# TRUST ICoCo Tutorial V1.7.6

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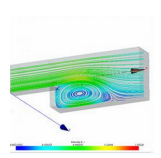
- 1 Introduction to code coupling and ICoCo
- 2 TRUST initialization
- Basic test case
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# Why code coupling?

## Code coupling ... what for ?

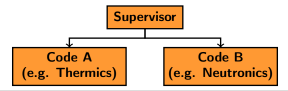
- Traditionnaly numerical simulation codes focus on a single physics
  - One code for thermics
  - Another code for mechanics,
  - Etc...
- Real life studies require the simulation of different physics
  - E.g. nuclear reactor simulations require a blend of: thermics, neutronics, mechanics.
- Solution? Code coupling!
  - Have different codes communicating one with another...
  - And having each of them dealing with its own area of expertise



# Code coupling

### First approach

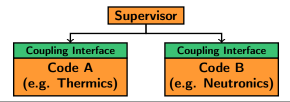
- An entity driving the complete computation is needed: the supervisor
  - Initializes code A and code B
    - Loop through code A and code B
    - Centralize the exchanges and conversions between A and B
    - Supervisor is often usually written from scratch as a C++ file, or a Python script
- In a dummy approach: supervisor needs to know both A and B API
  - o Becomes cumbersome if more than 2 codes to couple ...
  - Or if one supervisor is meant to run with different pairs of codes (code C, D, E
     ...)



# Code coupling

#### Better: common interface

- <u>Idea</u>: define a unique interface to which each code must comply
  - o Only an interface, i.e. a contract (=an API) that each code fulfills
  - Implementation (=plugging the wires) of this interface is tightly linked to the details of the code...
- Advantages:
  - All codes implementing the (unique) interface can be passed to the supervisor (almost) without modifying it! Versatility
- Cons:
  - Need to implement the interface for each code to be coupled



# **ICoCo**

## Interface for COde COupling

- ICoCo is such a coupling interface
- Stands for Interface for Code Coupling
- Written in C++
- Initially designed for simulation codes exhibiting an iterative time loop
- Presents a set of standard methods whose signature is fixed, with no implementation by default:
  - initializeTimeStep()
  - validateTimeStep()
  - abortTimeStep()
  - o ...
- Uses the notion of field for the exchange of data between codes
  - A field is a set of values supported by a mesh
  - A possible implementation: MEDCouplingFieldDouble, from SALOME's MEDCoupling library

# Further reading

## Meaningful documentation

- ICOCO presentation in TRUST documentation:
  - "An Interface for Code Coupling ICoCo v1.2":
    - \$ source /home/triou/env\_TRUST\_X.Y.Z.sh
    - \$ evince \$TRUST\_ROOT/doc/Kernel/ICoCo\_V1.2.pdf &
  - Ask the "APIProblem.pdf" note to trust@cea.fr

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# Initialisation of TRUST & ICoCo environment

- Source the TRUST environment:
  - \$ source /home/triou/env\_TRUST\_X.Y.Z.sh
- To know if the configuration is ok and where are the sources:
  - \$ echo \$TRUST\_ROOT
- Source the ICoCo environment:
  - \$ source \$TRUST\_ROOT/Outils/ICoCo/ICoCo\_src/env\_MEDICoCo.sh
- Test if ICoCo is compiled:
  - \$ 1s \$exec
- If you obtain "Is: cannot access \*\*/Outils/ICoCo/ICoCo\_src/ICoCo\_opt: No such file or directory", then compile ICoCo\_src:
  - \$ cd \$TRUST\_ROOT/Outils/ICoCo/ICoCo\_src
  - \$ baltik\_build\_configure -execute
  - \$ make optim debug
  - \$ source full\_env\_MEDICoCo.sh

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# First with TRUST:

#### Launch calculation with TRUST

- We create a first reference test case, launched with TRUST.
- Then we want to launch it with ICoCo and compare the results.
- Copy the following test case in your repository:
  - \$ mkdir ICoCo\_exercises
  - \$ cd ICoCo\_exercises
  - \$ trust -copy Vahl\_Davis\_hexa
- Rename the folder to sort your tests:
  - \$ mv Vahl\_Davis\_hexa Vahl\_Davis\_hexa\_trust
  - \$ cd Vahl\_Davis\_hexa\_trust
- Rename the data file to be consistent with the name of the repository:
  - \$ mv Vahl\_Davis\_hexa.data Vahl\_Davis\_hexa\_trust.data
- Launch calculation:
  - \$ trust Vahl\_Davis\_hexa\_trust
- You obtained a Vahl\_Davis\_hexa\_trust.lml file. We will use it in the following part.

