



MEDLoader - Reading and writing MED Files

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Agenda

MEDFile/MEDLoader Overview

Basic API

Advanced API

Families and Groups

Conclusion



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- Overview of the MEDLoader, fundamental differences with MEDCoupling model
- Basic API
 - Quick export to VTU (visualization)
 - Reading and writing a MED file
- Advanced API
 - Some insights into the MED file format
 - Advanced concepts
 - Groups
 - Profiles

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MEDLoader, from a memory model to a file

The tool to save / load what you do in memory with MEDCoupling

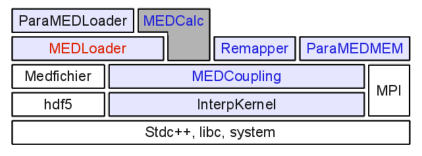


Figure 1: MEDLoader in the MEDCoupling distribution

MEDLoader, not only a "loader"

- Why do we need a loader/writer?
 - Communicate between processes / tools
 - Load / backup results
- Very unfortunate name
 - Can obviously read, but also write meshes and fields . . .
 - ... into a file with the MED type (".med" extension): "Modèle Echange de Données" (Data Exchange Model)
- Part of the MEDCoupling library, requires:
 - MED-file library (currently version 3.0.8) to compile and run
 - MED-file is the low level C/Fortran API to deal with MED files
 - HDF5 (prerequisite of MED-file library)
 - MPI if you're working in parallel
- You can use the core structures of MEDCoupling without MEDLoader
 - The reverse does not work
- Like before, most of it in C++, but wrapped in Python
 - Most of the algorithms are actually pure MEDCoupling core



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MEDLoader Services

- Read / write
- Groups manipulation
 - A collection of entities on the mesh (volumes, faces, points . . .)
- Geometric algorithms
 - convertAllToPoly()
 - unPolyze()
 - zipCoords()
 - duplicateNodes()
- Those services often have a MEDCoupling equivalent
- MFD file conversion tools
 - Between MED versions, from SAUV, ...



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Differences with MEDCoupling, more flexibility

- Warning: MEDFilexxx ≠ MEDCouplingxxx
 - MEDCoupling tries to rationalize the large flexibility provided by the MED-file format
 - Loosing some advanced functionalities, but more user-friendly
- Capabilities of MED-file that are "restricted" in MEDCoupling:
 - A mesh:
 - can have multiple dimensions
 - can have groups
 - A field:
 - can be defined only on a part of the mesh: "profiles"
 - can have more than one spatial discretization (ON_CELLS, ON_NODES,...)
 - A MEDFile field can have multiple time-steps (MEDCouplingFieldDouble = single step)
 - Solution: use multiple MEDCouplingFieldDouble
 - Solution: use MEDCouplingFieldOverTime
- Rule of thumb: try to do it with the MEDCoupling data model only

Differences with MEDCoupling, some constraints

- "Objects" need to have a name
 - Mesh and fields need to be named
 - setName() method
 - MED-file has a max length of 64 characters
- Cells need to be ordered by geometric type
 - E.g. if you mix TRIA3 and QUAD4 in your mesh (sometimes called a hybrid mesh)
 - Invoke sortCellsInMEDFileFrmt()
 - Try to keep cell types ordered if you can (will avoid save/load time)
 - Or convert everything to polyhedrons!
 - MEDCouplingUMesh::convertAllToPoly()
 - Only one cell type
 - Can be converted back with unPolyze()

MEDFile - Mesh Dimension

Up to 4 dimensions for a single MEDFileUMesh

- Start point = most upper level here 3D
- levelRelativeToMax = 0 (volumes)



■ levelRelativeToMax = -1 (faces)



■ levelRelativeToMax = -2 (edges)



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Basic API - Overview (1/2)

I need a quick look

- VTK/VTU export
 m = <some mesh/field I just created>
 m.writeVTK("/tmp/foo.vtu")
- Visualization with ParaView

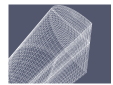


Figure 2: A vtu mesh

Basic API - Overview (2/2)

A step further – the basic MEDLoader API

- Directly reading/writing a mesh to a file on disk
 - WriteMesh() / ReadMesh()
- Directly reading/writing a field
 - WriteField() / ReadField()
 - WriteFieldUsingAlreadyWrittenMesh() if you have several fields on a single mesh
- All dealing with MEDCoupling objects (e.g. MEDCouplingUMesh, ...)
- Only static methods (i.e. no internal state). File reopened each time!



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Basic API - Reading a multi-dimensional mesh

Reading 3 levels of meshes (volumes, faces, vertex)

```
import medcoupling as mc
medFile = 'file.med'
meshName = 'mesh'
mesh3D = mc.ReadUMeshFromFile(medFile, meshName, 0)
mesh2D = mc.ReadUMeshFromFile(medFile, meshName, -1)
mesh1D = mc.ReadUMeshFromFile(medFile, meshName, -2)
```

- Works fine, but:
 - The 3 meshes have 3 independent coordinate arrays
 - Could be shared see later slide on mesh dimension ...

