r}$  shape functions), is decreasing when mesh size near the crack front is refined, which has a negative effect on the rate of convergence. If this is indeed a problem, *geometrical enrichment* can be used, where all nodes within a target user-defined distance of the crack front are automatically added to the K set, and are given full  $\sqrt{r}$  type of enrichment.

In the current Zébulon release (8.5), some restrictions on the use of this method still exist, that will be removed in the next versions:

- only 3D linear elements are currently officially supported,
- extensive capabilities do exist to use this scheme within the context of crack propagation (with automatic remeshing if needed), but those options are still under developemnt and will not be addressed here.

#### Syntax:

The syntax is as follows:

```
[ ***xfem_gtheta ]
 ***xfem_crack_mode
[ **elset ename ]
 **discontinuity 3d_levelset
  ( no_option |
       (
       [ *psi_file fpsi ]
       [ *phi_file fphi ]
       [ *fit_to_vertice eps ]
```

```
)
)
[ **geometrical_radius rad ]
[ *set_mpc ]
```

where:

- keyword \*\*\*xfem\_gtheta is mandatory for connection with the \*\*\*compute\_G\_by\_gth command (computation of 3D stress intensity factors by means of the  $G \theta$  method 3.86).
- optional command **\*\*elset** ename may be used to specify the name of an elset containing candidate elements of the enrichment process. By default, all elements in the FE mesh are concerned, and are indeed enriched when cut by the crack or located within the geometrical radius definition.
- commands **\*phi\_file**, **\*psi\_file** allow to specify the name of ASCII files defining the  $\phi, \psi$  levelsets on nodes of the FE mesh (see 2.31 for the mesher command used to create those files). Default names for those files are "**phi.dat**", "**psi.dat**".
- the optional command \*fit\_to\_vertices may be used to define the critical distance value eps, that controls if the crack does cut an element (ie. the element needs to be splitted), or simply pass through one of its corner nodes (vertice). Note that eps is a *relative* value, scaled against the size of the element edge cut by the discontinuity (default value is eps=0.05, ie. 5% of the length of the corresponding element edge).
- command **\*\*geometrical\_radius** specifies the size rad of the domain used for geometrical enrichment: int his case, all elements with nodes within a distance of less than rad of the crack front will be enriched with  $\sqrt{r}$  shape functions. This option enhance the rate of convergence of the xfem method (mesh size sensitivity), but should be used with caution since the size of the problem is strongly increased (12 additional dof by nodes concerned in 3D). Default mode is topological enrichment only (rad=0).
- when geometrical enichment is activated, the optional **\*set\_mpc** may be used to automatically add linear relationship between the geometrically enriched zone and the rest of the mesh, in order to insure continuity of displacements, and improve the rate of convergence.

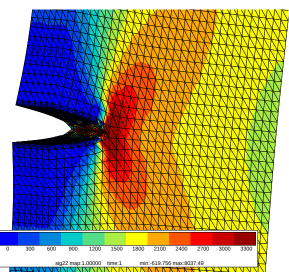
# Example:

% Basic example : calculation with xfem enriched elements
\*\*\*\*calcul
% activation of the Z8 output format mandatory
% for Zmaster handling of XFEM results
\*\*\*global\_parameter
Solver.OutputFormat Z8
Zmaster.OutputFormat Z8
...
% levelset definition in default phi.dat, psi.dat files
Zest\_Nonlinear material

```
***xfem_crack_mode
 **discontinuity 3d_levelset no_option
...
% special xfem output requests allowed in the Z8
% output mode
***output
 **xfem_split_integ
 **xfem_split_node
 **xfem_split_contour
...
****return
```

### Example:

```
% use with the g-theta module
****calcul
...
% mandatory to link xfem with gtheta
***xfem_gtheta
***xfem_crack_mode
**discontinuity 3d_levelset
...
% gtheta commands for SIF calculation
% (note the **xfem keyword)
***compute_G_by_gth gtheta_a
**xfem
**behavior paris
...
****return
```



Example of Zmaster capabilities to handle XFEM results and draw the solution on a *cracked* mesh as if the discontinuity was explicitly introduced in the model

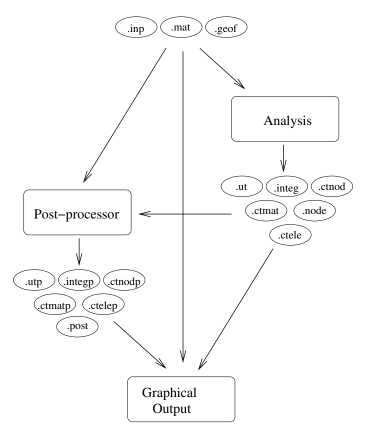
# Chapter 4

# Post calculations

# Introduction to Post Computations

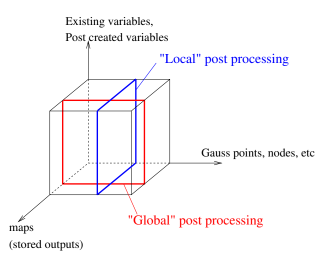
This mode of operation is used to apply post calculations on results obtained by the finite element method, and with simulations. Because the data input for post processing are in the calculation results files, the file name resulting from the post-processor are constructed from the original name, by adding **p** to the different file name extensions. For example, the treatment of integration points in the problem *problem.integ* will produce a file named *problem.integp*. A file *problem.utp* is automatically generated in order to allow graphical visualization of the new variables.

All of the results (calculation and post-calculations) are simultaneously available in the Zmaster graphical program, and the Zmaster batch program.



Two basic types of operations are defined:

- A post treatment so-called *local* which is used to evaluate a criterion at nodal or Gauss point locations throughout the mesh, or in a part of it. The calculation is based only on the data contained at the point(s) of concern.
- Post calculations which are *global* in that they are used to evaluate a criterion over the whole structure, or in a sub-part of it, at a given instant. The calculation therefore normally accounts for data distributed in space.



#### Local post-processing:

The local post computations is used for example to predict the component life of a structure using simple monotonic loading (e.g. Rice-Tracey models), under creep or fatigue (low or high cycle models). It can also used used to determine some derived data relative to the local history at each point, such as the maximum stress temperature or temperature during the loading history.

According to the users choice, the computation will be made at integration (Gauss) points (from the file *problem.integ*) or at nodal points from data originally saved at the nodal points (files *problem.node* and *problem.ctnod*).

The local post treatments give the user the ability to make connections between Z-set computations and other codes as well. This can be achieved for example using the format command to create formatted ASCII text output, or using and external program or script to calculate a particular user post computation. Note also that the post-processor is fully extensible using the Z-set plug-ins.

### <u>Global post-processing</u>:

Global post processing is applied when, for example, one wishes to calculate the mean values of a variable in the entire structure. The majority of the time, the structure's geometry must be known, and an integration volume is defined. Another example is a Weibull criterion for brittle fracture prediction.

The majority of global post computations produce output which is reduced to a scalar value for each time step of the calculation. These values are stored with some description in an ASCII file *problem.post*. Nevertheless, some of the global post processors produce local results (all the time using a non-local algorithm), which will be stored in the appropriate file (node or integration values).

## General rules:

For the two types of post-calculation, the user defines the *context* operation: groups of elements, of nodes, or of Gauss points on which to calculate. The output numbers may also be taken into account. The context can be redefines during the calculation. It is important to remember that a given computation or criterion will be applied to the last context which was defined. The individual computations will be applied one after another in the order of appearance in the input file. One can alternate arbitrarily local and global post computations.

The variables for which the post computation is applied can be:

- FEA output: their names appear in the file *problem*.ut
- Post-computation output calculated in the course of the same execution., That is, once a post computation is applied, the new variables generated by that application are henceforth added to the list of available variables.

Recursive computations are available using the newly generated variables in postcomputations. An example would be to calculate an equivalent strain as the combination of strains at each point, and then calculate the maximum of these equivalent strains. The global results (stored in the file *problem.post*) are not however re-usable.

## The data source:

In version 8.3 and greater Z-post has the capability to run off of different data source formats than the default Zebulon solver files. Notably there is the possibility to import from ABAQUS, ANSYS Z-sim and arbitrary ASCII files. With this mesher commands can be applied to the results files as necessary to allow new selections to be available. The commands to do that import are described in the section for **\*\*\*data\_source** on page 4.14. An example showing typical use follows.

```
****post_processing
 ***data_source rst
                                        % load the ansys results
   **open t-base_model_fillet.rst
                                        % and write a native
                                        % mesh file (GEOF format)
   **write_geof
   **elset LOOK
                                        % with some set generation
    *elements
      2205 2163 311 325 339 353 381 367 395 409 423 437 479 521
   **nset edge1
    *nodes
   2396 2397 1 2394 2395 2393 2392 2390 2391 2389 2388 2386 2387
    2385 2384 2382 2383 2381 2380 2379 2373
   **bset edge1
   *use_nset edge1
    *use_dimension 2
 ***precision 3
 %-----
                      _____
 ***local_post_processing
                                        % here we copy the displacements
     **output_number 1-999
                                        % so the post generated views
                                        % can be deformed
     **file node
     **output_to_node
     **nset ALL_NODE
     **process copy *list_var U1 U2 U3
 ***local_post_processing
                                        %
   **output_number 1-999
                                        % Ansys results are contours
   **file ctele
                  **elset ALL_ELEMENT
                                        % by element (CTELE)
                                        % where to find coefs.. (here we
   **material_file post1.inp
                                        % use the same physical file)
   **process transform_frame
    *tensor_variables sig
                                        % "sigma-material"
    *output_variables sigm
    *use_element_rotations
 ***return
```

## The data output:

×

In version 8.3.3 and greater Z-post has the capability to output to a number of different output formats, with multiple selections of localized output formats from your posts.

In every case, the standard "-p" files will also be written because these files are used for access to intermediate variables in the course of several posts relying on each other. An example use of this post processing with an ansys input and abaque output follows:

****post_processing	
***data_source rst	% load ansys rst format
**open ansys_model.rst	% in the file ansys_model.rst
***data_output odb	
<pre>*problem_name abaqus_post_results</pre>	% makes abaqus_post_results.odb file
<pre>*elset outer_surface</pre>	% make odb of this elset only
***local_post_processing	
•••	

With the release of Z 8.3.6 in March 2006 the data outputting also consist of a format for zebulon files. This output format has following additional features which may be preferable in some cases:

- the data is written to disk only in a single large write, without seek calls which can cause network latency.
- the output can be specified to a list of elsets, from which a reduced mesh is given. this data output can therefore be used to reduce storage size.
- can be used to write to a "zebulon" calculation format (non-p suffixed files) and therefore can form the basis of a subsequent series of post calculations.

## Interface with Zmaster:

The post calculations has been embedded in Zmaster in addition to their use by generating additional output files. That is, the post process computations can be run directly on the selected data in Zmaster. After adding a post computation in the Zmaster environment, the .mast file needs to be re-saved.

### Material coefficients:

Many of the post computation models use coefficients to specify parameters which vary for different materials. There are 2 ways of specifying these coefficients, each of which may be more convenient for different situations.

The first method is to have the material data separated from the process definition. This allows re-use of the coefficients for multiple computations by creating a repository of material values. An example input with two creep entries follows.

```
**material_file creep.inp
**process creep
 *var sig
**process creep
 *var sig
 *express_life_as time
```

Both creep process computations would then use the creep entry in a material file creep.inp:

```
***post_processing_data
    **process creep
    r 5.
    A 1500.
    S0 0.
***return
```

Another method (new to Z8.0) is to simply put the coefficients in-line with the process command. This is obviously easier for "one-off" post processing:

```
**process creep
 *var sig
 *model_coef
 r 5.0
 A 1500.0
 S0 0.0
```

#### Pointers and stacked post computations:

Like all of the Z-set packages, the post processing software is build on objects which perform specific tasks. In the post case however there are a large number of cases where certain calculations such as a thermo-mechanical fatigue analysis employing plastic range, oxidation, and creep damages will need to re-use other computations which themselves can act as standalone computations. This is to say, very often a post computation will require pointers to other post input files defining those additional procedures. The following is a detailed example of such input (from the test lcf.inp in Post\_test/INP

The analysis is defined by the "primary" section, which will be the first instance of **\*\*\*\*post\_processing** in the input file (or *N*-th instance in the case of an -N input switch).

```
****post_processing
   ***local_post_processing
       **file integ
       **elset ALL_ELEMENT
       **material_file ../MAT/test_simple
       **process LCF
                                 \% LCF auto creates a stress range making NC_S NF_S
         *mode NLC_ONERA
                                 % creates NR_NLC_ONERA
                                 % this means use fatigue_S post
         *fatigue fatigue_S 2
                            2
                                 % and creep post in post .inp file #2
         *creep
                  creep
       **material_file ../MAT/test_with_a
       **process LCF
                                 % there is already NC_S NF_S, new are NC_S_n1 NF_S_n1
         *mode NLC_ONERA
                                 % creates NR_NLC_ONERA_n1
         *fatigue fatigue_S 3
                                 % look in 3rd ****post_processing segment
         *creep
                creep
                            3
       **material_file ../MAT/test_simple_norm
       **process LCF
                                 % now NC_S_n2 NF_S_n2
         *mode NLC_ONERA
                                 % NR_NLC_ONERA_n2
         *fatigue fatigue_S 4
         *creep
                 creep
                            4
       **material_file .../MAT/test_with_a_norm
       **process LCF
                                 % and finally NC_S_n3 NF_S_n3
                                 % NR_NLC_ONERA_n2
         *mode NLC_ONERA
         *fatigue fatigue_S 5
         *creep
                  creep
                            5
   ***global_post_processing
                                 % average values are taken for the validation
       **output_number 1
                                 % purposes. One can look at a field in Zmaster
       **process average
         *list_var NC_S NF_S NF_S_n1 NF_S_n2 NF_S_n3
       **process average
         *list_var NR_NLC_ONERA NR_NLC_ONERA_n1 NR_NLC_ONERA_n2 NR_NLC_ONERA_n3
****return
```

Note that the active material file is being switched throughout the input structure, and each calculation uses a pointer for the **\*fatigue** and **\*creep** parts. That is, the LCF model is a method of combining fatigue and creep damages (with for example the nonlinear combination method of ONERA), and the ways that those damages are calculated is defined elsewhere. In this sense we are following the "object-oriented" sense found in all of Z-set, and the damages are being treated as abstractions. They become concrete through the additional objects instanced in the different pointed to data files.

For this example, the following sections are included in the same input file, after the above "primary" section.

```
%section 2
test_simple
****post_processing
    **process fatigue_S
      *var sig
      *mode simple
    **process creep
      *var sig
****return
%section 3
test_with_a
****post_processing
    **process fatigue_S
      *var sig
      *mode with_a
    **process creep
      *var sig
***return
%section 4
test_simple_norm
****post_processing
    **process fatigue_S
      *var sig
      *mode simple
      *normalized_coeff
    **process creep
      *var sig
***return
%section 5
test_with_a_norm
****post_processing
    **process fatigue_S
      *mode with_a
      *var sig
      *normalized_coeff
    **process creep
      *var sig
***return
```

 $\dots$  continued

Z-set — Non-linear material & structure analysis suite

For the material files, there will be a succession of **\*\*process** sections with the types corresponding to the types of calculation in the input file. For example, in the file .../MAT/test\_simple referenced above we have:

```
%
% fatigue coefficients for the formula without a, and
% non normed coefficients
%
***post_processing_data
 **process fatigue_S
                           % coefficients relating to the
   М
              14782.065
                          % fatigue_S defined in %section 2
              2.5
   beta
   sigma_l
              40.5
   sigma_u
             170.
             0.0003
   b1
             0.0003
   b2
                         % coefficients relating to
**process creep
   A 452.
                         % creep defined in %section 2
   SO 0.
   r 10.
   k 30.
**process LCF
                         % coefficients controlling the LCF
                         % from section 1, mode NLC_ONERA
   а
       0.1
   k
       30.
   beta 2.5
   beta 2.5
***return
```

In the following LCF sections, there is a similar coefficient file with similar structure. These different files get ready because of the sequence of **\*\*material\_file** options swapping the effective material file name as the main input is read.

# \*\*\*\*post\_processing

#### **Description**:

This command marks the start of an input section for the post processor. More than one section of this type can be in the same input file *problem.inp*. By default, it is the first section which is executed, and commands are read until the next command which begins with 4 asterisks.

The execution command is (see options for Zrun in zman zmaster):

Zrun -pp prob

In oder to execute other post processing sections in the same file, use the -N option. For example:

Zrun -N 3 -pp prob

will execute the third section of **\*\*\*\*post\_processing** in the file **prob.inp**.

#### Syntax:

A post processing file will generally include **\*\*\*** star commands which adjust the operation of the post computations, and different sections of **\*\*\*local\_post\_processing** and **\*\*\*global\_post\_processing** sections defining the operations to be performed.

```
****post_processing
***precision num
***data_source source-type
***data_output output-type
***global_parameter
***suppress_p_on_post_files
***post_file_prefix prefix
***local_post_processing
...
***global_post_processing
...
****return
```

Commands which are essentially flags or adjustments to the **\*\*\*\*post\_processing** process follow:

- **\*\*\*global\_parameter** After this option one can include global parameter statements as if they were in a **zsetrc** file. Please see the Z-set/Release Notes manual for more information uder the Reference/Adjustable parameters section.
- **\*\*\*mesh** This option is used to re-map element types in the mesh to a different (post-processable) type. The command takes a series of *elset-name* and *elem-type* pairs to do the mapping.
- **\*\*\*post\_file\_prefix** used to set what the output files should be named as. This option makes the output look like another problem from the original input problem.

**\*\*\*precision** set the precision for formatted output of real values.

\*\*\*suppress\_p\_on\_post\_files this option indicates that the post data files should not have a suffix -p in the set of files used. This option can be used to stack multiple post computations together when combined with the \*\*\*post\_file\_prefix command. One can also use the \*\*\*data\_output command to a similar effect.

The following additional commands will often be used in multiple instances, and are all discussed in their own documentation sections. The local and post processing commands follow other **\*\*\*** options because they both have many sub-options for all the computation types.

- **\*\*\*data\_source** allows the user to ask the post-processor to read its input data from one of many different source formats (see documentation in following sections).
- **\*\*\*data\_output** more than one instance of this option allows outputting results to different formats.
- **\*\*\*global\_post\_processing** Start a sequence of parameters and commands defining the type of global post treatment to be applied.
- **\*\*\*local\_post\_processing** Start a sequence of parameters and commands defining the type of local post treatment to be applied.

## Example:

A complete post processing file follows to show the basic structure of the input options all together.

```
****post_processing
    ***precision 6
    ***local_post_processing
       **file integ
       **material_file creep.inp
       **elset ALL_ELEMENT
       **output_number 1-100
       **process mises
         *var sig
       **process eigen2
         *var sig
       **process trace
         *var sig
    ***global_post_processing
       **process average
         *list_var sigmises sigp1 sigp2 sigp3 sigii
    ***local_post_processing
       **output_number 1-100
       **process creep
         *var sig
       **process creep
         *var sig
         *express_life_as time
    ***global_post_processing
       **output_number 1
       **process average
         *list_var NC_S TC_S
```

\*\*\*return

# \*\*\*data\_source

# **Description**:

This procedure is used to set the source of data for post processing. The selection should come before any other **\*\*\***-level commands.

## Syntax:

\*\*\*data\_source type
[ \*\*open file ]
 ...
possible mesher commands

The following data source types are currently available.

CODE	DESCRIPTION
Z7	standard Zebulon results for versions up to 8.3
d3plot	LS-DYNA binary format
fil	ABAQUS .fil file format (unix or win32 binary)
fin	ABAQUS .fin ASCII file format
fin	ABAQUS .odb database format (use $\verb+odb$ switch when launching $\texttt{Zrun})$
ideas	I-DEAS format
neu	FEMAP neutral format
odb	Abaque ODB database (run with -odb switch)
rst	ANSYS .rst file (unix or win32 binary)
sim	Z-set simulator format
t16	MARC t16 results format
ascii	ASCII file input

When using the **\*\*\*data\_source** option one gets the chance to add additional meshing operations before the post processing begins. A particularly useful application of this is to add node or element sets which can then be used to specify the location for post processing.

# <u>Note</u>:

For ABAQUS FIL formats we recommend using the POSITION=AVERAGED AT NODES \*EL FILE option. There are some limitations still with the fil reader. Please submit a results file which is giving trouble. As of 8.3.3 the ODB format is much more robust and the preferred method. The ODB format is completely implemented for all node/element node/integration point data formats.

# Note:

When using the **\*\*\*data\_source** option the post processing will generally pass silently if a requested field variable is available in the results database, but not with the specified location

(e.g. element nodes **ctele**, integration points **integ**, etc). In those cases the data used will most likely be null (in some instances an automatic interpolation or extrapolation is used).

# Example:

A typical example follows. For import examples there are numerous .pst files in the Z-mat test directory and the Zansys directory. Also for Zebulon, Z-sim, and ASCII data sources look in the Post\_test directory.

```
****post_processing
  ***data_source Z7
    **open plast3_util
  ***precision 6
  ***global_post_processing
   **file node
   **output_number 1-999
   **nset ALL_NODE
   **process curve plast3_util.test
    *precision
                 3
    *node look
                 U2
    *nset press RU2
    *node look
                 eto22 sig22 epcum
****return
```

 $\dots$  continued

# Example:

Output from Z-sim or even ASCII files can be used for post processing (even with 1D data). An example of of loading the binary results from the simulator follows:

```
****post_processing
***data_source simulator
**dimension 2 % sets the pb dimension
**file_name sim_pb % data discribed in sim_pb.uti
***local_post_processing
**file integ % sim data is always integ
**elset ALL_ELEMENT % always need a location
**process format
*file sig.txt
*list_var sig11 sig22 sig33 sig12
****return
```

An example load from ascii file follows:

```
****post_processing
 ***data_source ascii
 **dimension 2
 **file_name result.dat
 ***local_post_processing
 **file integ
 **elset ALL_ELEMENT
 **process format
 *file sig.txt
 *list_var sig11 sig22 sig33 sig12
****return
```

In this case, the post-processor expects **result.dat** to be an ascii file in column order containing a first line which describes the contents of all columns. This file could contain, for instance:

```
# time sig11 sig22 sig33 sig12
0. 0. 0. 0. 0. 0.
10. 0. 112. 0. 0.
```

There are several validation examples in testPost\_test/INP/ for ascii and simulation data sourcing. Abaqus fil and odb formats are used int the .pst files under the Z-mat directories.