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Advanced API - Introduction



- Dealing with multiple time-steps
- Dealing with multiple mesh dimensions
 - Typically a volumic mesh (space dim == mesh dim == 3)
 - And some boundary conditions on the faces (mesh dim = 2)
 - The two mesh share the same nodes
- Dealing with mesh groups
 - A group is a named set of cells in a given mesh
 - Frequently used for boundary conditions assignment
- Dealing with partial support (rare case): profiles

Advanced API - Class diagramm

Helps navigate the advanced API

All names prefixed with % actually start with MEDFile (e.g. MEDFileData)

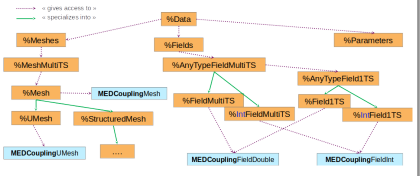


Figure 3: Links with the MEDCoupling data model

Advanced API – Important options

Writing a file

- Write options:
 - 2: force writing from scratch (an existing file will be reset)
 - 1: append mode (no corruption risk if file already there).
 - 0: overwrite mode: if the file and the MED object exist, they will be overwritten, otherwise write from scratch

Reading a field (MEDFileField1TS, MEDFileFieldMultiTS)

- Read everything in the file by default
- For large size fields:
 - Set the boolean loadAll to False in constructors
 - Use loadArrays() or loadArraysIfNecessary() to load data on demand
 - Use unloadArrays() Or unloadArraysWithoutDataLoss() to free memory



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Advanced API - Example

Example

```
medFile = "myWrittenFile.med"
coo = mc.DataArrayDouble([...])
mesh1D = <MEDCoupling object>
mesh1D.setCoords(coo)
mesh2D = <MEDCoupling object>
mesh2D.setCoords(coo)
mesh3D = <MEDCoupling object>
mesh3D.setCoords(coo)
mesh = mc.MEDFileUMesh.New()
mesh.setName("myMesh")
mesh.setMeshAtLevel(0, mesh3D)
mesh.setMeshAtLevel(-1, mesh2D)
mesh.setMeshAtLevel(-2, mesh1D)
mesh.write(medFile, 2)
```

Better than basic API

- All 3 meshes share the same coordinates array
- Saving space and write/load time
- Ensure consistency

Basic/Advanced API - Summary

Basic API is well suited for

- Meta information of a MED-file, without loading everything
 - E.g.: GetMeshNames, GetComponentsNamesOfField, GetFieldIterations
- Reading / Writing single instances of MEDCoupling objects
- Reading / Writing MEDCoupling meshes (without groups nor families)
- Reading / Writing MEDCouplingFieldDouble (without profiles)

You need the advanced API to

- Handle (part of) a MED-file in memory (e.g. to rewrite part of it only, a single time-step, . . .)
- Deal with families and groups
- Deal with field profiles



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Groups and families definition

Families

- Families do not need to be manipulated directly in normal usage !
- In a mesh cells are partitioned by families
- Each cell has a unique family ID (reverse not true)
- MEDLoader advanced API gives access to families
 - getFamilies(), getFamiliesArr(),...

Groups

- A group is a list of families
- MEDLoader advanced API gives access to groups
 - getGroups(),...

Families and Groups - Illustration



Figure 4: Families and groups example

Table 1: Families and cells associated to groups

Group A	Group B
Families 2 & 3	Families 7 & 3
Cells 0, 1, 4	Cells 2, 1

Recording a Group

Example

```
myCouplingmesh = ...
tabIdCells = mc.DataArrayInt()
tabIdCells.setName("meshGroup")
tabIdCells.setValues(...)
mfum = mc.MEDFileUMesh()
mfum.setName("Name")
mfum.setDescription("description")
mfum.setCoords(myCouplingmesh.getCoords())
mfum.setMeshAtLevel(0, myCouplingmesh)
mfum.setGroupsAtLevel(0, [tabIdCells])
```



Reading a Group

Example

```
mfum = mc.MEDFileUMesh(fname)
gpNames = mfum.getGroupsNames()
# mc.DataArryInt
myGroupArr = mfum.getGroupArr(0, gpNames[0])
# mc.MEDCouplingUMesh
myGroup = mfum.getGroup(0, gpNames[0])
Either get the result:
```

- as a DataArrayInt
- as a sub-mesh



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Conclusion

- Any question ?
- Let's get ready for the exercises!

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Advanced API – MED-file format (1/2)

What does a MED file look like?

- A specialization of the HDF5 standard
 - Take a look at http://www.hdfgroup.org/HDF5/
 - Focus on (potentially) big amount of data
 - Parallelism in mind
- Official documentation of the MED-file format available on-line at
 - \$\text{MEDFILE_ROOT_DIR}/\share/\doc/html/index.html (in French)}
- Reminder: MED-file denotes both
 - A file format on disk (I/O)
 - A comprehensive low level C library to read/write MED files
 - Now with a Python wrapping
 - Author: Eric FAYOLLE (EdF R&D)
 - MED-file existed before MEDCoupling!



Advanced API – MED-file format (2/2)

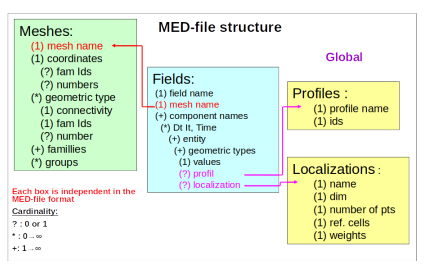


Figure 5: alt text