# \*\*\*data\_output

## **Description**:

This procedure is used to add a destination output of the post processing data of the following commands. The selection should normally come at the start of the **\*\*\***-level commands, except for a **\*\*\*data\_source** command which should always be first.

# Syntax:

\*\*\*data\_output type
\*problem\_name name
\*elset eset-name
type-specific-options

- \*problem\_name is used to specify the root-name of the problem which would have been run in the native code. Usually the output file will be this name appended with a dot suffix of the file type.
- **\*elset** specifies that the output problem is to be a sub-mesh made up of the given element set elements. This can be used as a general results file reduction capability with

The following data source types are currently available.

CODE	DESCRIPTION
odb	ABAQUS odb database format (use command switch -odb
	when executing Zrun)

Currently there are no additional *type-specific* options.

 $\dots$  continued

# Example:

A typical example follows. The data output capability has test cases in the test/Z-mat/zebu\_interface with a full assortment of stacked input and output runs are done.

```
****post_processing
  ***data_source Z7
    **open zpost_matsim.utp
  ***data_output odb
     *problem_name lifetime
     *elset MANIFOLD
  ***local_post_processing
     **output_number 1-9999
     **file node
     **nset manifold_nodes
     **material_file post_coefs.mat
   **process neu_sehitoglu_evi
    *total_strain_for_range
   **process copy
                         % for validation between input/output
    *list_var evcum
    *out_var evcum
    *last_only
****return
```

# \*\*\*local\_post\_processing

## **Description**:

This procedure is used to define the local post computations to apply at nodes or Gauss points. We recall that the local post computations are performed across all stored time points at each location. These operations are therefore primarily temporal, whereas a global post is used for spatial operations such as averaging over volumes. The global operations *can* however generate similar fields as local computations.

# Syntax:

The following syntax summary applies:

```
***local_post_processing
[ **duplicate | **no_duplicate ]
[ **elset eset ]
[ **file file-key ]
[ **ipset ipset ]
[ **material_file fname ]
[ **nset nset ]
[ **output_number out-num-list ]
[ **output_to_node|**no_output_to_node ]
[ **packet_size size ]
[ **process type ]
....
```

The sub options define geometrical groups of concern, time period, material files, and the post treatments to apply. Any number of **\*\*process** commands can be added in a **\*\*\*local\_post\_processing** section, and they will all normally have their own sub commands and parameters to enter. The following table summarizes the function of each of these commands, and the pages which follow give the detailed command input syntax.

- **\*\*duplicate** switches if we allow post processors to create variables with the same name or if the duplicate names are appended with \_n# in the **\*\*no\_duplicate** case (default).
- **\*\*elset** define the element set of concern.
- **\*\*file** define the *type* of file where the data will be read. (*problem.integ*, *problem.node...*).
- **\*\*ipset** define the Gauss point group of concern.
- **\*\*material\_file** indicates that a separate material file which is to be used for reading material coefficients if no **\*model\_coef** is entered in a **\*process** command.
- **\*\*nset** define the node set of concern.
- **\*\*output\_number** define the period of time which will be used. The default is the full time period of existing results.
- **\*\*at** is an alternative to **output\_number**: specific maps can be selected through their time.

- **\*\*output\_to\_node** indicates that the post output should be directed to a **node** file and not a **ctnod** file. The distinction is that if copying displacement variables the new files can have deformed geometry displayed in Zmaster.
- **\*\*packet\_size** can be used for large output results files to decrease physical memory requirements. In this case post computations are performed by packets of *size* elements/nodes. **Note:** This option is not compatible with all data-output formats.
- **\*\*process** add a new post computation and begin reading the input for it.

## Note:

In the first instance of **\*\*local\_post\_processing** or **\*\*global\_post\_processing** the full description of the file/localization (e.g. elset)/output range, etc. must be specified. Subsequent processing entries will however use the last entered data, so that easy switching of local and global post computations can take place.

### Example:

The following is another example input from the test Rainflow\_test/INP/fatigue.inp:

```
****post_processing
```

```
***precision 5
    ***local_post_processing
       **file integ
       **elset ALL_ELEMENT
       **output_number 1-40
       **material_file fatigue.mat
       **process range
         *alpha 0.001
         *var sig
       **process fatigue_S
         *mode simple
         *var sig
       **process multirange
         *var sig
         *reverse 5
       **process fatigue_rainflow
         *var sig
         *mode simple
         *reverse 5
    ***global_post_processing
       **output_number 1
       **process average
         *list_var Dsig NF_S ncyc D1sig NF1
****return
```

# \*\*output\_number

# **Description**:

This command is used to define the period of time which will be active for the post computation.

# Syntax:

|--|

The user indicates the number of "maps" in the form of a list of integers, or intervals of integers. The elements of the list are separated by commas or blank spaces. The intervals are specified with two integers separated by a dash (-).

In the absence of an **\*\*output\_number** entry, the post-processor will use all the solution maps in the file *problem.ut*.

# Example:

Output numbers have to be specifically asked for. The following input file snippets demonstrate the syntax allowable for this.

% to get outputs 1,2,3,4,5,20,35
\*\*output\_number 1-5,20,35
% to get outputs 1 and 20
\*\*output\_number 1 20
% to get outputs 1, 20 and 30
\*\*output\_number 1,20 30

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*at

\*\*at

## **Description**:

This command selects which maps which will be active for the post computation by their time, rather than by their number. Thus, it is an alternative to **\*\*output\_number**.

### Syntax:

## \*\*at t1 t2 ... tN

Note that specified time steps that do not exist in the results files are ignored. Explicitly specifying  $t1 t2 \dots tN$  in the **\*\*\*resolution** bloc ensures that such maps will exist.

\*\*nset

# **Description**:

This command defines the nodes to be treated by specifying a node group (nset). One can use the mesher (**\*\*\*\*mesher**) mode of operation to generate these sets.

# Syntax:

**\*\*nset** *nset* 

The groups of nodes *nset* must of course be defined in the geometry file *mesh.geof* designated after the instruction **\*\*meshfile file** in the file *problem.ut*. All the nodes of this group will be taken into account for the post calculation. To simply designate all the nodes in the structure, one may always use the pre-defined node set ALL\_NODE. The nset selection can be modified at any time with a new **\*\*nset** instruction.

# Example:

\*\*nset surface
% with the following definition in the .geof file:
\*\*nset surface
10 11 12 13 19 20 22 30 31
...

\*\*elset

# **Description**:

This option defines the elements to be treated with post computations by specifying an element group (elset). Elsets can be generated in a mesh using the batch mesher (chapter 4).

# Syntax:

\*\*elset *elset* 

The group of elements *elset* must of course be defined in the geometry file *mesh.geof* designated after the instruction **\*\*meshfile** in the file *problem.ut*. All the elements of this group will be taken into account for the post calculation; that is to say that the post computations will be evaluated at all the elements Gauss points. To simply designate all the elements in the structure, one may always use the pre-defined element set ALL\_ELEMENT. The elset selection can be modified at any time with a new **\*\*elset** instruction.

```
**elset wheel
% with the following definition in the .geof file:
** elset wheel
1 2 3 4 5 6 7 8
...
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*ipset

\*\*ipset

## **Description**:

This command is used to select the integration point set (ipset) to treat with post computations.

# Syntax:

\*\*ipset ipset

The group of Gauss points *ipset* must be defined in the geometry file *mesh.geof* designated after the instruction **\*\*meshfile** in the file *problem.ut*. Gauss points are specified in the *mesh.geof* file under the group section **\*\*ipset** by a series of element/integration point couples (*nel/nip* with *nel* the element id and *nip* the integration point number indexed from 1). The ipset selection can be modified at any time with a new **\*\*ipset** instruction.

# Example:

\*\*ipset failure
% with the following definition in the .geof file:
\*\* elset failure
1/1 1/2 2/1 2/2 3/1 3/2 4/1 4/2
...

\*\*file

## **Description**:

This command is used to specify on what types of file the post calculation will be calculated. **There is no selection by default**.

# Syntax:

**file	filetype
--------	----------

where *filetype* is the selection of files to read, among the possible files which were output by the finite element calculation. In the general case, the type of output file produced is the same as the input data file (i.e. if the input is an .integ file the output .integp file has the exact same format with different variables).

The following table is a summary of the different possibilities:

filetype	file read	file created
integ ctele ctmat node	problem.integ problem.ctele problem.ctmat problem.node	problem.integp problem.ctelep problem.ctmatp problem.ctnodp
	+ problem.ctmod	+ problem.ctnodp

Certain post-processors do not respect this generality. For example node\_interpolation reads its input data in the file *problem.*node but produces results in the file *problem.*integp.

The "active" file selection can be modified at any instant with a new **\*\*file** instruction.

# Example:

```
% to select the file extension .integ
% the results will be written in a file with a .integp
% extension
**file integ
```

Ş

# \*\*material\_file

## **Description**:

This command is used to specify the name of a material file containing coefficients when they are necessary. This file can be updated during the progression through the post calculations if it is necessary. The current material file is the one selected by the last instance of this command.

# Syntax:

\*\*material\_file file

# Example:

\*\*material\_file aluminum

# Material file syntax:

As this material file can be the same for the finite element, simulation and post processing calculations, it is necessary to localize the post-processor's specific data. This is done with an entry of the command **\*\*\*post\_processing\_data**<sup>1</sup>. In this data-entry section, different entries for the different post processors requiring coefficients should be entered. Each section will contain coefficients specific to the post processor given. Coefficients use the COEFFICIENT objects given in the Z-mat manual in Chapter 5.

Note that the COEFFICIENT can depend on the internal variables, auxiliary variables, flux and grad variables, or the external parameters (if the **save\_parameter** option was used for the later).

# Syntax:

```
***post_processing_data
 **process PROCESS [ option1 ... optionP ]
 name1 COEFFICIENT
 ....
 nameN COEFFICIENT
```

The optional input parameters *option1* ... *optionP* can be used to specify the particular use of the given post computation by giving an application option keyword, the tensor or variable on which it operates, etc.

The names *name1* ... *nameN* are the local names of the coefficients necessary to define for the corresponding calculation. These coefficient names are given for each process computation in the manual sections which follow.

<sup>&</sup>lt;sup>1</sup>Other behavior commands such as **\*\*\*behavior** terminate at the next command of level **\*\*\***. This effectively distinguishes the components of the material file, just as the **\*\*\*\*** level commands do in the input file

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*material\_file

```
***post_processing_data
**process fatigue_S sig
       2400.
   М
   beta 5.
   sigma_l 120.
   sigma_u 450.
***post_processing_data
 **process creep xtv
   А
       temperature
   1400. 1050.
   2350. 900.
   5000. 700.
        5.
   r
```

Ş

# \*\*process

## **Description**:

This keyword introduces a post-computation to be added to the set of available results. most post computations take one or more *subject* variables which will provide the basis of the calculations (e.g. finding principal values of a tensor takes a tensor name as the subject, which could be **sig** to find the principal stresses).

Post-computations which take subject variables introduced by the key word **\*list\_var** are applied uniquely to scalar variables. All the character strings following the key word are interpreted as such.

Certain post-computations which use the keyword **\*var** to introduce the subject variable are applied to either scalar or tensorial variables. In this case, there will also be a command option named **\*type** which is provided to specify the type of variable. Nevertheless, in order to simplify the input, the following variable name convention is given:

- All character strings which end in 11, 22, 33, 12, 23 and 13 will be interpreted as the name of a scalar variable.
- Otherwise the character string is interpreted as a tensorial variable name.

The following local post-processor types are available in the standard distribution.

HCF base_fatigue delay ductile_failure	LCF copy derive eigen2	adiabatic_temperature creep deviator evcum_sum
external	fatigue_E	fatigue_EE
fatigue_S	fatigue_rainflow	fmax
fmin	format	function
harmonic	hyper_visco	inc_creep_damage
inelastic	initiation	integrate
load	local_frame_axes	mat_sim
max	min	mises
multirange	neu_sehitoglu	norm
oxidation	rainflow	range
trace	transform_frame	tresca
triax	z7p	

Ş

# \*\*process copy

## **Description**:

The copy post processor is used to re-name a variable as a copy into the results database files. There are several occasions where this operation would be useful:

• For Z-mat runs where all the state variables are named SDV# (ABAQUS) or SVR# (AN-SYS), etc. The copy operation can be used importing the original input file using a **\*\*data\_source** (see page 4.14) and re-naming them in a more meaningful way. Note also that if the output naming respects the 11 22 33 12 23 31 ordering those sub-variables will be considered as tensors in further Z-post operations.

Remember that the SDV# name mapping can be determined using the Zpreload command or by looking at the calculation message files.

- To copy a database into a different format, but making certain to respect important fundamental names (such as **U1 sig11** etc. Using the copy functionality can therefore be used to reduce the size of a results set, or to translate between different supported formats.
- To fake the name of a tensor if needed. For example one could open an Abaqus ODB file, and copy the plastic strain tensor to be the total strain tensor and submit that to another code requiring total strain only.

There are no doubt other uses for such a capability, but these are the primary intent of the command.

## Syntax:

\*\*process copy
 \*\*list\_var var1 var2 ... varN
[ \*\*out\_var o-var1 o-var2 ... o-varN ]

The variables will be copied in the order given. Memory can be reduced for large models by breaking up man copies into several **\*\*process copy** operations.

## Note:

If the **\*\*out\_var** option is not given, the output names will be the same as the input names, so that should only be used when the post results are desired to "stand alone."

 $\dots$  continued

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process copy

### Example:

The following copies an Abaqus odb file to a new one while re-naming the state variables (in this case a **\*\*\*data\_storage** command was used to reduce the SDV allocations to just 2 variables).

```
****post_processing
 ***global_parameter
   ODB.MaxSteps 100
 ***data_source odb
   **open my_abaqus_run.odb
   **write_geof
   **nset interesting_nodes *elset MAIN_ELSET *function 1.0;
 ***data_output odb
   *problem_name zpost_copy_odb
   *elset MAIN_ELSET
 ***local_post_processing
     **output_number 1-999
     **file node
     **nset interesting_nodes
    % DO NOT USE PACKET SIZE HERE
     **process copy
     *list_var SDV1
                         SDV2
                                 NT11
     *out_var evcum
                         Dsum
                                 temperature
     **process copy
     *list_var sig11 sig22 sig33 sig12 sig23 sig31
     **process copy
     *list_var LE11 LE22 LE33 LE12 LE23 LE31
     *out_var eto11 eto22 eto33 eto12 eto23 eto31
```

\*\*\*\*return

## \*\*process creep

#### **Description**:

This post calculation calculates creep damage at an instant t in relation to a variable T, such as the integration of time between  $t_1$  and t:

$$I(t) = (k+1) \int_{t_1}^t \left(\frac{S - S_0}{A}\right)^r$$

with

$$S = (1 - \alpha - \beta)\mathbf{Mises}(T) + \alpha \mathbf{Eig1}(T) + \beta \mathbf{Trace}(T)$$

where Mises(T) indicates the second invariant of the deviatoric part of a tensor T, Eig1(T) the maximum principal value, and Trace(T) the trace (first invariant.

In accordance with the users choice, the result is given in terms of time to rupture or number of cycles to rupture. Two cases are envisioned:

- I(t) attains 1 for a value of time between  $t_1$  and  $t_2$ : the corresponding value is the time to rupture, and the number of cycles to rupture is set to one.
- $I(t_2)$  is less than 1: the time to rupture is obtained by superimposing a cyclic time and linear cumulation law. The number of cycles to failure is simply the inverse of  $I(t_2)$  and the time to rupture is obtained by multiplying this number by the length of loading considered.

# Syntax:

```
**process creep
 *var name
[*type scalar | tensor ]
[*express_life_as cycle | time ]
[*delay ]
[*scale lin | log ]
[*precision prec ]
```

name is the name of the variable to treat. The option **\*express\_life\_as** is used to specify the output mode to be number of cycles or time to failure. With the option **\*delay** a scaled stress measure is used:  $dS' = (S - S')/\tau$ .

The user must furnish the coefficients S0, A, and r. Coefficients k, alpha et beta are optional with zero value by default. If it is used, the coefficient k must be a constant. One more supplemental coefficient tau is expected with the option \*delay.

In the file *problem*.utp, we call the variable for time to rupture TC\_S, and the number of cycles to rupture NC\_S. If the output was put on a logarithmic scale, the names will be LTC\_S and LNC\_S. We generally prefer to work with log values when generating contour plots.

Creep damage is integrated using a second-order runge-kutta method. The precision required for integration can be specified by the optional **\*precision** keyword. Default value is 1.e-4.

```
\% Use stress tensor to compute time to failure and give
% a logarithmic output
**process creep
  *var sig
  *scale log
  *express_life_as time
\% The following syntax is used in the material file :
**process creep
   A 1130.
        7.
   r
        0.
   S0
   alpha
           0.3
   beta
           0.
\% ... but the coefficients may be non constant
**process creep
   A temperature
   1130.
           1050.
   2000.
            700.
   . . . .
```

# \*\*process cycle

#### **Description**:

The cycle post processor may be used to apply the same post-processing operations on all cycles of a cyclic calculation. Typical application is to monitor the evolution of critical quantities (stress amplitude, mean stress, number of cycles to failure, etc...) from one cycle to the other.

This post-processor takes an arbitrary number of sub-processors as arguments. The output maps available in the results file that correspond to each cycle are automatically selected, and given as input to the embedded sub-processors. The definition of the time period of the cyclic calculation is used as a basis to sort out which maps do belong to a particular cycle.

A single map of post-processing results will be generated for each cycle detected in the sequence of output maps available in the input results files. In this context two types of embedded sub-processors may be distinguished:

- (i) post-processors that generate one map of results for each input map (eg. \*process mises, \*process function, etc ...)
- (ii) post-processors that generate one single map for the whole range of input cards (eg. \*process max, \*process range, etc ... and all the damage post-processor in general)

Type (ii) needs no further specification, since the results given for each cycle correspond to a standard calculation on the associated cycle input maps (that may be selected by a repeated use of the **\*\*output\_number** command). On the opposite, for type (i) post-processors 2 options are available to specify if eiher the *max* of *average* value should be retained for each cycle.

#### Syntax:

```
**process cycle
 *period period
[ *start tstart ]
[ *end tend ]
 *process proc1 [ typ1 ] [sec1 ]
 *process proc2 [ typ2 ] [sec2 ]
...
*process procn [ typn ] [secn ]
```

where *period* if the time period value of the cycle.

The **\*start** command may be added if a preload sequence is stored in the results file, that needs to be skipped before scanning for cyclic results.

Similarly, the optional **\*end** command will stop the scanning of cyclic results for maps whose time values are greater than the *tend* specified.

Embedded sub-processes are specified by an arbitrary number of \*process commands. Argument *proci* is the name of the sub-process, and *seci* is the number of the input file section where the actual \*\*process *proci* definition will be given. Default value for this section number is 2. For type (ii) post-processors (see description above) argument *typ1* can be either max or average (default value is max). Variables written in the post-processing files are the cycle number (variable cyc) and output variables associated to the various sub-processes. For the latter, the "\_cyc" character string is appended to the conventional variable name.

### <u>Note</u>:

Specification of a material file is mandatory when using **\*process cycle**, even if embedded sub-processors don't need any material coefficients. In the latter case an empty material is required:

\*\*\*post\_processing\_data
\*\*\*return

```
The following commands can be used to compute
the max value of the von mises stress invariant
the mean value of the average stress, and the
multiaxial stress amplitude for each
cycle stored in the FE results files.
The process format allows to store the result
in an ascii file: note the "_cyc" character
string added to the name of the sub-process
output variables.
% first section
****post_processing
  . . .
  % note the need of an empty material file
  **material_file fake
  **process cycle
   *period 10.
   *process mises max 2
   *process trace average 2
   *process range 2
  **process format
   *list_var cyc sigmises_cyc sigii_cyc Dsig_cyc
   *file cycle.post
****return
% second section: sub processes definition
****post_processing
**process mises
*var sig
**process trace
*var sig
**process range
*var sig
***return
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process cycle

The \*process fatigue\_S should in theory be applied on the stabilized cycle. The next example illustrates how to use \*process cycle to evaluate the influence of an incomplete material stabilization on the number of cycles to failure predicted by the fatigue\_S model.

```
% first section
****post_processing
...
**material_file fatigue_S_coefs
**process cycle
*period 50.
*process fatigue_S
**process format
*list_var cyc NF_S_cyc
*file nf_evolution.post
*****return
```

```
% second section: sub processes definition
****post_processing
**process fatigue_S
*var sig
****return
```

# \*\*process cycle\_projection

# **Description**:

This post is used when we have a multiaxial loading which is close to proportional and it is desired to see the primary stress-strain hysteresis loops. The user will pick a specific point in a cyclic loading (or series of points at similar conditions) and the requested tensors will be projected onto the eigen directions determined at that point.

The user can specify the point from which we take the eigen values/vectors from:

- A map number can be given. This will be indexed base 1 from the start of the selected cards in the active post section (that is if we select maps 20-200, a selection of 20 will be number 40 in the global maps).
- A time can be given. If the time is between 2 maps the values will be interpolated.
- The post calculation uses a POST\_CYCLE\_TOOL object to determine the cycle positions, from which we can use a cycle number combined with start/mid/end tokens. The default is this method, with cycle 1/mid selected.

Using the eigen vectors of the ordered principal calculation of the given **\*primary\_variable** selection we project the selected tensors into that frame and output them with either the given list of output names, or as *tname\_cpj*. If the primary tensor is desired as output then it should also be given in the **\*var** listing.

## Syntax:

```
**process cycle_projection tool-type
 *primary_variable name
 *var name1 name2 ... nameN
[ *output name1 name2 ... nameN ]
[ *time time-val ]
[ *time time-val ]
[ *map map-index ]
[ *cycle num (start|mid|end) ]
[ *remap_each_cycle ]
```

# Note:

The POST\_CYCLE\_TOOL expects that temperature be defined as a field variable. If it does not, one can use the make\_field local post computation in order to do so.

 $\dots$  continued

# Example:

A common usage is to use the hot\_spot post computation to identify critical regions, and then study the mechanical cycles at that location.

In this example we have a nset named min\_life.nset stored via a hot\_spot (see page 4.113). That nset is split in the beginning mesher command \*\*split\_nset run when loading the data source so we can specify node locations symbolically. The worst case node is then named min\_life\_1.

Note that the tmf keyword after the **\*\*process cycle\_projection** command identifies the type of cycle identifying tool to use. Currently the reader should refer to the **sehitoglu** life prediction methods (page 4.77) for a discussion. Future versions of the documentation will have a more expansive discussion on that.

```
****post_processing
  ***data_source Z7
    **open zpost_matsim.utp
    **nset min_life
     *nodes_in_file min_life.nset
    **split_nset min_life
  ***local_post_processing
     **output_number 1-9999
     **file node
     **nset min_life_1
     **process cycle_projection tmf
      *primary_variable sig
      *var
             sig eps_me evi
      *out
             sig_x eps_x evi_x
      *cycle 1 mid
      *remap_each_cycle mid
      *total_strain_for_range
  ***global_post_processing
   **process curve curves_1.test
    *precision
                 3
    *node min_life_1 eps_x11 eps_x22 eps_x33 eps_x12 eps_x23 eps_x31
                      sig_x11 sig_x22 sig_x33 sig_x12 sig_x23 sig_x31
                      evcum temperature
***return
```

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process derive

# \*\*process derive

## **Description**:

This post calculation computes the derivative of some fields with respect to a variable (time or another variable). It is computed using a first order formula:

$$\dot{f}(t_i) = \frac{f(t_i) - f(t_{i-1})}{t_i - t_{i-1}}$$
 if the field is derived wrt. time  
$$\frac{\partial f}{\partial x}(t_i) = \frac{f(t_i) - f(t_{i-1})}{x(t_i) - x(t_{i-1})}$$
 if the field is derived wrt. another variable

# Syntax:

\*\*process derive

\*list\_var names base

names is the list of variables to derive with respect to base.

# Example:

The following example computes the velocity in each direction. It will generate the fields  $D_time_U1$ ,  $D_time_U2$  and  $D_time_U3$ .

```
****post_processing
***local_post_processing
**file node
**nset ALL_NODE
**process derive
*list_var U1 U2 U3 time
****return
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process deviator

# \*\*process deviator

### **Description**:

This post calculation calculates the deviatoric stress tensor. The prefix  $dev_{-}$  is prepended to the tensor name.

### Syntax:

\*\*process deviator

\*var name

```
****post_processing
***local_post_processing
**file integ
**elset ALL_ELEMENT
**process deviator
*var sig
****return
```

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process ductile\_failure

## \*\*process ductile\_failure

#### **Description**:

This post treatment is applied to predict ductile rupture under monotonic loading. The defines a damage variable D at the time t which is the integration of time between  $t_0$  and t:

$$D(t) = \exp\left[\int_{t_0}^t a \exp\left(b\frac{\operatorname{Trace}(S)/3.}{\operatorname{Mises}(S)}\right) dp\right]$$

where S is a tensor and p is a scalar variable (normally the equivalent plastic strain).

One output is generated for each solution "map." As the evaluation of this criterion is based on an integration, a sufficient number of result outputs are required for a precise answer. The user should therefore ask for a complete set of intermediate outputs (see output\_number on page 4.21).

#### Syntax:

```
**process ductile_failure
 *stress name1
 *strain name2
[*rice_tracey]
```

name1 is the name of the tensor, and name2 that for the scalar. With the option **\*rice\_tracey** the criterion is evaluated with the stress tensor **sig** and the cumulated plastic deformation<sup>2</sup>.

The default values for the coefficients a and b are those used by the Rice-Tracey criterion: a=0.283 and b=1.5.

The variable names generated (in the *problem.utp* file, and appearing in the Zmaster results) is DAMAGE in the case of a general ductile criterion, and  $R_RO$  if the particular case of Rice-Tracey was chosen.

```
% Applying the Rice-Tracey criterion needs only:
**process ductile_failure
    *rice_tracey
% The following instructions allow the user to also compute
% Rice-Tracey criterion. sig and epcum are respectively
% the names of the stress tensor and of the
```

 $<sup>^{2}</sup>$ this assumes the name epcum for the cumulated plastic strain. Note that gen\_evp behaviors name this variable according to the users input after a potential keyword

```
% cumulated plastic strain in the prob.utp file.
**process ductile_failure
  *stress sig
  *strain epcum
\% The following syntax must be used in the material file :
**process ductile_failure
       0.283
  а
  b
       1.5
% This is another criterion:
**process ductile_failure
  *stress x1v
  *strain evcum
\% ... with other values in the material file:
**process ductile_failure
       0.33333
  а
  b
       2.
```

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process eigen2

# \*\*process eigen2

# **Description**:

This process is used to compute the eigenvalues of a symmetric second order tensor. No coefficients are needed. The process provides three values for each point. The eigenvalues are given in decreasing order, starting by the larger one. Their name in the *problem.utp* file is built from the name of the tensor by adding respectively p1, p2 and p3.

# Syntax:

\*\*process eigen2

\*var name

```
% this will provide sigp1, sigp2, sigp3
% with sigp1 >= sigp2 >= sigp3
**process eigen2
*var sig
```

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process fatigue\_E

## \*\*process fatigue\_E

### **Description**:

This post-processor is used to predict fatigue lifetime using a strain measure. This strain measure can be a plastic strain, or another one such as the total strain. The critical variable is the amplitude of the strain (plastic) for uni-axial loading with a generalization pertinent for multi-dimensional loading (see the post computation range). If the loading is well known, the user can give simply the output "maps" necessary to calculate this amplitude (mode n1 n2 in \*\*output\_number). In the other case, a period of loading must be input (mode n1-n2 in \*\*output\_number).

The strain amplitude is denoted  $\mathbf{DEQ}$ . The post calculation will generate a single result record for the entire loading history. The number of cycles to failure is defined as follows:

$$N_f = \left(\frac{\mathbf{DEQ}}{A}\right)^{-\alpha}$$

Syntax:

\*\*process fatigue\_E
 \*var name
 [\*type scalar | tensor ]
 [\*scale lin | log]
 [\*range section]

name is the name of the variable to use as input to the computation.

For multi-dimensional loading the amplitude calculation will be made using a **range** type post-computation. In order to be able to input options to the range processor, the user can give a section number for that user input after the **\*range** keyword.

The computation requires input of the coefficients A and alpha in the material file.

Output is given as number of cycles to failure in the variable NF\_E. If the user specifies a logarithmic scale using the \*scale option, the output variable will be LNF\_E.

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process fatigue\_EE

# \*\*process fatigue\_EE

### **Description**:

This post-computation is used to determine component life by a criterion dependent on a strain measure (elastic and plastic). The critical variable is the amplitude of the strain for uniaxial loading, with a generalization for multi-dimensional loading (see the post processor range). If the loading is well known, the user can give simply the output "maps" necessary to calculate this amplitude (mode n1 n2 in \*\*output\_number). In the other case, a period of loading must be input (mode n1-n2 in \*\*output\_number).

The strain amplitude is denoted  $\mathbf{DEQ}$ . The post computation will generate a single output record for the total loading history input. This is the number of cycles to failure, given by:

 $\mathbf{DEQ} = \mathbf{A} \, \mathbf{N_f}^{-1/\alpha} + \mathbf{B} \, \mathbf{N_f}^{-1/\beta}$ 

Syntax:

```
**process fatigue_EE
 *var name
[*type scalar | tensor ]
[scale lin | log]
[range section]
```

name is the name of the subject variable of the computation.

For multi-dimensional loading the amplitude calculation will be made using a **range** type post-computation. In order to be able to input options to the range processor, the user can give a section number for that user input after the **\*range** keyword.

The computation requires input of the coefficients  $\tt A$  and  $\tt B, \tt alpha$  and  $\tt beta$  in the material file.

Output is given as number of cycles to failure in the variable NF\_EE. If the user specifies a logarithmic scale using the \*scale option, the output variable will be LNF\_EE.

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process fatigue\_EE

alpha	З.
В	1.7
beta	1.5

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process fatigue\_rainfl

# \*\*process fatigue\_rainflow

## Description:

This command is used in replacement of **\*\*process fatigue\_S** for rainflow counting evaluations of fatigue life. The model applied is the same as the one described for **\*\*process fatigue\_S**, but a number of cycles to failure  $N_f$  is calculated for each subcycle detected by the **\*\*process multirange** post-processor.

# Syntax:

```
**process fatigue_rainflow
 *var name
[*reverse ] number ]
[*type scalar | tensor ]
[*mode with_a | simple ]
[*normalized_coeff ]
[*use_mises ]
[*scale lin | log ]
[*range section ]
```

The only difference with **\*\*process fatigue\_S** is the **\*reverse** keyword, that allows to specify the maximum number of sub-cycles retained in the calculation, within the ones detected by the rainflow counting algorithm (see the **\*\*process multirange** command). Default value is *number*=3. Note that **\*\*process multirange** gives out cycles with maximum amplitude first, such that most damaging cycles are always taken into account in the calculation. Damage cumulation can be performed by the **\*\*process rainflow** command.

Material parameters are the same as the ones described for the **\*\*process fatigue\_S**.

By default the equivalent max stress SMAX is the maximal eigen stress  $S_I$ . Using the option \*use\_mises makes SMAX equal the von Mises stress. Caution: in the \*\*process fatigue\_S the default equivalent max stress SMAX is the Mises stress.

Output is as follows:

- ncyc the number of cycles detected. If the algorithm detects more cycles than the number requested by the **\*reverse** number keyword, the code returns ncyc = number.
- the number of cycles to failure NF*i*, associated with each sub-cycle *i*. If a logarithmic scale is requested by means of the option \*scale the output is renamed LNF*i*.

# \*\*process fatigue\_S

### **Description**:

Using the stress as critical variable, the formula implemented by this option predicts a fatigue life (Wöhler curve). Even though the criterion can be applied to any variable contained in the problem, the description of the equations is made in terms of stress.

The model is capable of taking into account a mean stress and a multi-axial loading history. The critical variables are the stress amplitude and the maximum stress (with generalization pertinent for multi-axial values), calculated over the time period specified by the user.

The model is written in terms of reduced stress (The components of the stress divided by the last stress under monotonic loading  $\sigma_u$ ). The stress amplitude is noted **SEQ**, and the equivalent maximal stress is **SMAX**, and the mean trace is **TRMEAN**. The post computation returns a rupture in one cycle if **SMAX** attains the ultimate stress  $\sigma_u$ , and an infinite number of cycles if **SEQ** is less than the fatigue limit  $\sigma_l$ . Otherwise the number of cycles to failure  $N_f$  is defined as follows:

$$N_f = \frac{1}{a(\beta+1)} \left\langle \frac{\sigma_u - \mathbf{SMAX}}{\mathbf{SEQ} - \sigma_1'} \right\rangle \left( \frac{\mathbf{SEQ}}{\overline{M}} \right)^{-\beta}$$

with  $\sigma'_l = \sigma_l (1. - b_1 * \mathbf{TRMEAN})$ ;  $\overline{\mathbf{M}} = \mathbf{M}(1. - \mathbf{b}_2 * \mathbf{TRMEAN})$ 

## Syntax:

```
**process fatigue_S
 *var name
[*type scalar | tensor ]
[*mode with_a | simple ]
[*normalized_coeff ]
[*use_eigen2 ]
[*scale lin | log ]
[*range section ]
```

name is the name of the subject variable if the computation (one would expect sig here). Using the extra keyword simple after the option \*mode, the post-computation uses a simpler formulation where the ratio  $1/a(\beta + 1)$  is included in the coefficient M.

The coefficients sigma\_1, M, b1 et b2 can be optionally given in normalized form. If they are normalized (using the option \*normalized\_coeff) the expected names in the material files are changed to N\_sigma\_1, N\_M, N\_b1 and N\_b2.

The other coefficients necessary to input are **beta** and **a** if the option with\_a is given.

By default the equivalent max stress SMAX is the von Mises stress. Using the option  $*use\_eigen2$  makes SMAX equal the maximal eigen stress  $S_I$ . Caution: in the  $**process fatigue\_rainflow$  the default equivalent max stress SMAX is the maximal eigen stress  $S_I$ .

Number of cycles to failure is given in the output, with the variable named NF\_S. The user can ask that a logarithmic scale be used with the option \*scale. If so the output will be named LNF\_S.

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process format

# \*\*process format

### **Description**:

This post processor is used to produce an "export file" containing the specified variables for all the maps being run. The output file is a formatted ASCII file. The variables should exist in the file specified by the **\*\*file** command (see page 4.26). For instance, the command **\*list\_var U1** should not be given with **\*\*file** integ. Note: this post processor also exists for **\*\*\*global\_post\_processing**, with similar options (see page 4.112).

### Syntax:

\*\*process format
 \*file file
 \*list\_var name1 ... nameN
[ \*precision digits ]
[ \*optimizer ]
[ \*blank\_line ]

where file denotes the name of the output file, and name1, ..., nameN is a list of scalar variable names to output. The values are stored point by point for each map. The option \*optimizer inhibits the writing of the character string # time name1 ... nameN. The option \*precision determines the number of *digits* with which the results will be written (default is 10), and the option \*blank\_line writes an empty line between the output of each location. Some plotting programs, for instance gnuplot, may need this.

## Example:

The following example

```
**output_number 1-3
**process format
*file myfile
*list_var U1 U2
```

will produce a file called myfile like this:

#	time	U1	U2
	1	0	0
	2	0	0
	3	0	0
	1	0	0.01
	2	0	-0.01
	3	0	0
	1	0.01	0.01
	2	-0.01	-0.01

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process format

3 0 0 ...

Adding the **\*blank\_line** and **\*optimizer** options will give

1	0	0
2	0	0
3	0	0
1	0	0.01
2	0	-0.01
3	0	0
1	0.01	0.01
2	-0.01	-0.01
3	0	0

• • •

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process function

## \*\*process function

#### **Description**:

This post processor allows the user to write an interpreted function of scalar variables.

## Syntax:

\*\*process function
 \*output name
 \*expression expression

name is the name of the resulting variable to be generated by the function evaluation. expression is the function expression to be input (see the function reference on page 5.2). A semicolon must always mark the end of the function.

```
% this example computes FMAX_sig ...
**process fmax
*list_var sig
% and uses it in the calculation of the variable Pr.
**process function
*output Pr
*expression 1.-exp(-(FMAX_sig)/412.);
```

## \*\*process HCF

#### **Description**:

This post-computation gives an evaluation of the equivalent stress to compare with the fatigue limit, in order to define the high cycle fatigue (HCF) resistance.

Four different criteria are implemented for HCF. The variables which provide the basis of the criterion are the hydrostatic pressure (for each of them), and a stress amplitude in terms of the von Mises invariant (for three of them), or in terms of a shear (in one case). In the following, the stress amplitude is designated Dsig (and defined in relation to stresses tensor),  $p_{max}$  the maximum hydrostatic pressure, and  $p_{mean}$  the mean value of hydrostatic pressure ( $p_{mean} = 0.5(p_{max} + p_{min})$ .

The following criteria are implemented for the HCF model:

• The criterion from Sines (mode SI) uses a coefficient b to calculate the equivalent stress:

 $\sigma_e q = Dsig + bp_{mean}$ 

• The criterion due to Crossland (mode CR) uses a coefficient b to calculate its equivalent stress:

 $\sigma_e q = (1-b)Dsig + bp_{max}$ 

- The criterion from Dang Van uses a coefficient *b*. Its characteristic, if we compare the two first two models, resides in the combination of two variables at the same time. Two versions are implemented:
  - The classical model of Dang Van (mode DV) searches for the maximum value in all the physical space directions  $(\vec{n})$ , at all instants  $t_i$  of the equivalent stress. This is constructed from the current shear stress amplitude  $\tau$  and from the hydrostatic stress:

$$\sigma_e q = \max(\vec{n}) \min(t_i) (\tau(t_i) + bp(t_i))$$

- An alternative formulation of the Dang Van criterion (mode DV2) is also implemented. Following the same philosophy as previously, this modification provides a simpler evaluation because it uses a von Mises stress measure. Knowing the stress amplitude one can calculate the value  $\sigma_0$  corresponding to the loading path "center." The critical variable  $DJ_2(t_i)$  then corresponds to the the von Mises invariant of the difference in current stress and  $\sigma_0$ :

$$\sigma_e q = \max(t_i)(DJ_2(t_i) + bp(t_i))$$

#### Syntax:

\*\*HCF

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#### \*mode SI | CR | DV | DV2

The user must choose the *mode* of operation (SI, CR, DV, or DV2). The coefficient b must be defined in the material file. The post-computation produces an output for each input point at each time, which is the equivalent stress. The output variable name is HCF\_mode.

#### Example:

\*\*process HCF
\*mode DV2
% With the following definition in the material file
\*\*process HCF DV2
b 0.3

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process initiation

## \*\*process initiation

### **Description**:

This post processor is used to estimate the time to crack initiation. For a cycle the initiation damage is calculated by the following expression:

$$I_a = \frac{1}{c} \int_{cycle} \left\langle \frac{\langle SII - Sla(1 - Dox)}{(1 - Dox) - S_{max}^{eq}} \right\rangle^b dN$$

with

$$Sla = Sla0(1 - h\overline{\mathbf{Trace}(S)})$$

and  $Dox = \sqrt{N/Nox}$  if we're taking into account the effect of oxidation, Dox = 0 otherwise.

A single output map is generated for the total history of loading. The number of cycles to initiation  $N_a$  which corresponds to the moment where the damage attains a value of 1. The variable name generated is Na in the *problem.utp* file. The number of oxidation cycles ahead of the start of initiation (stress lower than the fatigue limit) is also saved with the name Na-ox.

## Syntax:

```
**process initiation
 *var name
 *type scalar | tensor ]
 [*normalized_coeff]
 [*oxidation [section1]]
 [*range section2]
```

name is the variable name. It is systematically normalized by the coefficient sigma\_u given in the material file.

The option **\*oxidation** is optional. If it is present, it can be followed by the number of section containing the user input for oxidation. Otherwise the oxidation will be initialized with its default values.

For multi-dimensional loading the amplitude calculation will be made using a **range** type post-computation. In order to be able to input options to the range processor, the user can give a section number for that user input after the **\*range** keyword.

The coefficients Sla0 and h are given normed or not. If they are normalized (the option \*normalized\_coeff was given), the expected names in the material file will be N\_Sla0 N\_h. The other coefficients which must be input are b and c.

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process initiation

% a complete example \*\*process initiation \*var sig \*oxidation \*norm \*normalized\_coeff %with the following syntax in the material file : \*\*process initiation sigma\_u 130. N\_h 5.2 N\_Sla0 0.023 b 2. c 3500.

## \*\*process LCF

#### **Description**:

This process is used to apply a cumulation model for the life prediction under creep-fatigue interaction. The reference number of cycles to failure in pure fatigue and in pure creep must have been previously computed. The number of cycles to failure under creep-fatigue loading is defined as  $N_r$ , from the number of cycles to failure under creep  $N_c$  and the number of cycles to failure  $N_f$ . Several cumulation rules can applied, according to users's choice:

• linear cumulation, LC

$$\frac{1}{N_r} = \frac{1}{N_c} + \frac{1}{N_f}$$

• bilinear cumulation, BLC

A "knee-point" is defined in the cumulative diagram, from the numerical value of the material coefficients Kc and Kf (both values must be between O and 1).

if 
$$\frac{N_c}{N_f} \ge \frac{K_c}{K_f}$$
:  $\frac{1}{N_r} = \frac{1}{N_f} + \frac{1 - K_f}{K_c} \frac{1}{N_f}$  (1)

if 
$$\frac{N_c}{N_f} \le \frac{K_c}{K_f}$$
:  $\frac{1}{N_r} = \frac{1}{N_c} + \frac{1 - K_c}{K_f} \frac{1}{N_c}$  (2)

The resulting cumulation rule must :

- be equivalent to the linear cumulation rule if  $K_c + K_f = 1$
- predict lower life if  $K_c + K_f < 1$
- predict longer life if  $K_c + K_f > 1$

• nonlinear cumulation, NLC

The cumulation rule uses the quantities  $C = \frac{1}{N_c}$  and  $F = \frac{1}{N_f}$  to compute damage evolution from  $D_i$  to  $D_f$  in each cycle, according to:

$$C = (1 - D_i)^{k+1} - (1 - D)^{k+1}$$
(3)

$$F = \left[1 - (1 - D_f)^{\beta + 1}\right]^{1 - \alpha} - \left[1 - (1 - D)^{\beta + 1}\right]^{1 - \alpha}$$
(4)

The resulting cumulation rule predicts lower lives than linear cumulation does.

• another nonlinear cumulation, NLC\_ONERA

This is the same cumulation rule as previously given. The difference rests in the the coefficient  $\alpha$  which is constant in the mode NLC and calculated as follows in this mode:

$$\alpha = 1 - a * \left(\frac{Dsig/2 - \sigma_l}{\sigma_u - \sigma_{max}}\right)$$
(5)

with

$$\sigma_l = \sigma_{l0}(1 - b1 * \overline{tr(\sigma)}) \tag{6}$$

#### Syntax:

The syntax for this criterion is as follows.

```
**process LCF
 *mode LC | BLC | NLC | NLC_ONERA
 *fatigue name1 section1
 *creep name2 section2
[*initiation name3 section3]
[*scale lin | log ]
```

The user chooses his method of cumulation among the key words LC, BLC, NLC and NLC\_ONERA after \*mode. name1 is the name of the fatigue processor to apply, and name2 the name of the creep processor. Those which are currently available for fatigue are fatigue\_S, fatigue\_E and fatigue\_EE. The current creep processor is uniquely creep. Plug-ins could however be created for new models of either of these. For each the user indicates after the processor type a number of post processing section containing the input for that processor.

The user can add an initiation phase, of which the name and the post processing section are specified after the option \*initiation. Only the post-computation initiation is currently available. The number of cycles  $N_a$  is then taken into account in the cumulation rule.

The model LC does not require any material coefficients. Two coefficients Kc and Kf are necessary for the model BLC, three coefficients k, alpha and beta for the model NLC, and five coefficients k, beta, a, sigma\_10 and b1, are required for the model NLC\_ONERA. Some of these coefficients appear in the criterions for creep or fatigue. In this case the reading will attempt to find the coefficients in the respective creep or fatigue sections. For example, the coefficient k of the mode NLC will be read in the section **\*\*process creep**, if it is present, otherwise in the section **\*\*process LCF** of the material file.

The number of cycles to failure  $N_r$  is called NR\_mode in the output variables. The user can also ask for a logarithmic scale which will re-name the output to LNR\_mode. The number of cycles to failure  $N_f$ ,  $N_c$  and  $N_a$  are also stored in the output (see these processors to find the specifics of variable naming).

#### Example:

A complete example with non linear cumulation rule follows:

Z-set — Non-linear material & structure analysis suite

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process LCF

```
**process LCF
  *mode NLC
  *fatigue fatigue_S 2
  *creep creep 2
  *scale log
% with in an other ****post_processing section
**process creep
  *var sig
**process fatigue_S
  *var sig
  *range 2
**range
  *var sig
  *method 2
  *alpha 0.2
\% With the following definition in the material file
\% beta and k are readed in their specific section
**process LCF
  alpha 0.9
**process fatigue_S
  М
          980.
          2.5
  beta
  sigma_1 70.
  sigma_u 180.
  b1
          0.003
          0.003
  b2
**process creep
  SO 0.
  A 420.
    10.
  r
  k 30.
```

## \*\*process make\_field

## **Description**:

This post is used to create field variables from some other source than directly from a results database. For example, a uniform temperature can be imposed (satisfying post computations requiring that variable be present), or for doing analytical solutions or graphing over a mesh body.

## Syntax:

The following syntax summary is available:

```
**process make_field
 *var name1
[ *single_value val ]
[ **single_field scale-val fname ]
[ *constant_values ]
   time# val#
   ...
[ *constant_field ]
   time# scale-val# fname#
```

Multiple instances of **\*constant\_values** and **\*constant\_field** can be used to mix uniform fields and fields loaded in from an external file.

## Example:

There are examples of this process in the tests plast3.inp and hot\_spot.inp under test/Post\_test/INP. An excerpt of the hot\_spot.inp test case follows, and is a good example of how one can use FEA post processors as a general visualization tool for spatial functions. The data for that test is generated in a small perl script hot\_spot.pl.

```
****post_processing
 ***data_source mesh_only
  **format Z7
  **open hot_spot.geof
  **maps 0. 1. 2. 3.
 ***local_post_processing
  **process make_field
   *var X
   *constant_values
           0.2
     0.
   *constant_field
     1.
           2.
                 hot_spot.dat
     З.
           1.
                 hot_spot.dat
```

## \*\*process mat\_sim

## **Description**:

This post-processor re-simulates material behavior given a strain and parameter (e.g. temperature) history. This post computation is most useful with very large structures and with imported results files. For example, ABAQUS ODB files are much larger than the Z-post files for the same stored results. In that case only the strain need be stored from the FEA run, and additional material state variables can be generated as need-be using this process. This can also be used to re-establish "named" material variables consistent with the material model instead of having SDV indexed names.

## Input:

The full strain history is required in the variable name eto## listed as full tensor. The dimension will be consistent with the finite element geometry dimension. If the name eto is not directly available (for example with abaqus calculations using \*NLGEOM we have LE##) a copy post process can be used beforehand to re-map the name.

If a \*parameter option is given the parameters must be in input results as well. The user has the option to re-map those names using the \*change\_param\_name. For example with abaqus results one would specify \*parameter NT11 and then re-map that with \*change\_param\_name NT11 temperature.

## Output:

The output variables are selected by the user. Before running, the **Zpreload** command can be run on the material file to get a listing of available material variables.

## Syntax:

Basic options for this post are:

```
**process mat_sim
[ *change_param_name from-name to-name ]
[ *every_increment ]
[ *file fname ]
[ *file fname ]
[ *integration INTEGRATION ]
[ *parameter p1 p2 ... pN ]
[ *p_init pval1 pval2 ... pvalN ]
[ *rotation ROTATION ]
[ *save v1 v2 ... vN ]
```

The command descriptions are given below:

\*every\_increment selects if the output will be for every solution map, or just the result at the end of the calculation (or selected maps). If this is not careful use of \*\*output\_number 1-999 can allow the user to investigate full material variables at specific points in time. \*change\_param\_name used to re name a parameter to be consistent with Z-mat names.

\*file specifies the material file (as in standard Z-mat or Z-set input procedures).

\*integration specifies the material integration rule to use rather than the default.

\*parameter specifies parameter names which are to be added and having constant values.

\*p\_init specifies parameter values in sequence that they have been defined which represent the initial values. This is important because often the first FEA result map has already had some loading, and thermal strain for example will be calculated based on the temperature difference from the rest state before that results map.

**\*rotation** specifies the material rotation.

\*save gives a listing of the variables which are desired to be stored in the new results files.

#### Example:

The following example is from Post\_test/INP/disk6\_matsim.inp

```
***local_post_processing
 **output_number 1-999
 **file integ
 **elset ALL_ELEMENT
 **process mat_sim
    *every_increment
    *file disk6_matsim.inp
    *parameter temperature
               800.
    *p_init
    *integration theta_method_a 1.0 1.e-8 100
    *save_variables
      evi11 evi22 evi33 evi12 evi23 evi31
      eto11 eto22 eto33 eto12 eto23 eto31
      sig11 sig22 sig33 sig12 sig23 sig31
      evcum
```

## <u>Note</u>:

The material simulation should always be run from the beginning of the loading history, and following the complete loading path. This is because the material state variables will be tracked, and has substantial impact on the results.

## \*\*process max

## **Description**:

This post computation gives at each node or Gauss point, the maximum value of a variable in the course of time. One output is produced for the totality of the time period for each variable specified.

## Syntax:

\*\*process max
\*list\_var name1 ... nameN

The names name1, ..., nameN designate the list of scalar variables to treat. Outputs appearing in the file *problem.utp* will be named MAX\_name1, ..., MAX\_nameN.

```
% compute the maximum value of the U1 and U2 displacements
**process max
 *list_var U1 U2
% compute von Mises invariant of the stress tensor, ...
**process mises
 *var sig
% ... then its maximum value
**process max
 *list_var mises
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process fmax

## \*\*process fmax

### **Description**:

This post computation produces for each variable specified an output per "map" of time, which corresponds to the maximum value achieved in the loading instant.

## Syntax:

\*\*process fmax
\*list\_var name1... nameN

The names name1, ..., nameN designate the scalar variables to be treated. The output names which are generated will be of the form FMAX\_name1, ..., FMAX\_nameN (see the file problem.utp).

## Example:

% this will provide FMAX\_U1 and FMAX\_U2
\*\*process fmax
\*list\_var U1 U2

## \*\*process min

### **Description**:

This post computation gives at each node or Gauss point, the minimum value of a variable in the course of time. One output is produced for the totality of the time period for each variable specified.

## Syntax:

\*\*process min
 \*list\_var name1 ... nameN

The names *name1*, ..., *nameN* designate the list of scalar variables to treat. Outputs appearing in the file *problem.utp* will be named MIN\_*name1*, ..., MIN\_*nameN*.

```
% compute the minimum value of the temperature
**process min
 *list_var TP
% compute eigenvalues of the stress tensor, ...
**process eigen2
 *var sig
% 3 new variables are now available for the processing,
% sigp3 <= sigp2 <= sigp1
% ... then the minimum value of the two smaller ones
**process min
 *list_var sigp2 sigp3</pre>
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process fmin

## \*\*process fmin

## **Description**:

This post computation produces for each variable specified an output per "map" of time, which corresponds to the minimum value achieved up to each loading instant.

## Syntax:

\*\*process fmin
 \*list\_var name1... nameN

The names name1, ..., nameN designate the scalar variables to be treated. The output names which are generated will be of the form FMIN\_name1, ..., FMIN\_nameN (see the file problem.utp).

```
% this will provides FMIN_TP
**process fmin
    *list_var TP
```

## \*\*process mises

## **Description**:

This process is used to compute the von Mises invariant of a symmetric second order tensor. One value per node or Gauss point is obtained for each requested record. If output\_name is not specified, the name of the output variable is obtained by suffixing mises to the input name.

## Syntax:

\*\*process mises
\*var name

[ \*output\_name output\_name ]

```
% this will provide sigmises
**process mises
*var sig
```

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process multirange

## \*\*process multirange

### **Description**:

This post-processor can be used to analyze complex tridimensional loading paths in order to extract and count cycles (rainflow counting). It can only be applied to tensorial input variables.

The method is based upon multisurface theories of plasticity to detect closed loops (cycles) in the input stress or strain histories and their corresponding amplitude. It is an extension to the general case of multiaxial non-proportional loading paths of the well-known "rainflow technique" conventionally applied to uniaxial loadings. This post-processor can be used in conjunction with post-processors post\_fatigue\_rainflow (calculation of fatigue life for each sub-cycle) and rainflow (cumulation of damage generated by each sub-cycle).

## Syntax:

\*\*process multirange
 \*var name
[\*center ]
[\*reverse number]

name is the input (tensorial) variable which is the subject of the range calculation.

number is the maximum number of sub-cycles given as output (default value is 3). The code gives a warning in the case where the number of cycles detected is greater than the one requested for output.

With the option **\*center**, the code sends back additionally the center of the sphere containing the loading path associated with each subcycle.

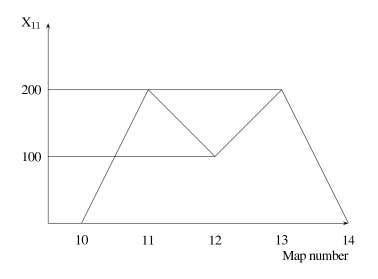
The output is the following:

- ncyc the number of cycles detected. If the algorithm detects more cycles than the number requested by the **\*reverse** number keyword, the code return ncyc = number.
- for each cycle i:
  - Diname the amplitude of cycle *i* for input variable name,
  - niiname is the number of the output map (within the ones selected by the \*\*output\_number command) corresponding to the beginning of cycle i,
  - nfiname is the number of the output map (within the ones selected by the \*\*output\_number command) corresponding to the end of cycle i,
  - optionally if the **\*center** keyword is included, the components of tensor Ciname at the center of the sphere including the loading path for cycle *i*.

Output cycles are stored by decreasing values of amplitude, such that the most meaningful cycles are included even if the number of requested cycles (specified by the **\*reverse** keyword)

is smaller than the one detected by the rainflow algorithm. In the the event where the number of requested cycles is larger than the one detected, all corresponding quantities are zero.

## Example:



Applying the following commands to the loading path of the above figure:

```
**output_number 10-14
**process multirange
 *var X
 *reverse 3
 *center
```

will yield as output:

- ncyc = 2
- D1X = 200, ni1X = 1, nf1X = 4, C1X11 = 100, C1X22 = 0 ...
- D2X = 100, ni2X = 1, nf2X = 3, C2X11 = 150, C2X22 = 0 ...
- $\mathtt{D3X}=0,\,\mathtt{ni3X}=0,\,\mathtt{nf3X}=0,\,\mathtt{C3X11}=0,\,\mathtt{C3X22}=0\ldots$

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process norm

## \*\*process norm

### **Description**:

The norm post processor is used to norm a tensor or scalar variable. It returns a variable of the same type (tensor or scalar) with name constructed by pre-fixing the original variable name(s) with an N.

### Syntax:

\*\*process norm
 \*var name
 [\*type tensor|scalar]

The user must give a material coefficient sigma\_u. The coefficient can be constant or depend on other parameters.

```
% this will provide 4 components :
% Nsig11, Nsig22, Nsig33 and Nsig12
**process norm
    *var sig
% the following syntax must be used in the material file:
**process norm
    sigma_u 170.
```

## \*\*process oxidation

### **Description**:

This post-processor can be used to quantify the effects of oxidation which contributes to the mechanisms of damage.

We calculate, for a cycle, the oxidation damage  $I_{ox}$  with the following expression:

$$I_{ox} = \int_{cycle} \left(\frac{K0}{e0} exp\left(\frac{-Q}{RT}\right) \left[1 + \left\langle\frac{S - S_{lox}}{B}\right\rangle\right]^m\right)^2 dN$$

with

$$S = (1 - \alpha - \beta)\mathbf{Mises}(\sigma) + \alpha \mathbf{Eig1}(\sigma) + \beta \mathbf{Trace}(\sigma)$$

and

$$Slox = Slox0(1 - h\overline{\mathbf{Trace}(\sigma)}) + \overline{\mathbf{Trace}(\sigma)}$$

The output is generated as a single map for the whole loading history. The cycle number is  $N_{ox}$ , defined as the inverse of oxidation damage per cycle and denoted Nox in the output.

## Syntax:

```
**process oxidation
 *var name
[*type scalar | tensor ]
[*delay ]
[*norm
[*normalized_coeff ]]
[*isotherm ]
```

name is the variable name to treat. If the option **\*norm** is specified the subject variable will be normed by the coefficient **sigma\_u** furnished in the material file.

With the option \*delay the lag stress S' will be used. The lag stress is computed from

$$dS' = \frac{S - S'}{\tau}$$

The coefficients B, SloxO, and h are to be given in the material file regardless of the use of normalization. If they are normalized (option \*normalized\_coeff) the expected names in the material file will be  $N_B$ ,  $N_SloxO$  and  $N_h$ .

With the option **\*isotherm** the temperature is read in the material file using a "coefficient" **temperature**. Otherwise the temperature must be included in the output files (use the **\*\*save\_parameter** output option in the FEA calculation.

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process oxidation

alpha and beta are optional coefficients, having a zero value by default. The other coefficients which must be input are m, KO, eO and Q. The additional coefficient tau must be supplied if the \*delay option was input.

```
% a simple example
**process oxidation
  *var sig
  *norm
% The following syntax must be used in the material file :
**process oxidation
  sigma_u 850.
  В
          46.5
          2.5
  m
  Slox0
          0.12
  h
          3.5
          50.e-06
  e0
  KO
          1.42
  Q
          214.
```

## \*\*process rainflow

#### **Description**:

This command can be used in replacement of **\*\*process LCF** to perform damage cumulation within a rainflow counting procedure. Cumulation rules and material parameters are the same as in the case of the **\*\*process LCF** post-processor.

## Syntax:

\*\*process rainflow
[\*reverse number]
 \*mode LC | NLC | NLC\_ONERA
 \*fatigue fatigue\_rainflow section1
[\*creep name2 section2]
[\*scale lin | log ]
[\*load\_cycle cb1-ce1 cb2-ce2 ... cbn-cen ]

The main difference with **\*\*process LCF** is the **\*reverse** keyword, that allows to specify the maximum number of sub-cycles retained in the calculation, within the ones detected by the rainflow counting algorithm (see the **\*\*process multirange** command). Default value is *number*=3. Note that **\*\*process multirange** gives out cycles with maximum amplitude first, such that cumulation is always performed for the most damaging cycles.

Contrary to **\*\*process LCF**, the only fatigue model allowed is the one implemented in the **\*\*process fatigue\_rainflow** post-processor (see this command).

Note also, that the creep damage part is optional in this post-processor, and that cumulation can be restrained only to the fatigue damage induced by the sub-cycles detected by the rainflow algorithm.

The optional command  $*load_cycle$  can be used to define a preloading and calculate the associated initial damage, before cycling occurs. Preloading definitions have the form *cbi-cei*, where *cbi* and *cei* are the map numbers defining respectively the beginning and the end of the preloading sequence number *i*. An arbitrary number of sequence can be given. When using this command, the last sequence *cbn-cen* given as input, is used to define the cycle that will be repeated until failure. The number of cycles to failure calculated are thus the number of times sequence *cbn-cen* can be repeated, taking into account an initial damage corresponding to sequences *cb1-ce1* to *cbn-1-cen-1*. Note that map numbers given as input are in fact *relative*, and depend on the \*\*output\_number command specified before the current \*\*process rainflow definition.

Output is as follows:

• ncyc the number of cycles detected. If the rainflow algorithm detects more cycles than the number requested by the **\*reverse** number keyword, the code returns ncyc = number.

- the number of cycles of failure NFi corresponding to fatigue damage associated with each sub-cycle i.
- if the option \*creep is included the number of cycles to failure NC associated with creep damage,
- the number of cycles to failure  $\mathtt{NR\_mode}$  obtained by culmulation of the previous damage mechanisms,
- the initial damage Dpreload corresponding to preloading sequences when the command \*load\_cycle is used.

# \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process range

## \*\*process range

#### **Description**:

This post-processor calculates the amplitude of a scalar or tensorial variable from its history.

In the case of a tensorial variable, an invariant of the type von Mises is used to calculate the space distance in the six different dimensions, and the result is the diameter of the smallest sphere encompassing the point of interest's path during the loading.

#### Syntax:

```
**process range
 *var name
[*type scalar | tensor ]
[*method num ]
[*alpha valeur ]
[*center ]
[*delta ]
```

name is the input variable which is the subject of the range calculation.

Two methods are available. By default, the first method is applied. If the first method fails, it will automatically roll over to the second method. This later depends on a coefficient alpha which must be between 0 and 1. Its default value is 0.2.

With the option **\*center**, the code sends back additionally a tensor corresponding to the center of the sphere containing the loading path.

With the option **\*delta**, implemented only with the **scalar** keyword, the code just deliver the difference of the prescribed variable between the last two increments.

The generated output will consist of the amplitude named after the input subject prefixed with a D. If the center tensor is output, the prefix C is added.

```
% this will provide Dtemper
**process range
  *type scalar
  *var temper
% this will provide DX
% using directly the second method
% and the center (CX11, CX22, CX33, CX12)
**process range
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process range

\*var X
\*method 2
\*alpha 0.15
\*center

## \*\*process neu\_sehitoglu

### **Description**:

This post-processor implements the thermo-mechanical fatigue model proposed by Neu and Sehitoglu [Neu89]. This model is a bit of a departure from the rest of the fatigue analysis post-processing in that it does not re-use components for LCF, oxidation and creep damages, but rather implements the specific forms defined in the referenced paper.

The damage is a linear summation of three components as per standard practice in thermomechanical fatigue studies.

$$D = D_{fatigue} + D_{oxidation} + D_{creep}$$

And the number of cycles to failure can be found from the cumulated damage over a number of studied cycles,  $N_c$ :

$$N_f = N_c/D$$

The interesting part of the Neu and Schitoglu is their assignment of a damage phase factor, which accounts for the tendency for creep damage in in-phase loading (such as found in rotating machinery or pressure vessels at high temperature), and oxidation cracking in out-of-phase loading (such as around notches in constrained geometries).

$$\phi_{\text{ox}} = \frac{1}{t} \int_{0}^{t_{c}} \exp\left[-\frac{1}{2} \left(\frac{\dot{\epsilon}_{th}/\dot{\epsilon}_{mech}+1}{\xi_{ox}}\right)^{2}\right] dt$$
$$\phi_{\text{cr}} = \frac{1}{t} \int_{0}^{t_{c}} \exp\left[-\frac{1}{2} \left(\frac{\dot{\epsilon}_{th}/\dot{\epsilon}_{mech}-1}{\xi_{cr}}\right)^{2}\right] dt$$

So the coefficients xi\_ox and xi\_cr control the width of oxidation and creep effects about pure out-of-phase and pure in-phase loading respectively.

#### Fatigue part:

The fatigue part can be determined in several ways. In general we look at the strain range as being the maximum shear strain range in from the cycle calculated as follows:

$$\Delta\gamma = \frac{1}{2}(\Delta e_1^{loading} - \Delta e_3^{loading}) + \frac{1}{2}(\Delta e_1^{unloading} - \Delta e_3^{unloading})$$

with the  $\Delta e_1$  and  $\Delta e_3$  terms being the 1st and 3rd ordered eigenvalue of the strain increment from across the cycle. Note that the default method does not take the critical plane of all loading points in the cycle, but rather just the points of the major temperature cycle. If the **\*use\_loading** or **\*use\_unloading** option is given, we use only one term above instead of the average. Normally for a stabilized cycle the loading and unloading parts should be symmetric.

In the above strain range calculation, we also allow for either the total mechanical strain to be used (model neu\_sehitoglu) or the plastic strain range (model neu\_sehitoglu\_evi), and currently there is no HCF (stress based) part taken into account.

Lifetime is predicted via:

$$N_{f}^{fatgue} = \frac{1}{D_{fatigue}} = \frac{1}{2} \left[ \frac{\Delta \gamma}{2\epsilon_{c}'} \right]^{1/c}$$

where we have the coefficients  $eps_f$  and c. If temperature dependent coefficients are given for the fatigue part, the values calculated at the cycle average temperature are used.

## **Oxidation part**:

The oxidation damage is calculated via the following equation:

$$D_{ox} = \frac{1}{N_f^{oxide}} = \left[\frac{\overline{h}_f \delta_o}{B\phi_{ox} K_p}\right]^{-1/\beta} \frac{2(\Delta \epsilon_{mech})^{2/\beta+1}}{\dot{\epsilon}^{1-a/\beta}}$$

Which uses a term for the critical oxide depth:

$$\overline{h}_f = \frac{\delta_0}{(\Delta \epsilon_m)^2 \, \phi_{ox} \, \dot{\epsilon}^a}$$

And an assumed parabolic oxidation rate:

$$K_p = \frac{1}{t_c} \int_0^{t_c} D_o \exp\left[\frac{-Q}{RT(t)}\right] dt$$

For the oxidation part the coefficients to be entered are D\_0, Q, B, beta, delta\_0, a, h\_cr. Note that increasing D\_0 or B increases the damage effect, bigger Q will emphasize an abrupt change in high temperature degrading (but require a change in B to compensate), while decreasing either delta\_0 or h\_cr will increase the oxide damage. These later parameters should be calibrated from primary metallurgical parameters from specialized oxidation tests. If the model is being used empirically then there is quite a great deal of redundancy in these parameters and the user should use caution.

#### Creep part:

The creep damage equation is:

$$D_{cr} = \frac{1}{N_f^{creep}} = \phi_{cr} \int_0^{t_c} A e^{-\Delta H/RT} \left[ \frac{\alpha_1 J(\boldsymbol{\sigma}) + \alpha_2 \frac{1}{3} Tr(\boldsymbol{\sigma})}{K} \right]^m dt$$

with  $J(\boldsymbol{\sigma})$  being the von Mises equivalent stress. The creep damage will occur only in tensile loading under uniaxial conditions if  $\alpha_1 = 1/3$  and  $\alpha_2 = 1$  (these are the defaults).

For the creep part the coefficients to be entered are alpha1, alpha2, A, K, m, dH which should be obvious names from the above equation.

#### Output:

A large output set is generated by this processor, but with only one solution map for the whole history analyzed. The following summarizes the output variables:

Nc the number of cycles detected for validation. There may be zones where the number is incorrect in totally unloaded portions of a model.

- Nf, log\_Nf the life prediction based on the history so far, and log base 10 of that lifetime for flatter contour plots.
- D, Dox, Dcr the "damage" so far in the loading history, with 0 being undamaged, and 1 full failure. D is total, Dox oxidation part, Dcr the creep part.
- extr\_Nf-1 extr\_log\_Nf-1 extrapolations of the lifetime to account for the trend in evolving strain ranges.
- extr\_N-1 the number of cycles forward in the extrapolation.
- range\_ave average value of the plastic strain range (or effective range used for LCF calculations).
- phi\_ox\_ave, phi\_cr\_ave "phase factors" for the oxidation and creep effects as defined by sehitoglu. zero means that effect is not considered because of the type of loading.

## Syntax:

Basic options for this post are:

```
**process neu_sehitoglu
[ *extrapolation_cycles n1 n2 n3 ]
[ *extrapolation_sequence n1 n2 n3 ]
[ *max_cycles maxc ]
[ *model_coef ]
    ...
[ *R R-value ]
[ *skip_first skip ]
[ *small_strain_rate sm-rate ]
[ *use_loading ]
[ *use_unloading ]
```

- \*extrapolation\_cycles gives absolute cycle numbers which will be used to extrapolate the trend in fatigue life. This is used when cycles are not yet stabilized.
- \*extrapolation\_sequence gives a series of cycles ending with the last detected cycle which will be used to extrapolate the trend in fatigue life. This is used when cycles are not yet stabilized.
- \*max\_cycles limits output cycle number as essentially infinite life. This is convenient to limit the ranges for contour plotting.
- \*R sets the gas constant in case of different units. The default value is 8.3144e-3 KJ/(mol K).

- **\*skip\_first** causes the first *skip* cycles to be left from consideration in order to limit damage prediction from large initial transient cycles.
- \*small\_strain\_rate gives a small strain rate which will be used in several locations to limit divide by zero conditions, including the strain rate effect and prediction of the constraint average.
- **\*use\_loading** indicates that only the loading portion of the strain range will be used for fatigue. This would normally be detected as the heat-up side of a TMF cycle.
- **\*use\_unloading** indicates that only the unloading portion of the strain range will be used for fatigue.

Additional controls are allowed for the CYCLE\_TOOL part. Currently these are<sup>3</sup>.

[ \*width wid ]
[ \*major\_tensor tens-name ]
[ \*range\_tensor tens-name ]

\*width gives a size control for detecting cycles. If too few cycles are detected the number is probably too large; too many cycles detected and the size is probably too small (default  $10^{-4}$ ). Units will be that of the major tensor.

\*major\_tensor select the name of the variable to be used for cycle detection (default sig).

**\*range\_tensor** select the name of the variable to be used for strain range calculations (default **eto**).

For the TMF cycle tool (default) the following additional commands are available (however \*major\_tensor and range\_tensor are disallowed).

```
[ *total_strain_for_range ]
```

[ \*mechanical\_strain\_for\_range ]

- [ \*temperature\_unit celcius|fahrenheit|kevin ]
- \*total\_strain\_for\_range detect the cycles based on the total strain. Note for fully
  constrained test cases this will always be zero so the mechanical range should be used.
  Conversely for basically unconstrained conditions using the total strain will be a better
  choice.

\*mechanical\_strain\_for\_range use the mechanical strain for the range calculation.

#### <u>Note</u>:

If the coefficients D\_O, Q, B, beta are not entered the oxidation part will be skipped without mention. Likewise if alpha1, alpha2, A, K are not entered the creep damage will be skipped. Partially entering these coefficients will cause an error message.

<sup>&</sup>lt;sup>3</sup>section to be separated in next version

**Example**: The following example is from Post\_test/INP/sehitoglu1.inp

```
***local_post_processing
  **file integ
  **elset ALL_ELEMENT
  **output_number 1-99999999
  **process neu_sehitoglu_evi
   *total_strain_for_range
   *model_coef
     eps_f 0.20
           -0.64
      С
               1.0
      b
      sig0_E 0.0
      xi_ox
                    1.0
                   0.20
      xi_cr
      a 0.0
      D_0
                2000.
      Q
               200.
      В
           7.00e-03
      beta
                  1.5
      delta_0
                 2.00e-07
                  500.
      h_cr
      alpha1
                    0.33
                    1.0
      alpha2
      А
           1.00e+08
      K temperature
       1.00e+06 0.
       1.00e+06 450.
       20.0
                 600.
       20.0
                 1200.
            2.0
      m
            200.
      dH
```

## \*\*process swt

#### **Description**:

This process allows to calculate number of cycles to failure according to a unified strain-stress fatigue model based on the Smith-Watson-Topper criterion.

The fatigue life is calculated by solving the following equation:

$$\sigma_{eff} = F(N_f)$$

where the effective stress  $\sigma_{eff}$  is calculated as a product of the stress and strain amplitudes multiplied by a correction factor depending on the mean stress:

$$\sigma_{eff} = \sqrt{E\frac{\Delta\sigma}{2}\frac{\Delta\epsilon}{2}}f(\frac{\Delta\sigma}{2},\bar{\sigma})$$

In the above formula E is the material Young's modulus, while:

- $\frac{\Delta\sigma}{2}$  is the multiaxial stress amplitude (measured in terms of the von mises J2 stress invariant) as calculated by the **\*\*process range** post-processor,
- a special treatment is needed for the calculation of the multiaxial strain amplitude  $\frac{\Delta \epsilon}{2}$  in order to be compatible with the unixial case (because of nonzero strains that arise in directions orthogonal to the uniaxial stress direction). The expression used to calculate  $\frac{\Delta \epsilon}{2}$  is the following one:

$$\frac{\Delta \epsilon}{2} = \frac{\Delta \sigma}{2E} + \frac{\Delta \epsilon^p}{2}$$

where the plastic strain amplitude  $\frac{\Delta \epsilon^p}{2}$  is computed either using the **\*\*process range** post-processing, or by means of a cyclic hardening relation  $\Delta \epsilon^p = f(\Delta \sigma)$ , depending on the option selected (see syntax). Note that when the **range** post-processor is used, the distance measuring the amplitude in the deviatoric strain space is calculated by:

$$J2^*(\underline{\epsilon}^p) = \sqrt{\frac{2}{3}\operatorname{dev}(\underline{\epsilon}^p) : \operatorname{dev}(\underline{\epsilon}^p)}$$

the  $\frac{2}{3}$  factor being used instead of the conventional  $\frac{3}{2}$  of the mises stress invariant, for compatibility with the uniaxial case:

$$\epsilon^{p} = (\epsilon^{p}_{11}, \epsilon^{p}_{22}, \epsilon^{p}_{33}, \epsilon^{p}_{12}, \epsilon^{p}_{23}, \epsilon^{p}_{31}) = (\epsilon^{p}_{11}, -0.5\epsilon^{p}_{11}, -0.5\epsilon^{p}_{11}, 0, 0, 0)$$

such that: $J2^*(\underline{\epsilon}^p) = \epsilon_{11}^p$  in this case.

• in the multiaxial case, the mean stress value  $\bar{\sigma}$  is calculated as the mean trace of tensor  $\sigma$ :

$$\bar{\sigma} = \frac{1}{2} \left[ \max(\operatorname{Trace}(\sigma)) + \min(\operatorname{Trace}(\sigma)) \right]$$

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• the mean stress correction factor  $f(\frac{\Delta\sigma}{2}, \bar{\sigma})$  is defined as the following function of  $\bar{\sigma}$ :

– traction :  $\bar{\sigma}>0$ 

$$f(\frac{\Delta\sigma}{2},\bar{\sigma}) = \left(1 + \frac{\bar{\sigma}}{\left(\frac{\Delta\sigma}{2}\right)}\right)^n$$

where n is a model coefficient.

– compression :  $\bar{\sigma} < 0$ 

$$f(\frac{\Delta\sigma}{2},\bar{\sigma}) = \left(1 - \frac{\bar{\sigma}}{\left(\frac{\Delta\sigma}{2}\right)}\right)^{-1}$$

which yields an effect similar to the one predicted by the Sines criterion, or alternatively  $f(\frac{\Delta\sigma}{2}, \bar{\sigma}) = 1$  when the \*skip\_compression option is specified. The latter option ignores any beneficial effect of a compressive stress, and may be too pessimistic.

Several definitions of  $F(N_f)$  are allowed depending on the options:

• the first possibility (keyword \*expression manson) is to derive  $F(N_f)$  from a prior Manson-Coffin model calibration on symmetric cyclic tests (loading factor R=-1,  $\bar{\sigma} = 0$ ).

$$\frac{\Delta \epsilon}{2} = \frac{\Delta \epsilon^p}{2} + \frac{\Delta \epsilon^e}{2} = A N_f^{-\alpha} + B N_f^{-\beta}$$

which yields by replacing  $\frac{\Delta \epsilon^p}{2} = A N_f^{-\alpha}$  and  $\frac{\Delta \epsilon^e}{2} = B N_f^{-\beta}$  in the effective stress formula:

$$F(N_f) = E\sqrt{ABN_f^{-\alpha-\beta} + B^2N_f^{-2\beta}}$$

In that case model coefficients A, B,  $\alpha$ ,  $\beta$  are directly known from the Manson-Coffin calibration, and the only additional coefficient still left to define is the n exponent involved in the mean stress correction factor,

• a second possibility lets the user define an arbitrary expression for  $F(N_f)$ , using the keyword \*expression <FUNCTION>, where <FUNCTION> is a valid function definition. For example, an user-defined expression of  $F(N_f)$  equivalent to the manson option could be specified by:

```
*expression E*(A*B*Nf^(-alpha-beta) + B*B*Nf^(-2.0*beta)^(0.5);
```

Note that it is mandatory that a variable named Nf or nf should be used in the function definition. Other function coefficients are assumed to be model coefficients and should be given in the post-processing material file.

## Introduction of a fatigue limit effect

An additional model coefficient  $\epsilon_D$  may be used to add a fatigue limit effect when the manson option is selected. In this case the Manson-Coffin model used during calibration is rewritten:

$$\frac{\Delta \epsilon}{2} = \frac{\Delta \epsilon^p}{2} + \frac{\Delta \epsilon^e}{2} + \epsilon_D$$

such that  $N_f = \infty$  when  $\frac{\Delta \epsilon}{2} < \epsilon_D$ .

In that case the  $F(N_f)$  expression used in the swt model becomes:

$$F(N_f) = E\sqrt{ABN_f^{-\alpha-\beta} + B^2N_f^{-2\beta} + \epsilon_D\left(2BN_f^{-\beta} + AN_f^{-\alpha} + \epsilon_D\right)}$$

Syntax:

```
**process swt
    *var name_sig
    [*type scalar | tensor ]
    [*expression manson | func_exp ]
    [*derivative func_deriv ]
    *plastic_strain name_ep
| *cyclic hardening function func_hard
| *cyclic hardening file fname sig_col eto_col
    [*precision prec ]
    [*iter iter ]
    [*skip_compression ]
```

name\_sig is the name of the variable used to store the stress in the results file (one would expect sig here).

The \*expression command allows to define the  $F(N_f)$  function. Default is \*expression manson that defines a  $F(N_f)$  expression derived from a Manson-Coffin calibration as detailed above. A user-defined expression may alternatively be given by means of argument func\_exp.

The \*derivative command allows to define the derivative  $\frac{dF(N_f)}{dN_f}$  of  $F(N_f)$  in the func\_deriv argument, and may be needed when a user defined function is given after \*expression. In this case a Newton-Raphson method is used to solve the  $\sigma_{eff} = F(N_f)$  equation to calculate  $N_f$ . Otherwise, a slower dichotomy method is used. Note that the \*derivative command is not needed in the default \*expression manson mode, and that the Newton-Raphson method is always used in this case.

The \*plastic\_strain command is used to define the name of the variable where the plastic strain  $\epsilon_{\sim}^{\mathbf{p}}$  is stored in the results files. In this case the  $\frac{\Delta \epsilon^{p}}{2}$  amplitude needed in the effective stress expression will be calculated by applying the range post-processor on the variable specified by the argument name\_ep.

Alternative specification of  $\frac{\Delta \epsilon^p}{2}$  calculation can be given by means of the **\*cyclic\_hardening function** command. In that case a function  $\frac{\Delta \epsilon^p}{2} = f(\frac{\Delta \sigma}{2})$  is expected for the *func\_hard* argument.

 $\frac{\Delta \epsilon^p}{2}$  may also be interpolated from cyclic hardening curve results by means of the **\*cyclic\_hardening file** command. In this case argument *fname* is the name of the file containing the cyclic hardening curve, and *sig\_col*, *eto\_col* are the column numbers in this file used to store  $\frac{\Delta \sigma}{2}$  and  $\frac{\Delta \epsilon^p}{2}$  repectively. Note that the half-amplitude is expected for both stress and strain values.

The optional \*precision command defines the precision required when solving the  $\sigma_{eff} = F(N_f)$  by either dichotomy or Newton-Raphson. Default value is  $10^{-4}$ . Similarly, the maximum number of iterations allowed may be given after \*iter command (default is 500). In general, the Newton-Raphson method converges in at most a few dozens of iterations, while the dichotomy method may need larger values depending on the precision required.

When the optional \*skip\_compression command is given, the mean stress correction factor  $f(\frac{\Delta\sigma}{2}, \bar{\sigma})$  is set to 1.0, negative value of the mean stress  $\bar{\sigma}$ .

Default output gives the number of cycles to failure in a variable named Nf, while the effective stress is stored in a variable named "name\_sig\_swt", obtained by concatenation of the stress tensor name "name\_sig" specified with the "\_swt" character string.

The optional **\*full\_output** command will add the following results to the output:

- the stress amplitude under the name "name\_sig\_alt"
- the average stress under the name "name\_sig\_ave" where "name\_sig" is the name of the stress tensor specified
- the elastic and plastic amplitudes in variables eel\_alt and ein\_alt

#### Example:

```
% standard definition using the default *expression manson option
**process swt
    *var sig
```

```
*plastic_strain ev
*full_output
```

% The following syntax must be used in the material file in this case: \*\*\*post\_processing\_data

## \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process swt

```
**process swt
  young 98000.00
  alpha 0.7438 % mandatory Manson-Coffin coefficients
  A 165.e-2
  beta 0.0745
  B 0.8015e-2
  epsd 0.13e-2 % optional fatigue limit coefficient
  n 0.65
                 % mandatory mean stress correction coefficient
\% user definition of the F(Nf) function
**process swt
   *var sig
  *expression Au*Nf^(-au)+Bu*Nf^(-bu);
  % optional derivative specification to allow faster newton-raphson resolution
  *derivative -au*Au*Nf^(-au-1.0)-bu*Bu*Nf^(bu-1.0);
  *plastic_strain ev
  *full_output
\% The following syntax must be used in the material file in this case:
\% note the Au, au, Bu, bu coefficients needed depend of the user expression
***post_processing_data
 **process swt
  young 98000.00
  au 0.7438
  Au 165.e-2
  bu 0.0745
  Bu 0.8015e-2
  n 0.65
% definition of a cyclic hardening function
**process swt
  *var sig
  % note that the function variable should be called "dsig"
  *cyclic_hardening function (dsig/K)^m;
  *full_output
\% The following syntax must be used in the material file in this case:
% note the K, m coefficients needed depend on the *cyclic_hardening user definition
***post_processing_data
 **process swt
  young 98000.00
  alpha 0.7438
  A 165.e-2
  beta 0.0745
  B 0.8015e-2
  epsd 0.13e-2
  n 0.65
```

\*\*\*\*post\_processing
\*\*\*local\_post\_processing
\*\*process swt

K 1000. m 0.07

## \*\*\*\*post\_processing \*\*\*local\_post\_processing \*\*process trace

## \*\*process trace

## **Description**:

This post-computation calculates the trace of a second order tensor. One value per node or per Gauss point is added to the output records, for each solution map. The new variable name is constructed using the tensor name suffixed with ii.

## Syntax:

\*\*process trace
 \*var name

```
% this will provide eplii
**process trace
 *var epl
```

## \*\*process transform\_frame

### **Description**:

This post calculation calculates existing variables in a new local frame. Variables can be either tensors or vectors.

## Syntax:

\*\*process transform\_frame

\*local\_frame type

[ \*tensor\_variables tensor\_var1 tensor\_var2 ...]

```
[ *vector_variables vector_var1 vector_var2 ...]
```

[ \*suffix suffix ]

[ \*output\_variables name1 name2 ... ]

\*local\_frame specifies the local frame type. type can be euler, cartesian, cylindrical, spherical or be specified by a z7p script. Refer to \*\*local\_frame (page 3.146) for more details and syntax.

**\*tensor\_variables** is the list of tensor variables to transform.

**\*vector\_variables** is the list of vector variables to transform.

**\*suffix** this suffix is added to the variable names (default is **-rot**);

\*output\_variables can alternatively be specified to specify a different output name.

```
****post_processing
***local_post_processing
**file node
**nset ALL_NODE
**process transform_frame
    *local_frame cylindrical
        (1. 2.)
        *tensor_variables sig eto
        *vector_variables U
****return
```

## \*\*process tresca

## **Description**:

This processor computes the Tresca criterion of a symmetric second order tensor. One output variable is generated for each point at each map. The output variable name is constructed from the input variable by adding tresca.

## Syntax:

\*\*process tresca

\*var name

```
% this will provide sigtresca
**process tresca
*var sig
```

## \*\*process triax

## **Description**:

This computes the triaxiality of a symmetric second order tensor. The expression is as follows

## **SMEAN/SEQ**

One value is produced per input point. The new variable name will be constructed from the input variable by adding triax.

## Syntax:

\*\*process triax

\*var name

## Example:

% this will provide sigtriax
\*\*process triax
\*var sig

## \*\*\*global\_post\_processing

## **Description**:

This command is used to define "global" post processing treatments to apply over sets of nodes and integration points. Global post processors produce global values saved in the file *problem*.post and/or in local (field) variables in the node or integration point files. This distinction is indicated by the sign <sup>G</sup> for global variable producing processors, and <sup>L</sup> for local variable producing processors. The reason for the distinction is that file data will be loaded in a manner which may be less efficient than the local post processing methods.

## Syntax:

```
***global_post_processing
[ **deformed ] GLOBAL OPTION
[ **elset eset ]
[ **file file-key ]
[ **ipset ipset ]
[ **ipset ipset ]
[ **material_file fname ]
[ **nset nset ]
[ **output_number out-num-list ]
[ **process type ]
....
[ **undeformed ] GLOBAL OPTION
```

The different sub-options define the geometrical groups of concern, the time period, materials files, and the post treatments themselves. The majority of these options are exactly the same as for **\*\*\*local\_post\_processing**. The following *additional* options are relevant to the global post processing.

**\*\*deformed** Use the mesh in its deformed configuration.

**\*\*undeformed** Use the mesh in its undeformed configuration.

The following global post computations are available in the current release:

average	average_around	average_in_element
beremin	beremin_max	continue_curve
coordinates	сору	couple
crack_front	curve	cylindrical
extract2d3d	format	gil_sevillano
gp_vol	gp_xyz	gradient

input_damage	max	min
mm_localization	momentum	node_interpolation
selective_int	static_torsor	surface_normals
test_gp	torque	transform_frame
volume	volume_above	weibull
z7p		

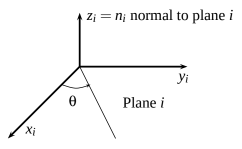
### \*\*process anisotropic\_failure

#### **Description**:

This process is an extension of the weibull or batdorf models to the case of anisotropic brittle materials. A probability of failure  $P_f^i$  is evaluated on an arbitrary number of characteristic planes for the material (eg. particular cristallographic planes), and the total probability of failure  $P_f$  is then computed in the following way:

$$P_f = 1 - \prod_i \left(1 - P_f^i\right)$$

A particular plane *i* is characterized by its normal vector  $\underline{\mathbf{n}}_i$  and the weibull model parameters (strength function  $\sigma_u^i(\theta)$  and modulus  $m(\theta)$ ) depend on an angle  $\theta$  that defines a particular direction in this plane:



The probability of failure  $P_f^i$  associated to failure on plane *i* is then calculated by means of one of the following equations:

• Weibull mode:

$$P_f^i = 1 - exp\left(-\int_V \frac{dV}{V_0} \frac{1}{\theta_{max}} \int_0^{\theta_{max}} \left(\frac{\sigma_e(\theta)}{\sigma_u^i(\theta)}\right)^{m^i(\theta)} d\theta\right)$$

In the above equation, the equivalent stress  $\sigma_e(\theta)$  is chosen as the normal stress in direction  $\theta$ :

$$\sigma_e(\theta) = \underline{\mathbf{d}}_{\theta} \cdot \underline{\sigma} \cdot \underline{\mathbf{d}}_{\theta}$$

where  $\sigma_{\alpha}$  is the stress tensor at the current integration point.

The modulus  $m^i(\theta)$  and the scaling function  $\sigma_u^i(\theta)$  are arbitrary functions of  $\theta$  that should be defined by the user. Coefficients of this function are automatically added as model coefficients by the post-processor, and need to be given in the material file. For example, one can express  $\sigma_0^i(\theta)$  in term of the strength s0 ( $\theta = 0$ ) and s90 ( $\theta = \frac{\pi}{2}$ ) in the following way:

$$\sigma_u^i(\theta) = s0 \cos^2(\theta) + s90 \sin^2(\theta)$$

with material coefficients s0 and s90 declared in the material file.

```
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```

## • Batdorf mode:

This mode is a slight modification of the previous one, where Batdorf-type statistics are preferred (integration over all possible crack sizes, characterized by their critical stres  $\sigma_c$ ):

$$P_f^i = 1 - exp\left(-\int_V \frac{dV}{V_0} \frac{2}{\pi} \int_0^{\frac{\pi}{2}} \left\{\int_0^{\sigma_I} P(\sigma_c, \theta) g(\sigma_c, \sigma_e(\theta)) d\sigma_c\right\} d\theta\right)$$

where:

- \* The equivalent stress  $\sigma_e(\theta)$  is taken as the normal stress in direction  $\theta$  (same definition as in the previous mode)
- \*  $P(\sigma_c, \theta)$  is the crack density with critical stress  $\sigma_c$ . As in the conventional batdorf model this quantity is expressed as a power law, but depends on  $\theta$  by means of the anisotropic scale parameter  $\sigma_u^i(\theta)$ :

$$P(\sigma_c, \theta) = \left(\frac{\sigma_c}{\sigma_u^i(\theta)}\right)^{m_i}$$

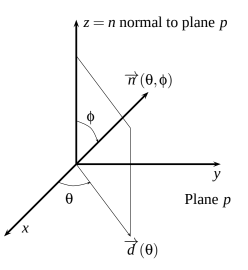
\*  $g(\sigma_c, \sigma_e(\theta))$  is equal to 1 when the equivalent stress is greater than the critical stress  $\sigma_c$ :

$$g(\sigma_c, \sigma_e(\theta)) = 1$$
 if  $\sigma_e(\theta) >= \sigma_c$ , else  $g(\sigma_c, \sigma_e(\theta)) = 0$ 

• Full Batdorf mode:

This mode fully generalizes the Batdorf approach to the anisotropic case, and allows some flexibility in the choice of the equivalent stress  $\sigma_e$ . Integration is taken over the whole space, using 2 angles  $\theta$  and  $\phi$  as represented in the following figure, where:

- $-\overrightarrow{d}(\theta)$  is the direction at angle  $\theta$  in the cristallographic plane as before,
- $-\phi$  is the angle between the z axis (normal to the cristallographic plane) and a particular  $\overrightarrow{n}(\theta, \phi)$  direction in plane  $(\overrightarrow{d}(\theta), z)$



The probability of failure is then computed by the following equation:

$$P_f = 1 - exp\left[-\int_V \frac{1}{V_0} \, dV \, \int_0^{\sigma_I} d\sigma_c \left\{\frac{1}{2\pi} \int_0^{\pi} d\theta \int_0^{\pi} P(\sigma_c, \theta, \phi) \, g(\sigma_e(\theta, \phi), \sigma_c) \, \sin(\phi) \, d\phi\right\}\right]$$

where:

(

- for each gauss point, integration is performed on the half space only (which correspond to a solid angle of  $2\pi$ ) because of symmetry of the equivalent stress  $\sigma_e(\theta, \phi)$
- $P(\sigma_c, \theta, \phi)$  is the density of cracks having a critical stress  $\sigma_c$ :

$$P(\sigma_c, \theta, \phi) = \frac{1}{\sigma_u(\theta, \phi)} \left( \frac{\sigma_c}{\sigma_u(\theta, \phi)} \right)^{m(\theta, \phi)}$$

where coefficients  $\sigma_u$  and m depend on the angles  $(\theta, \phi)$  defining the current direction  $\overrightarrow{n}(\theta, \phi)$ .

 $-g(\sigma_e(\theta, \phi), \sigma_c)$  is equal to 1 when the equivalent stress is higher than the crack critical stress:

$$g(\sigma_e(\theta,\phi),\sigma_c) = 1$$
 if  $\sigma_e(\theta,\phi) > \sigma_c$ , else  $g(\sigma_e(\theta,\phi),\sigma_c) = 0$ 

- The equivalent stress  $\sigma_e(\theta, \phi)$  is allowed to depend both on the normal stress in direction  $\vec{n}(\theta, \phi)$  and the norm of the shear stress vector acting on the facet normal to  $\vec{n}$ :

$$\begin{aligned} \sigma_n(\theta,\phi) &= \overrightarrow{n}(\theta,\phi) \cdot \overrightarrow{\sigma} \cdot \overrightarrow{n}(\theta,\phi) \\ \overrightarrow{\tau}(\theta,\phi) &= \overrightarrow{\sigma} \cdot \overrightarrow{n}(\theta,\phi) - \sigma_n(\theta,\phi) \overrightarrow{n}(\theta,\phi) \\ \tau(\theta,\phi) &= \|\overrightarrow{\tau}(\theta,\phi)\| \end{aligned}$$

The implementation then allows to choose between several expression for the equivalent stress  $\sigma_e$ :

- \* normal stress ("ns") :  $\sigma_e(\theta, \phi) = \sigma_n(\theta, \phi)$
- \* critical energy ("ce") :  $\sigma_e(\theta, \phi) = \sqrt{\sigma_n(\theta, \phi)^2 + \tau(\theta, \phi)^2}$
- \* maximal strength ("ms") :  $\sigma_e(\theta, \phi) = \frac{1}{2} \left( \sigma_n(\theta, \phi) + \sqrt{\sigma_n(\theta, \phi)^2 + \tau(\theta, \phi)^2} \right)$

#### Syntax:

```
**process anisotropic_failure
 *var name
[ *mode weibull | batdorf | [ full_batdorf ( ns | ce | ms ) ] ]
 *plane
 [ tmin tmin ]
 [ tmax tmax ]
 normal <VECTOR>
 function <FUNCTION>
```

```
[ m_function <FUNCTION> ]
[ phi_function <FUNCTION> ]
[ phi_m_function <FUNCTION> ]
[ *plane
    ... ]
[ *orientation | *local_frame <LOCAL_FRAME> ]
[ *nb_constant_step steps ]
[ *nb_sigc_step sc_steps ]
[ *precision eps ]
```

name is the name of the variable used to store the stress in the results file (one would expect sig here)

The \*mode command allows to select the model used to calculate the probability of failure. Possible options, as detailed before, are: weibull (default), batdorf or full\_batdorf. In the full\_batdorf mode an additional keyword is needed to define the choice of the equivalent stress expression. As decribed before, candidates are: ns (normal stress), ce (critical energy) or ms (maximum strength).

The **\*plane** command is used to define the material (or crystallographic) planes characterizing anisotropy of the failure properties. Note that as many **\*planes** definitions as needed may be included. The total probability of failure is then computed as:

$$P_f = 1 - \prod_i \left( 1 - P_f^i \right)$$

where  $P_f^i$  is the probability of failure computed on a particular plane (numbered *i* in the previous equation). Subcommands of the **\*plane** keyword are the following ones:

• the optional commands tmin and tmax allow to define the bounds (tmin and tmax values in degrees) for the  $\theta$  angle integration:

$$P_f = 1 - exp\left\{-\int_V \left(\frac{1}{\theta_{max} - \theta_{min}} \int_{\theta_{min}}^{\theta_{max}} \dots \ d\theta\right) dV\right\}$$

Default values are tmin=0 and tmax=90.

• a VECTOR object is expected after the **normal** keyword to define the normal to the current plane.

```
*plane
normal (0.0 0.0 1.0)
```

• a FUNCTION object is expected after the function keyword to define the  $\theta$  dependence of the strength weibull parameter  $\sigma_u(\theta)$ .

```
*plane
function s0*(cos(theta)^2.0)+s90*(sin(theta)^2.0);
```

As is conventional in Zebulon input files, the function definition must be ended by a ";" character, and all libc mathematical functions are allowed (sin, cos, sinh, log, exp, ...). Note that in this particular context a "theta" character string is expected in the definition for the name of the function variable. All other parameters in the function definition are assumed to be coefficients (eg. s0 and s90 in the previous example) and need to be given in the material file.

- an optional m\_function keyword can be used to declare a dependence of the *m* weibull modulus in term of the direction in the current plane. Conventions for the function definition are the same as in the previous case. Without this option, *m* is assumed to be a constant in all directions. The name of the coefficients expected in the material file are then the following one: m in the case of a single plane, or m1, m2 ... when several planes are declared.
- In the full\_batdorf mode only, additional function definitions are needed to specify the values of the  $\sigma_u$  and m parameters in a particular  $(\theta, \phi)$  direction. Corresponding keywords are phi\_function (for  $\sigma_0$ ) and phi\_m\_function (for m).

```
*plane
function s0*(cos(theta)^2.0)+s90*(sin(theta)^2.0);
phi_function stheta*(sin(phi)^2) + sc*(cos(phi)^2);
```

Functions in the previous example will define  $\sigma_u(\theta, \phi)$  in the following way:

$$\sigma_u(\theta) = s0 \cos^2(\theta) + s90 \sin^2(\theta)$$
  
$$\sigma_u(\theta, \phi) = \sigma_u(\theta) \sin^2(\phi) + sc \cos^2(\phi)$$

where stheta is a predefined name denoting the result of the  $\sigma_0$  function in a ( $\theta, \phi = 0$ ) direction, and s0, s90, sc are model coefficients that need to be declared in the material file. As before, those coefficients are arbitrarily named by the user, and automatically added to the model definition after parsing the function expressions.

The optional command \*orientation or \*local\_frame command allows to transform the stress tensor  $\sigma$  from the global frame to a local material frame prior to volume integration. Such a material frame is in general more convenient to define the anisotropic failure coefficients.

The default method used to perform integration over the  $\theta$  angle is a simple constant step method with a midpoint rule. The default number of steps is 20, and this value can be modified by means of the **\*nb\_constant\_step** command.

Similarly, integration on the critical stress  $\sigma_c$  in the batdorf and full\_batdorf modes is done by a constant step method and the number of steps involved can be specified by means of the \*nb\_sigc\_step command (default value is 20). Note that when of value of 0 is used as argument of the previous two commands, a second-order runge-kutta method will be used to perform the corresponding integrations. This method is more precise and will automatically adapt the size of the steps to verify a target integration error. However, CPU times may be too long for practical applications, in particular in the batdorf or full\_batdorf modes, and this option should only be used in debug mode to check the implementation against analytical solutions. Experience has shown that a constant step method with a default value of 20 steps is in general sufficient to provide reasonable accuracy. However, if runge-kutta integration is defined (by setting \*nb\_constant\_step 0 and/or \*nb\_sigc\_step 0) the command \*precision eps may to set the precision of the runge-kutta method (default value is eps=1.e-4).

For each plane *i* the probability of failure  $P_{ip}^i$  associated to each integration point is stored in the results file under the name Pname\_plane*i*, where name is the variable name (specified by the \*var command) and *i* the plane index. The result of the volume integration of the probability of failure on each plane, and the total  $P_f$  for the complete set of planes is written in the .post file.

```
**file integ % mandatory
    **process anisotropic_failure
     *mode weibull
     *var sig
     *plane
      tmin 0.0 tmax 90.0
      normal (0.0 0.0 1.0)
                 sm*(cos(3.0*theta)^2.0)+sa*(sin(3.0*theta)^2.0);
      function
      m_function mm*(cos(3.0*theta)^2.0)+ma*(sin(3.0*theta)^2.0);
     *plane
      tmin 0.0 tmax 90.0
      normal (0.0 1.0 0.0)
                 sc*(cos(theta)^2)+sm*(sin(theta)^2);
      function
      m_function mc*(cos(theta)^2.0)+mm*(sin(theta)^2.0);
     *plane
      tmin 0.0 tmax 90.0
      normal (1.0 0.0 0.0)
                 sa*(cos(theta)^2)+sc*(sin(theta)^2);
      function
      m_function ma*(cos(theta)^2.0)+mc*(sin(theta)^2.0);
    % material file
***post_processing_data
  **process anisotropic_failure
    VO
        2.0
        4.0
    mm
        3.5
    ma
        3.2
    mc
        880.0
    \mathtt{sm}
```

\*\*\*\*post\_processing
\*\*\*global\_post\_processing
\*\*process anisotropic\_failure

sa 1200.0 sc 2000.0 \*\*\*return

```
**file integ % mandatory
    **process anisotropic_failure
     *mode full_batdorf ms
     *var sig
     *plane % m,a
      tmin 0.0 tmax 90.0
      normal (0.0 0.0 1.0)
      function
                 sm*(cos(3.0*theta)^2.0)+sa*(sin(3.0*theta)^2.0);
      m_function mm*(cos(3.0*theta)^2.0)+ma*(sin(3.0*theta)^2.0);
      phi_function stheta*(sin(phi)^2) + sc*(cos(phi)^2);
      phi_m_function mtheta*(sin(phi)^2) + mc*(cos(phi)^2);
    % material file
***post_processing_data
**process anisotropic_failure
    VO 2.0
       4.0
   mm
       3.5
    ma
        3.2
    mc
       880.0
    \mathtt{sm}
    sa
       1200.0
        2000.0
    sc
***return
```

# \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process average

## \*\*process average

## **Description**:

This post computation calculates the spatial average of variables specified over a group of elements (elset) for each solution step. The average of x is defined as:

$$\bar{x} = \frac{1}{V} \int_{V} x dV$$

## Syntax:

\*\*process average

\*list\_var name1 ... nameN

name1, ... nameN are scalar values to be averaged.

## Example:

\*\*process average
 \*list\_var sig11 epcum sigmises

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process momentum

## \*\*process momentum

## **Description**:

This post calculation is used to find then *n*-th order moment of specified variables over a group of elements (elset) at each solution time. The moment of order n > 1 of x is defined as:

$$\sigma_n(x) = \left(\frac{1}{V} \int_V (x - \bar{x})^n \, dV\right)^{1/n}$$

### Syntax:

\*\*process momentum
 \*order n
 \*list\_var name1 ... nameN

n is the order (integer value), and name1, ... nameN are scalar variable names to treat.

## Example: Example:

```
**process momentum
 *order 2
 *list_var sig11 epcum sigmises
```

\*\*\*\*post\_processing
\*\*\*global\_post\_processing
\*\*process average\_around

## \*\*process average\_around

### **Description**:

This post processor calculates an average around each integration point in an element set and assigns this average value to each integration point.

### Syntax:

\*\*process average\_around
 \*length value
 \*list\_var name1 ... nameN

value is a real value to indicating the distance to take into account around each integration point to make the average. *name1*, ... *nameN* are the scalar variable names to average. The output variable names are generated by adding the prefix **aa\_** to each input variable name.

Symmetries are not taken into account. The method is not valid for axisymmetric geometries.

## Ś

```
% this will provide aa_epcum
**process average_around
*list_var epcum
*length 1.5
```

## \*\*process average\_in\_element

## **Description**:

This post processor calculates the average of the specified variables in each element of an elset, and assigns the mean value to each integration point.

## Syntax:

\*\*process average\_in\_element

\*list\_var name1 ... nameN

where *name1*, ... *nameN* are the scalar variables to average. The output names are created by adding the prefix **ae\_** to each variable.

## Example:

% this will provide ae\_epcum
\*\*process average\_in\_element
\*list\_var epcum

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process batdorf

### \*\*process batdorf

#### **Description**:

This process computes a probability of failure for brittle materials according to the Batdorf model. This model is an extension of the Weibull model that may be more relevant in the case of multiaxial stress states, and also allows some flexibility as regards of the definition of the equivalent stress causing failure.

The probability of failure  $P_f$  is expressed in the following form;

$$P_f = 1 - exp\left(-\int_V dV \int_0^{\sigma_1} \theta(\sigma_c) \frac{\Omega}{4\pi}(\sigma_c) d\sigma_c\right)$$

where:

- the integration  $\int_0^{\sigma_1} \dots d\sigma_c$  is computed over all possible crack sizes, characterized by the critical stress  $\sigma_c$  ( $\sigma_1$  is the maximum principal stress et the current integration point),
- $\theta(\sigma_c)$  is the density of cracks having  $\sigma_c$  as critical stress A classical power law is chosen for  $\theta(\sigma_c)$ :

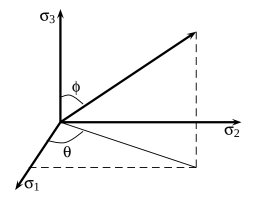
$$\theta(\sigma_c) = \frac{1}{V_0 \sigma_u} \left\langle \frac{\sigma_c - \sigma_0}{\sigma_u} \right\rangle^m$$

where  $V_0$ ,  $\sigma_u$ ,  $\sigma_0$  and m are material parameters.

•  $\Omega(\sigma_c)$  is the solid angle of the region of space where the equivalent stress  $\sigma_e$  is higher than the critical stress  $\sigma_c$ 

 $(\Omega = 4\pi$  for the whole space around the current integration point in the volume)

To evaluate the solid angle  $\Omega(\sigma_c)$  the following coordinate system that depends on the principal stress values  $\sigma_1$ ,  $\sigma_2$ ,  $\sigma_3$  ( $\sigma_3 \ll \sigma_2 \ll \sigma_1$ ) at the current integration point is defined:



and the following expression is used to calculate  $\Omega(\sigma_c)$ :  $\frac{\Omega(\sigma_c)}{4\pi} = \frac{1}{2\pi} \int_0^{\pi} \left( \int_0^{\pi} g(\sigma_e(\theta, \phi), \sigma_c) \sin\phi \ d\phi \right) \ d\theta$ 

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In the previous equation  $g(\sigma_e(\theta, \phi), \sigma_c)$  indicates if the equivalent stress  $\sigma_e$  is greater than the critical stress  $\sigma_c$ :

 $g(\sigma_e(\theta,\phi),\sigma_c)=1 \ \, \text{if} \ \, \sigma_e(\theta,\phi)>=\sigma_c \ \, , \ \, \text{else} \ \, g(\sigma_e(\theta,\phi),\sigma_c)=0$ 

- Various expression of the equivalent stress  $\sigma_e$  may be chosen:
  - normal stress mode ("ns") :  $\sigma_e = \sigma_n$
  - critical energy release rate mode ("ce") :  $\sigma_e = \sqrt{\sigma_n^2 + \tau^2}$
  - maximal strength mode ("ms") :  $\sigma_e = \frac{1}{2} \left( \sigma_n + \sqrt{\sigma_n^2 + \tau^2} \right)$

where  $\sigma_n$  an  $\tau$  are the normal and shear stress components on a facet with normal  $\vec{n}(\theta, \phi)$ . In the coordinate system of the above figure, the following expressions are used to evaluate  $\sigma_n$  and  $\tau$  from the principal stress values:

$$\sigma_n = \sigma_1 \cos^2\theta \sin^2\phi + \sigma_2 \sin^2\theta \sin^2\phi + \sigma_3 \cos^2\phi$$
  
$$\sigma_n^2 + \tau^2 = \sigma_1^2 \cos^2\theta \sin^2\phi + \sigma_2^2 \sin^2\theta \sin^2\phi + \sigma_3^2 \cos^2\phi$$

Syntax:

```
**process batdorf
 *var name
[ *mode ns | ce | ms ]
[ *steps st ]
[ *precision eps ]
[ *nb_constant_step const ]
[ *force_numeric ]
```

name is the name of the variable used to store the stress in the results file (one would expect sig here)

The **\*mode** command allows to select the type of equivalent stress used in the model as defined above. Mode **ns** (normal stress) is the default.

The optional command **\*steps** is used to defined the number of sub-steps involved in runge-kutta integrations of the various integrals defined above. The default value is st=4 (eg. 4 substeps of  $\frac{\pi}{4}$  radians for an integration from 0 to  $\pi$ ).

eps is the runge-kutta precision required for the previous integrations. Default value is eps=1.e-4, that may be decreased to 1.e-3 for faster integration at the expense of a lower precision of the failure probabilities calculated by the post-processor.

In the ns and ce modes, analytical expressions can be derived and are used to accelerate the calculation of the  $\Omega(\sigma_c)$  critical solid angle. In that case a double integration is needed (integration over  $\sigma_c$  and  $\theta$  only) to evaluate  $\Omega$ . In the ms mode, no analytical acceleration is possible, and an additional integration over angle  $\phi$  is needed. Use of runge-kutta for this inner integration is not manageable for efficiency reasons, and a simple constant step integration method is preferred in this case. The  $*nb_constant_step$  optional command may then be used to specify the number of sub-steps needed for  $\phi$  integration. The default value is const=20.

The optional command \*force\_numeric may be used in the ns and ce mode to replace the previous analytical  $\phi$  integration by the numeric constant step method.

The probability of failure  $P_{ip}$  associated to each integration point:

$$P_{ip} = \int_0^{\sigma_1} \theta(\sigma_c) \; \frac{\Omega}{4\pi}(\sigma_c) \; d\sigma_c$$

is stored in the results file under a name constructed by adding " $\_ba$ " to the variable name, while the total probability  $P_f$  obtained after integration on the volume of the component is written in the .post file.

```
**file integ % mandatory
**process batdorf
*mode ns
*var sig
% material file
**process batdorf
VO 1.0
sigma_u 200.0
m 3.0
```

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process beremin

## \*\*process beremin

### **Description**:

This command performs the Weibull stress and the rupture probability according to the Beremin model (see Beremin F.M., "A local criterion for cleavage fracture of a nuclear pressure vessel steel", Met. Trans. A, 14A, 2277–2287 (1983)). The Weibull stress is defined as:

$$\sigma_W = \left(\frac{1}{V_0} \int_{V, p > p_c} \sigma_I^m \, dV\right)^{1/m}$$

where  $V_0$ ,  $p_c$  and m are material parameters.  $\sigma_I$  is the stress tensor maximum principle stress. The integral is taken over the volume where the plastic strain is higher that a critical value  $p_c$ . The rupture probability  $P_r$  is given by:

$$P_r = 1 - \exp\left(-\left(\frac{\sigma_W}{\sigma_u}\right)^m\right) = 1 - \exp\left(-\frac{1}{V_0}\int_{V, p > p_c} \left(\frac{\sigma_I}{\sigma_u}\right)^m dV\right)$$

 $\sigma_u$  is a material parameter.

#### Syntax:

**process	beremin
*stress	name1
*strain	name2

name1 is the name of the stress tensor and name2 the name of the inelastic deformation measure (scalar). The material file must contain the values  $V_0$ ,  $p_c m$  and  $\sigma_u$ .

```
% input file
**process beremin
*stress sig
*strain epcum
% material file
**process beremin
V0 10.
m 20.
sigma_u 1200.
p_c 0.01
```

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process clip\_image

## \*\*process clip\_image

## **Description**:

This command is used to generate maps of Gauss point variables inside a predefined cut plane. Produced images are 256 colors gif files where the value for each pixel is the value of the closest Gauss point. It is also possible to obtain the mean of the element Gauss point values. This process can be quite long as the image is generated finding the element location of each pixel in the structure.

## Syntax:

\*\*process clip\_image

\*list\_var var1 var2 ...

\*output output-file-name

- \*P0 vector
- \*P1 vector
- \*P2 vector
- [ \*ortho ortho-dir-value ]
- [ \*step spatial-step-value ]
- [ \*dimx dimx-value ]
- [ \*dimy dimy-value ]

```
[ *color_map color-map-name ]
```

- [ \*min\_max mode ]
- [ \*mini min-value ]
- [ \*maxi max-value ]
- [ \*values type ]
- [ \*transparency background-color ]
- [ \*draw\_limits\_between\_elsets ]

```
[ *elsets_start_with prefix ]
```

[ \*latex orientation ]

\*list\_var specifies Gauss point variables to be plotted.

**\*output** is used to specify the gif output file name.

\*P0, P1, P2 specify points describing the cut plane: image will be plotted for all  $P = P0 + \alpha P0P1 + \beta P0P2$  with  $(\alpha, \beta) \in [0, 1]^2$ .

- \*ortho is used to impose an orthogonal (P0,P1,P2) base. If value 1 is given a new P2 point will be searched as the projection of the orginal P2 point on a direction normal to P0P1. If value 2 is given, the same process if done for P1 point respecting P0P2 direction.
- **\*step** is used to impose the spatial pitch of generated image pixels.
- \*dimx, dimy are used to imposed the number of pixel used in POP1 and POP2 direction (length and height of the gif image). If only one paramter is set, the other will be computed as to respect a spatial aspect ratio. Note that at least one of the commands step dimx dimy must be specified.
- \*color\_map specifies the color map that will be employed, choice are one of: default, bone, broken, BW, BW2, cool, copper, hot, HSV, inv\_default, inverse, jet, Low\_white, pink, prism, PS-default, White\_to\_dark (same as those in the Zmaster interface).
- \*min\_max is used to specify how the scalebar limits are determined and can be manual or auto. Default behaviour is auto. If manual mode is selected, user has to provide min and max values thanks to the \*mini and \*maxi keywords.
- \*values is used to specify if the values are taken as closest Gauss point (integration\_point, which is the default behavior), as the mean of the element Gauss point values (average\_in\_element) or as the values of the continuous field which has been extrapolated to the nodes from the Gauss points values (at the global scale, not element-wise) and then interpolated (interp).
- \*transparency is used to specify the background color in order to produce transparent pixels when no element are found. It can be white or black.
- \*draw\_limits\_between\_elsets print limits between elsets with background color (useful to visualize grain boundaries inside polycrystals).
- **\*elsets\_start\_with** is used specify what is the prefix of the involved elsets to print limits between. By defaults, all elsets are involved.
- **\*latex** requires portrait or landscape modifier. This command will generate latex and pdf files containing generated images with color bars, if a valid imagemagick and latex environment is present.

```
**process clip_image
*list_var sig23
*output cuts
*P0 (-1. -1. 0.)
*P1 (41. 0. 4.)
*P2 (0. 41. 4.)
*dimx 1000
*ortho 1
*latex portrait
```

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process coordinates

## \*\*process coordinates

## **Description**:

This post-processing is used to produce coordinates (X, Y, Z) as field variables; they can then be used by other post-processings. A "node id" field is also produced (resp. "element id" if the process is applied on an integ file).

## Syntax:

\*\*process coordinates

[ \*prefix prefix ]

Output fields are named X, Y, Z and id, optionnaly prefixed by prefix.

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process format

## \*\*process format

## **Description**:

This post processor is used to produce an "export file" containing the specified variables for all the maps being run. The output file is a formatted ASCII file. The variables should exist in the file specified by the **\*\*file** command (see page 4.26). For instance, the command **\*list\_var U1** should not be given with **\*\*file integ**. Note: this post processor also exists for **\*\*\*local\_post\_processing**, with similar options (see page 4.50).

## Syntax:

\*\*process format
\*file file
\*list\_var name1 ... nameN
[ \*precision digits ]
[ \*optimizer ]
[ \*blank\_line ]
[ \*write\_nodal\_coordinates coord | coord0 | none ]
[ \*write\_gp\_coordinates]

where file denotes the name of the output file, and name1, ..., nameN is a list of scalar variable names to output. For each map, the values at all locations (nodes, integration points) are stored. The option **\*optimizer** inhibits the writing of the output lines containing the character string **#** === time followed by the time associated with the current map, and the character string listing all variables. The option **\*precision** determines the number of digits with which the results will be written (default is 10). The option **\*blank\_line** writes an empty line between each output map. Some plotting programs, for instance gnuplot, may need this.

For variables associated to nodes, it is possible to write out the nodal coordinates as well by using the **\*write\_nodal\_coordinates** option. This options takes as argument either **coord** to indicate that the *current* coordinates should be written, or **coord**0 to indicate that the *initial* coordinates should be written. The default value **none** indicates that the nodal coordinates should not be written. Note: currently this option does not exist for the **\*\*\*local\_post\_processing \*\*format** version of this command.

For integrated variables, it is also possible to write out the Gauss point coordinates, by specifying the **\*write\_gp\_coordinates** option.

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process hot\_spot

## \*\*process hot\_spot

## **Description**:

This post processor is used to find locations which are local maxima or mimima of a given variable, and within a specified range, but isolated from other peaks by a given radius. This post computation should be useful for automating fatigue or critical stress analysis when meshes and loading conditions are often changing (see for example the usage with cycle\_projection on page 4.38).

## Syntax:

\*\*process hot\_spot
 \*format fmt-string
 \*function FUNCTION
 \*max\_val max
 \*min\_val min
 \*minimum
 \*minimum
 \*num fname
 \*radius rad
 \*screen\_output
 \*variable name
 \*write\_nodes fname

The following summarizes the different commands and their parameters:

- \*format is used to format the screen output for aesthetic value to the user. The command reads the characters remaining on the same line and uses that as a printf style formatting.
- **\*function** applies a FUNCTION to the selected field values before evaluating the hot spot condition (remember the semicolon after the function declaration).
- \*max\_val specifies a real value for the maximum values to be considered. This can be used to truncate the analysis to regions of reasonable scale.
- \*min\_val specifies a real value for the minimum values to be considered. An example use is
  when regions of a structure do not have a fatigue life prediction (and therefore values of
  zero), but we are interested in the minimum locations with values over 10 for example.
- \*minimum is a switch to indicate that we want minimum values as the hot spot. The default is to search for maximum absolute values.
- \*num specifies an integer for the number of values to report.

**\*radius** specifies a real value for the spatial radius separating the hot spot locations.

- **\*variable** specifies the scalar variable of interest. This can of course be a derived variable coming from previous post processing operations.
- \*write\_nodes is used to give a file name where a Zmesh compatible node set will be listed in order of the *num* hot spot points.

#### Note:

This command currently applies to nodal data only.

#### Example:

The following example is from the test case hot\_spot.inp in test/Post\_test/INP.

```
***global_post_processing
**process hot_spot
*radius 5.
*variable X
*num 2
*function -1.0*x;
*screen_output
```

Because we asked for  $*screen_output$  the following summary is printed to the standard out (and .msgp file). At the first timestep (t=0) the field is uniform so the first 2 nodes respecting the zone rule are printed. In the following the field is just scaled so the location results are the same. That would of course not be the case in a general FEA analysis.

```
+++++ HOTSPOT ANALYSIS X +++++
Time:0.000000
  1
                -2.00e-01
  7
                -2.00e-01
running : 25 %
+++++ HOTSPOT ANALYSIS X +++++
Time:1.000000
  152
                -2.80e+00
  309
                -2.15e+00
running : 50 %
+++++ HOTSPOT ANALYSIS X +++++
Time:2.000000
  152
                -2.10e+00
  309
               -1.61e+00
running : 75 %
+++++ HOTSPOT ANALYSIS X +++++
Time:3.000000
  152
                -1.40e+00
  309
               -1.08e+00
```

## \*\*process max

## **Description**:

This post-processor gives the maximal value of the specified variables for a group of elements (elset).

## Syntax:

\*\*process max

\*list\_var name1 ... nameN

name1, ... nameN are scalar variable names.

```
**process max
*list_var sig11 sig22
```

\*\*\*\*post\_processing
\*\*\*global\_post\_processing
\*\*process min

## \*\*process min

### **Description**:

This post-processor gives the minimum value of the specified variables for a group of elements (elset).

## Syntax:

\*\*process min

\*list\_var name1 ... nameN

name1, ... nameN are scalar variable names.

## Example:

\*\*process mises
 \*var sig
\*\*process min
 \*list\_var sigmises

## \*\*process node\_extrapolation

## **Description**:

This post-computation extrapolates integration point data to either the nodes (CTNOD format), or to the element nodes (CTELE).

## Syntax:

```
**process node_extrapolation
    *list_var name1 ... nameN
[ *output_var o-name1 ... o-nameN ]
```

name1, ..., nameN are the scalar variable names to treat.

The input data is read in the results files *problem.integ*. The calculated results are stored in the file *problem.ctnodp*.

These output variables are named gpname1, ... gpnameN, or o-name1 ... o-nameN if they were specified.

```
% Generates the variable gpsig11 at nodes
% from the sig11 variable in the .integ file
**file integ
**process node_extrapolation
    *list_var sig11
```

\*\*\*\*post\_processing
\*\*\*global\_post\_processing
\*\*process node\_interpola

## \*\*process node\_interpolation

### **Description**:

This post-computation interpolates nodal variables at the Gauss points.

## Syntax:

\*\*process node\_interpolation
 \*list\_var name1 ... nameN
[ \*output\_var o-name1 ... o-nameN ]

name1, ... , nameN are the scalar variable names to treat.

The input data is read in the results files *problem*.node or *problem*.ctnod (\*\*file node). The calculated results are stored in the file *problem*.integp.

These variables are named gpname1, ... gpnameN.

```
% provides gptemperature at the Gauss points
% from the temperature in .node file
**file node
**process node_interpolation
    *list_var temperature
```

## \*\*\*post\_processing \*\*\*global\_post\_processing \*\*process relocalised\_post

## \*\*process relocalised\_post

### **Description**:

This post computation interpolates the specified DOFs at specified coordinates.

#### Syntax:

\*\*process relocalised\_post
 \*variables list-of-variables
[ \*station vector ]
[ \*station\_file infile ]
[ \*output\_file outfile ]
[ \*tolerance value ]
[ \*locator\_type locator ]

\*variables is the list of DOFs to output.

- \*station is used if a single point is selected ; use \*station\_file if more than one point is required.
- \*output\_file specifies in which file output should be given. If it is omitted, results are printed on screen. If the filename ends with .vtk, the output file will be in vtk format; otherwise, standard Ascii file format suited to gnuplot is used.
- **\*tolerance** is passed to the element locator, to allow slightly out-of-elements point localization. Default is 0.
- \*locator\_type is the localization method. It defaults to bb\_tree (currently the optimal method available in Z-set).

```
****post_processing
***global_post_processing
**file node
% **at 1. 30. 365. 3650. 36500.
**output_number 0-999
**process relocalised_post
*station_file GPS_stations.dat
*variables U1 U2 U3
```

\*\*\*\*post\_processing
\*\*\*global\_post\_processing
\*\*process relocalised\_post

```
*output_file GPS-out.vtk
*tolerance 1. % in distance unit
****return
```

where the station file looks like (note that for convenience, all text after the 3rd column is ignored):

# X	Y	Z	site
-1.129764e+06	6.229049e+06	7.156934e+05	ARAU
-1.155688e+06	6.239179e+06	5.715508e+05	BABH
-1.834799e+06	6.058418e+06	-7.202233e+05	BAKO
-9.878078e+05	6.183675e+06	1.173047e+06	BANH
#[etc]			

## \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process static\_torsor

## \*\*process static\_torsor

## **Description**:

This post compute the static torsor equivalent to the sum of every nodal forces over a nset for each solution step. The equivalent static torsor is defined as:

$$\left\{ \begin{array}{l} \mathcal{T} \end{array} \right\}_{O} = \left\{ \begin{array}{l} \sum_{i \in \text{nset}} \vec{F_{i}} \\ \sum_{i \in \text{nset}} \vec{OO_{i}} \land \vec{F_{i}} \end{array} \right\}_{O}$$

## Syntax:

\*\*process average

\*point vector

\*file file

where vector is the position where the momentum is computed (O). The output gives the coords of point O, the 3 components of the resultant and the 3 components of the momentum for each time step. The output file is named *file*.

## Example:

\*\*\*global\_post\_processing
\*\*nset top
\*\*file node
\*\*process static\_torsor
\*point (1. 0.5 0.5)
\*file torsor.dat

## \*\*process surface\_normals

### **Description**:

This post processor computes the outer-pointing normals at nodes in the specified nset, and generates associated nodal fields. It currently works in 3D only.

Defining a normal direction at a node on a ridge is naturally ambiguous. Each node receives a contribution of its attached elements, *if all nodes of the elements are in the specified nset*. Thus, on ridges the normal is averaged over adjacent elements. Restricting the computation to a smooth subset of the surface will usually generate the desired field.

### Syntax:

\*\*surface\_normals
[ \*prefix prefix ]

prefix is used to name the output variable. Its default value is N so that normal vector components are N1, N2 and N3.

```
% This post processing calculates outer-pointing
% heat flux from a thermal calculation
****post_processing
 ***global_post_processing
  **file node
  **nset interior
  **process surface_normals
   *prefix N
    % Can be visualized in paraview with this calculator:
    % N1*iHat+N2*jHat+N3*kHat
 ***local_post_processing
  **file node
  **nset interior
  **process function
   *expression q1*N1 + q2*N2 + q3*N3 ;
   *output normalflux
***return
```

\*\*\*\*post\_processing
\*\*\*global\_post\_processing
\*\*process volume

## \*\*process volume

#### **Description**:

Calculate the volume of a group of elements (elset).

### Syntax:

\*\*process volume

## Example:

```
****post_processing
    ***precision 3
    ***data_source mesh_only
    **format Z7
    **open mymesh.geof
    **maps 1.
    ***global_post_processing
    **file integ
    %
    % volume of the whole structure
    %
     **elset ALL_ELEMENT
     **process volume
    %
    % volume of a part
    %
     **elset PART1
     **process volume
```

#### \*\*\*\*return

# \*\*\*\*post\_processing \*\*\*global\_post\_processing \*\*process volume\_above

### \*\*process volume\_above

#### **Description**:

The volume\_above processor is used to calculate the volume of elements of the active element set for which the value of the specified variable is greater than a criterion fixed by the used.

The result can be normalized to find the relative percentage of volume meeting the given criterion.

#### Syntax:

\*\*process volume\_above
\*var name
\*threshold value
\*normalize

where *name* is the name of a scalar variable. *value* is a real (floating point) value indicating the criterion to consider. With the option **normalize** the given result corresponds to the total volume fraction of elements meeting the criterion.

#### Example:

```
**process volume_above
 *var epcum
 *threshold 1.
 *normalize
```

#### \*\*process weibull

#### **Description**:

This post computation provides the means for doing a Weibull analysis on a structure. Two modes of operation are available:

#### eigenstress

$$\sigma' = \left(\frac{\sigma_1^M - \sigma_0}{\sigma_u}\right)^m$$

independent

$$\sigma' = \left(\frac{\sigma_1^M - \sigma_0}{\sigma_u}\right)^m + \left(\frac{\sigma_2^M - \sigma_0}{\sigma_u}\right)^m + \left(\frac{\sigma_3^M - \sigma_0}{\sigma_u}\right)^m$$

The Weibull stress is then defined as:

$$\sigma_W = \left(\frac{\sigma_u}{V_0} \int_V \sigma'^m \, dV\right)^{1/m}$$

where  $\sigma_1^M$ ,  $\sigma_2^M$  and  $\sigma_3^M$  are the post-computation results from a local fmax applied successively to the three principal stresses  $\sigma_1$ ,  $\sigma_2$  and  $\sigma_3$ , in descending order. These sub-posts are run automatically by the weibull processor.

 $V_0, \sigma_u, \sigma_0$  and m are material parameters.

The probability of failure  $P_r$  is is given by:

$$P_r = 1 - \exp\left(-\left(\frac{\sigma_W}{\sigma_u}\right)^m\right)$$

In addition to the output of  $\sigma_W$  and  $P_r$ . the values of  $\sigma'$  are stored at each Gauss point under a name constructed by adding \_wb to the variable name.

#### Syntax:

```
**process weibull
 *var name
 *mode eigenstress | independant
 *coefmin value1
 *coefmax value2
 *file namef
```

If the user has only specified a single map (with output\_number), the history is reconstructed from the values read after the options \*coefmin and \*coefmax.  $\sigma_1^M$  varies linearly over 100 maps of value1\* $\sigma_1$  to value2\* $\sigma_1$  (where  $\sigma_1$  is calculated at the specified map).

With the option **\*file**, the history is read in the file **namef**, of the form:

 0.01
 10

 0.02
 15

 0.03
 20

 0.04
 40

 0.05
 60

 ...

where the first column represents the time and the second the value of  $\sigma_1$ .

These two methods presented for the mode  $\verb"eigenstress"$  extend to the mode  $\verb"independant"$  as well.

#### Example:

```
**output_number 10
**process weibull
 *var sig
 *mode independent
 *coefmin 0.5
 *coefmax 2.5
% material file
**process weibull
 V0 10.
 m 20.
 sigma_u 1200.
 sigma_0 0.
```

## Chapter 5

## Reference

### Functions

#### **Description**:

The functions have been generalized in code versions greater than 7.1 to have a natural equation like form, and apply more widely throughout the program. In addition to coefficient definitions, the functions may now be used to define load waveforms, apply to the optimizer, etc. More applications will be added as well.

The functions follow a C-like syntax, and may be defined in terms of pre-defined functions (sin, log, etc) and named parameters of the problem. What parameters are allowed and available depends on application.

#### Syntax:

Objects which are functions may now be entered using a C-like format. This *must* include a semi-colon at the end of the function definition to terminate the reading of the function. Many operators exist, which have the same meaning and order of selection as in C. Coefficients which are functions still require the keyword **function** to follow the coefficient name, but are otherwise generally given in terms of the relevant parameters.

The following operators are defined in functions:

:= == >= <= + - \* / ^ > <

as are the following functions:

triangle floor asinh acosh atanh asin acos atan cosh sinh tanh ceil sqrt neg abs sqr sin cos tan exp ln Hx H urandom

Pre-defined functions may take complex arguments of other expressions or functions.

triangle function used to make a sawtooth waveform with period 1 of the parameter.

floor largest integral value not greater than x.

*H* Heavyside function. Equal to 1 for x > 0 and 0 otherwise.

Hx Heavyside of its parameter times its parameter. Equal to x for x > 0 and 0 otherwise.

urandom(a,b) Uniform random distribution that returns values between a and b

#### Example:

```
***function load_tab sin(6.28319*time);
***elasticity
young function 2.e6-100.*temperature -2.*temperature^2.;
poisson 0.3
```

## Degrees of Freedom (DOF)

#### **Description**:

The degree of freedom (DOFs) types in Z-set each have a particular name used for their identification in various parts of the code. These applications may be in specifying the boundary conditions (\*\*impose\_nodal\_dof takes a DOF name for example), specifying output curves, or use within the post-processor.

The names currently active in the code are summarized in the following table:

CODE	DESCRIPTION
U1	displacement along the axis 1
U2	displacement along the axis 2
U3	displacement along the axis 3
EZ	Deformation along the axis 3 for plane stress
R1	Rotation about the axis 1
R2	Rotation about the axis 2
R3	Rotation about the axis 3
W1	Micro-polar rotation about the axis 1
W2	Micro-polar rotation about the axis 2
W3	Micro-polar rotation about the axis 3
PR	Pressure
TP	Temperature

### **Element Geometries**

#### **Description**:

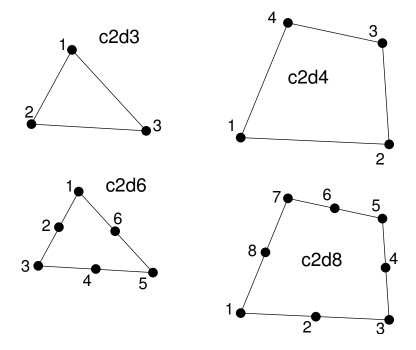
This section defines the ordering of node and Gauss point numbering for the different element geometries. The following figures display the element information with numbering corresponding to the order of definition in the .geof file.

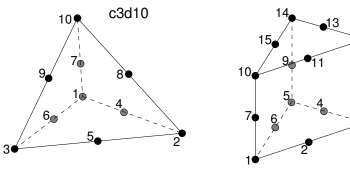
The following element geometries are available:

- 12d2, 12d2r, 13d2, 13d2r line elements. These are used for truss or spring elements with two nodes. There is only a force acting between the nodes.
- 12d1, 13d1 one node spring or dashpot elements. These oppose motion from their originl location.
- rve1d, rve2d, rve3d representitive volume element (RVE) to be used for material simulation. There are no nodes. and only one "integration" point.
- deb2 2D debonding element geometry.
- c2d3r, c2d3, c2d4r, c2d4, c2d6r, c2d6, c2d8r, c2d8 2D continuum elements. "r" denotes reduced integration.
- cax3r, cax3, cax4r, cax6r, cax6r, cax8r, cax8 axisymmetric 2D continuum elements.
- c3d4, c3d4r, c3d10r, c3d10, c3d6r, c3d6, c3d8r, c3d8, c3d20r c3d20, c3d15r, c3d15 3D continuum elements.
- sax3, sax3r, sax3rr, sax2, sax2r, sax2rr axisymmetric shells.

s3d3r, s3d3, s3d4r, s3d4, s3d6r, s3d6, s3d8r, s3d8 3D shells.

Node definitions for different element geometries are given on the following page. Note that all other information about the element chosen (formulation, material, real constants) are specified in terms of element sets in the *problem.inp* file.



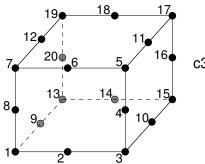




12

8•

3



c3d20

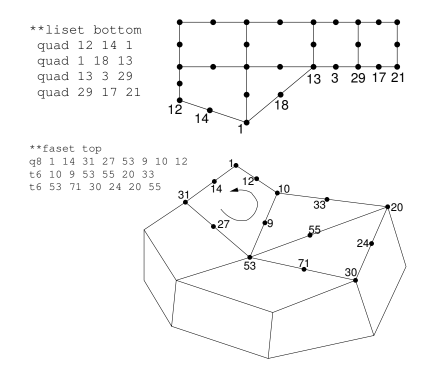
## **Boundary sets**

#### **Description**:

The boundary sets (faces in 3D or lines in 2D) must be ordered properly such that the normals may be calculated, positions interpolated, and surface integrals evaluated over them. Incorrect definitions of these sets are often responsible for boundary condition problems.

CODE	DESCRIPTION
line	linear $(2 \text{ node})$ line
quad	quadratic (3 node) line
t3	linear 3 node triangle face
t6	quadratic 6 node triangle face
q4	linear 4 node quadrilateral face
q8	quadratic 8 node quadrilateral face

The following boundary types are available:

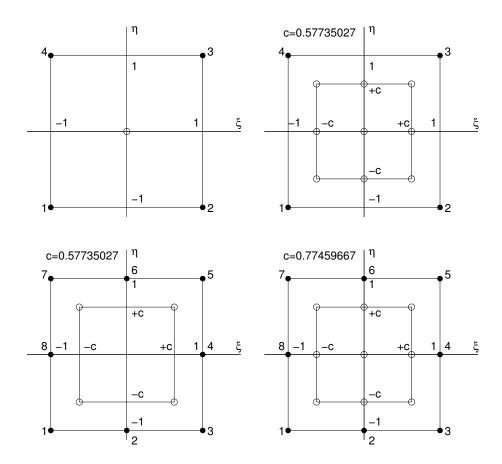


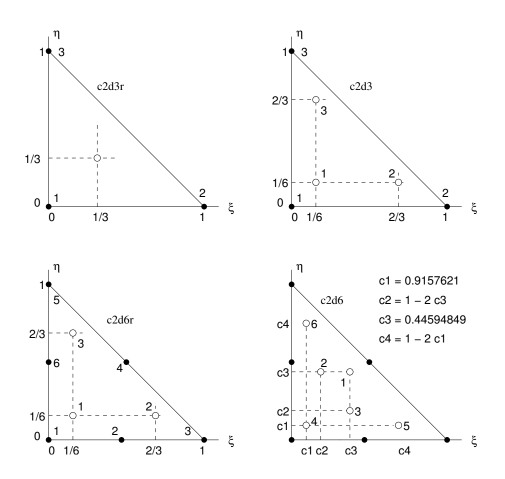
### **Element Integration**

#### **Description**:

This section defines the positions and numbering of element Gauss points used for spatial integration. These points are the *only* places where material values (e.g. stress and strain) exist. If you need accurate magnitudes of these variables, it is highly advised to use a **\*\*test** output (see page 3.151) to output them, or use the **\*\*value\_at\_integration** output for the structure (page 3.148).

Gauss points are defined in a local element coordinate system which is mapped to the global coordinates (real element geometry) using the element shape functions. The element is defined in the local coordinates by -1 to 1 on the  $\xi$  and  $\eta$  element axis.





### Structure of *problem*.geof

The structure of the ASCII file describing problem geometries is summarized below:

```
***geometry
**node
     dim
                            number of nodes, space dimension
n
      y1
x1
            [z1]
...
            [zn]
xn
      yn
**element
                            number of elements
j
                            element number, type, list of node numbers
1
   type1 n11 .....n1k
j typej nj1 .... njk
***group
                            name of a node set
**nset name
n1 n2 \dots nk
                            list of the k nodes in the node set
**elset name
                            name of an element set
n1 n2 ... nk
                            list of the k elements in the element set
**faset name
                            surface (3D face) group name
type1 n11 ... n1k
                            type of face, ordered list of nodes in the face
...
typej nj1 ... njk
**liset name
                            surface (linear) group name
type1 n11 ... n1k
                            type of line, ordered list of nodes in the liset
...
typej nj1 ... njk
***return
```

The position of the nodes must be given in Cartesian coordinates for 2D plane and 3D problems, or in cylindrical coordinates (r,z) for 2D axisymmetric problems.

The groups of nodes, faces, elements, and element lines segments must come after the

node and element definitions. These groups are defined between the keywords **\*\*\*group** and **\*\*\*return**.

## Z-set output formats

Z-set has two output formats. The default output format is Z7. A new output format has been implemented since Z8.4 version, to allow remeshing.

#### Z7 output format

problem.ut The description of the output files problem.node, problem.integ, problem.ctnod, and problem.ctele is given in the file problem.ut.

- After **\*\*node** the names of the nodal variables and parameters stored in *problem.***node** are given.
- After **\*\*integ** the names of integration (material) variables stored in *problem.integ*, *problem.ctnod*, and *problem.ctele* are given.
- **\*\*meshfile** gives the name of the file *problem*.geof used for the problem (presently without importance).
- **\*\*element** is not presently used.

Following the demarcated sections described above, a list of the output details is given. Each line represents the state of the calculation for one output. These lines contain the output number, the cycle number, the sequence number, increment, and corresponding time, in that order.

problem.node The file problem.node contains the nodal output variables. These variables consist of the nodal degrees of freedom (such as displacements, temperature, etc), the associated reactions to the nodal DOFs (forces, heat flux), and possibly the problem nodal parameters if the **\*\*save\_parameter** option was given after **\*\*\*output**. The stored variables will be the complete set of variables at each node, regardless if certain parameters do not exist at every node in the problem. For variables which do not exist at a given node, a zero value will be stored.

The *problem*.node file has the following structure:

 $\begin{array}{ll} [d_1, \dots, d_N]_1 \dots [d_1, \dots, d_N]_{n_d} & 1^{st} \text{ output} \\ [v_1, \dots, d_N]_1 \dots [v_1, \dots, d_N]_{n_d} & \\ [d_1, \dots, d_N]_1 \dots [d_1, \dots, d_N]_{n_d} & 2^{nd} \text{ output} \\ 1ere \ variable & variable & n_d \\ [\dots] & \end{array}$ 

where  $n_d$  is the number of DOFs given by **\*\*node** in the *problem.ut* file, and N is the number of nodes.

The *problem*.node file may be written in C with the following code segment:

```
for(card=1;card <= stored_cards; card++){
  for(variable=1; variable <= nodal_variables_stored;
  variable++) {
    for(node=1;node <= number_of_nodes_in_mesh ; node++) {
        fwrite(&composante[node][variable][card],4,1,fp);
    }
  }
}</pre>
```

problem.integ The file problem.integ contains the integration point values issued from the material behavior laws (see \*\*value\_at\_integration user \*\*\*output). In the event that certain variables do not exist at all integration points in the problem, a zero value will be stored in its place. Thus the number of variables is always the union of the different material variables in the problem.

Each record in the *problem*.integ file has the following format:

$v1_{pg1el1}, v1_{pg2el1} \ldots v1_{pgnel1}$	$1^{st}$ variable for element 1
$v2_{pg1el1}, v2_{pg2el1} \ldots v2_{pgnel1}$	$2^{dn}$ variable for element 1
$vnv_{pg1el1}, vnv_{pg2el1} \dots vnv_{pgnel1}$	variable nv for element 1
$v1_{pg1el2}, v1_{pg2el2} \ldots v1_{pgnel2}$	$1^{st}$ variable for element 2
$v2_{pg1el2}, v2_{pg2el2} \ldots v2_{pgnel2}$	$2^{nd}$ variable for element 2
$vnv_{pg1el2}, vnv_{pg2el2} \dots vnv_{pgnel2}$	variable nv for element 2
$v1_{pg1elNE}, v1_{pg2elNE} \ldots v1_{pgnelNE}$	$1^{st}$ variable for element NE
$v2_{pg1elNE}, v2_{pg2elNE} \ldots v2_{pgnelNE}$	$2^{nd}$ variable for element NE
$vnv_{pg1elNE}, vnv_{pg2elNE} \dots vnv_{pgnelNE}$	variable nv for element NE

where NE is the number of elements in the structure, and nv the number of values stored per integration point (given by **\*\*integ** in the file *problem.ut*.

Elements generally do not have the same number of integration points. Due to this, the storage of different element types in a problem will generate element data structures of different size within the same record.

The file *problem*.integ may be generated using the following C code:

```
for(card=1;card <= number_of_outputs; card++){
  for(ele=1; ele <= number_of_elements ; ele++) {
    for(var=1; var<= number_of_stored_variables; var++) {
      for(point=1;point <=
         number_of_integration_points_per_element[ele] ;point++) {
         fwrite(&component[point][var][ele][card],4,1,fp);
      }
    }
  }
}</pre>
```

problem.ctnod The problem.ctnod file contains the integration point variables (material variables) extrapolated to the nodes and will be generated when the \*\*contour option is selected under \*\*\*output. This file follows the storage format used for the problem.node files.

The structure of each record in the *problem*.ctnod file is the following:

 $v1_1, v1_2 \ldots v1_N$  variable 1

Z-set — Non-linear material & structure analysis suite

```
v2_1, v2_2 ... v2_N variable 2
...
vnv_1, vnv_2 ... vnv_N variable nv
where N is the number of nodes and nv the number of variables indicated by **integ
in the problem.ut file. A problem.ctnod may be generated using the following C code:
for(card=1;card <= nombre_de_carte_stockees; card++){
  for(var=1: var <= nombre_de_variables_stockees : var+t) {</pre>
```

```
for(var=1; var <= nombre_de_variables_stockees ; var++) {
   for(node=1;node <= nombre_de_noeuds_du_maillage ; node++) {
      fwrite(&composante[node][var][card],4,1,fp);
   }
}</pre>
```

problem.ctele The file problem.ctele contains the integration variables at nodes. The file duplicates the problem.ctnod data but uses on an element to element extrapolation basis. Element based storage retains the discontinuous fields between elements with different materials. The file will be generated by using the \*\*contour\_ele option under \*\*\*output. Output records in the problem.ctele file each have the following format:

$v1_{no1el1}, v1_{no2el1} \ldots v1_{nonel1}$	$1^{ere}$ variable for element 1
$v1_{no1el2}, v1_{no2el2} \ldots v1_{nonel2}$	$1^{ere}$ variable for element 2
$v1_{no1elNE}, v1_{no2elNE} \ldots v1_{nonelNE}$	$1^{ere}$ variable for element NE
$v2_{no1el1}, v2_{no2el1} \ldots v2_{nonel1}$	$2^{nde}$ variable for element 1
$v2_{no1el2}, v2_{no2el2} \ldots v2_{nonel2}$	$2^{nde}$ variable for element 2
$v2_{no1elNE}, v2_{no2elNE} \ldots v2_{nonelNE}$	$2^{nde}$ variable for element NE
$vnv_{no1el1}, vnv_{no2el1} \dots vnv_{nonel1}$	variable nv for element 1
$vnv_{no1el2}, vnv_{no2el2} \dots vnv_{nonel2}$	variable nv for element 2
$vnv_{no1elNE}, vnv_{no2elNE} \dots vnv_{nonelNE}$	variable nv for element NE

where NE is the number of the elements in the structure, and nv the number of values stored per integration point (as given by the **\*\*integ** section in the file *problem.ut*).

Different elements do not have the same number of nodes. The storage for different elements within the problem may therefore have different data structure sizes within the same record.

The file *problem*.ctele may be generated using the following C code:

```
for(card=1;card <= nombre_de_carte_sorties; card++){
  for(var=1;
  var <= nombre_de_variables_stockees_au_points_d_integration;
  var++) {
    for(element=1; element <= nombre_d_elements; element++) {
        for(node=1;node <= nombre_de_noeuds_de_l_element[element];
    }
}</pre>
```

```
node++) {
    fwrite(&composante[node][element][var][card],4,1,fp);
    }
  }
}
```

- *problem.eigen\_info* If the calculation is an eigen value problem, the resonant frequencies and associated energies for each mode are stored in the file *problem.eigen\_info*. This file is an ASCII formatted text file.
- problem.eigen When an eigen value calculation is run, the resonant vectors or modal displacements are stored in the file problem.eigen. The structure of the binary file is ordered mode number, frequency of the mode, and the energy associated to the mode. Thus we have:

$U1_{1},$	$U1_2$	 $U1_N$	displacement U1
$U2_1$ ,	$U2_2$	 $U2_N$	displacement U2
$[U3_1,$	$U3_2$	 $U3_N$	displacement U3 ]

end of the storage of the first mode.

The file *problem*.eigen may be generated using the following C code:

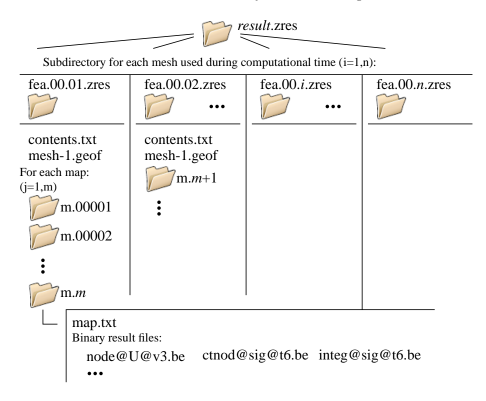
```
for(mode=1;mode <= nombre_de_modes_extraits; mode++){
   fwrite(&mode,4,1,fp);
   fwrite(&frequence_du_mode[mode],4,1,fp);
   fwrite(&energie_du_mode[mode],4,1,fp);
   for(dime=1; dime <= dimension_de_l_espace ; dime++) {
      for(node=1;node <= nombre_de_noeuds_du_maillage ;node++) {
        fwrite(&composante[dime][node][mode],4,1,fp);
      }
   }
}</pre>
```

#### Z8 output format

#### **Description**:

The Z8 output format has been implemented since Z8.4 version, to allow remeshing. While the default Z7 format uses *results.integ*, *results.node*, *results.ctnod* ... result files, the new Z8 format uses a tree allowing to read results for which mesh changes from one map to another.

The figure below shows the structure of the file system containing result and mesh files.



#### Syntax:

User can activate the new Z8 format by adding in the .inp file the following commands: \*\*\*\*calcul

```
...
***global_parameter
Solver.OutputFormat Z8
Zmaster.OutputFormat Z8
...
```

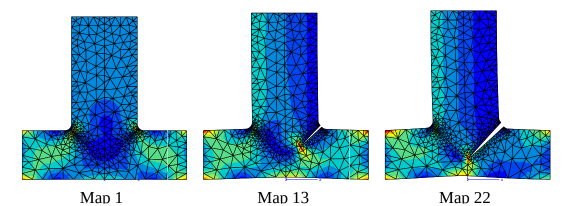
```
****return
```

The Z8 format file structure is organized as follows:

- one main directory named -here- result.zres (implying that the .inp file is result.inp, and the execution command "Zrun result");
- some sub-directories respectively named fea.00.01.zres, fea.00.02.zres ... created at each remeshing. Each *i*<sup>th</sup> remeshing generates a corresponding fea.00.*i*.zres sub-directory. Each sub-directory contains:
  - an ASCII file named contents.txt that describes the name, the type (scalar, tensor ...), and the location (at nodes, at integration points, extrapolated at averaged nodes ...) of the physical quantities stored in result files;
  - the current mesh file *i.e.* an ASCII file with a .geof extension;
  - a list of sub-directories named m.00001, m.00002 ..., corresponding to the list of output maps (computational times for wich user wants results to be written), and containing:
    - \* an ASCII file named map.txt that describes the current output map (cycle number, sequence number, increment number, time value ...);
    - \* binary files containing the results computed for each physical quantity to be written.

This new file system is directly machine-readable using **Zmaster**. User may from now on graphically analyze computational results for which mesh changes from one map to another.

The figure below shows a crack propagation during a computation implying remeshing.



## **Environment Variables**

The environment variables are used to personalize the Z-set program for individual users, and maintain a self-contained project structure. The function extends as well to manage multi-architecture sites, thereby allowing the transparent use of the program over a diverse network. The currently used environment variables are summarized as:

CODE	DESCRIPTION
Z7PATH	path to the head Z-set's directory structure
Z7_MAX_NB_DOF	number of DOFs above which disk storage is used
Z7_TMP_DIR	path for the temporary files if needed
Z7_LICENSE	path to a license file if not stored in \$Z7PATH/lib/Zebulon.License

Z7PATH variable which indicates the location of the Z-set distribution. This variable allows easy switching to different distributions. It also is a cause of much difficulty with users if not configured correctly.

cd /usr/local/Z8.3.6-Mar-15-2006 setenv Z7PATH 'pwd' source lib/Z7\_cshrc

- Z7\_MAX\_NB\_DOF This sets the maximum number of degrees of freedom before the global matrix is stored on disk. Because disk storage is not expensive, it is advisable to set this to a fairly low value (e.g. 5000). One can also use **\*\*\*file\_management** to comtrol the parameter.
- Z7\_TMP\_DIR directory where the temporary files are to be stored. Default is the problem directory. This can also be set with **\*\*\*file\_management**.
- Z7\_LICENSE fully qualified path to a license file. Extremely useful when there are multiple versions, or for running Zebulon directly off a CD-ROM where one can't put the license file in a read only lib dir.

#### Example:

In the C-shell, these variables are set using the **setenv** command. The variables may be directly in the user's .cshrc file, or in a separate file. For the later case, one may add the line to the .cshrc file: if (-f ~/lib/Z7\_vars) source ~/lib/Z7\_vars Examples of variable settings are given below:

% cat ~/lib/Z7\_vars
setenv Z7\_TMP\_DIR /home/disk/me/Z7
setenv Z7\_MAX\_NB\_DOF 5000
setenv Z7\_PRINTER\_COLOR\_A3 HPDESKJET

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