# With ICoCo

## Adjusting the data file

- Now lets do the same with ICoCo:
  - \$ cd ..
- You are now in your folder "ICoCo\_exercises", create a new directory in order to launch your test case with ICoCo:
  - \$ trust -copy Vahl\_Davis\_hexa
  - \$ mv Vahl\_Davis\_hexa Vahl\_Davis\_hexa\_ICoCo
- Rename the data file to be consistent with the name of the repository:
  - \$ cd Vahl\_Davis\_hexa\_ICoCo
  - \$ mv Vahl\_Davis\_hexa.data Vahl\_Davis\_hexa\_ICoCo.data
- Edit the data file and modify it to add ICoCo instructions:
  - Add the following line after the definition of the dimension: Nom ICoCoProblemName Lire ICoCoProblemName pb
  - Comment the "solve pb" line at the end of the data file.



# With ICoCo

## Creation of the main.cpp file

- Create the main.cpp which will launch the calculation:
  - \$ cp \$TRUST\_ROOT/doc/TRUST/exercices/ICoCo/main1.cpp main.cpp
- You can edit it and see the main method which creates the objects needed to do the information exchanges.
- You can use ICoCo with 1 or more processors.
- Here we use only one processor to solve the problem.

#### Creation of the makefile

- Create a makefile for your calculation on 1 proc:
  - \$ sh \$project\_directory/share/bin/create\_Makefile 1
- Compile it:
  - \$ make
- It creates an executable "couplage" and a data file "couplage.data".

# With ICoCo

#### Launch calculation

- Execute it:
  - \$./couplage
- You may obtain the same results as with TRUST executable.
- Compare it:
  - \$ compare\_lata Vahl\_Davis\_hexa\_ICoCo.lml
  - ../Vahl\_Davis\_hexa\_trust/Vahl\_Davis\_hexa\_trust.lml
- The files are the sames!

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## Adjusting the main.cpp file

- Copy the following test case in your repository:
  - \$ cd ICoCo exercises
  - \$ cp -r Vahl\_Davis\_hexa\_ICoCo Vahl\_Davis\_hexa\_ICoCo\_exchange
  - \$ cd Vahl\_Davis\_hexa\_ICoCo\_exchange
- Clean the repository and rename the data file:
  - \$ trust -clean
  - \$ mv Vahl\_Davis\_hexa\_ICoCo.data
  - Vahl\_Davis\_hexa\_ICoCo\_exchange.data
- We have to add a block in the main.cpp file to made exchanges.
- Copy the following file in your folder:
  - \$ cp \$TRUST\_ROOT/doc/TRUST/exercices/ICoCo/main2.cpp main.cpp

### Adjusting the data file

- Then modify the data file Vahl\_Davis\_hexa\_ICoCo\_exchange.data to made the input:
  - Change the line:
    - "Gauche Paroi\_temperature\_imposee Champ\_Front\_Uniforme 1 10." to
    - "Gauche Paroi\_temperature\_imposee ch\_front\_input { nb\_comp 1 nom TEMPERATURE IN DOM probleme pb }"
  - You can see that the name "TEMPERATURE\_IN\_DOM" is the one employed in the main.cpp file.

#### Launch calculation

- Now we can compile and launch the calculation:
  - \$ make
  - \$ ./couplage
- You may obtain the same results as with TRUST executable.
- Compare it:
  - \$ compare\_lata Vahl\_Davis\_hexa\_ICoCo\_exchange.lml
- ../Vahl\_Davis\_hexa\_trust/Vahl\_Davis\_hexa\_trust.lml
- The files are the sames! (but not for the first time!!!!!)

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## Adjusting the main.cpp file

- Copy the following test case in your repository:
  - \$ cd ICoCo\_exercises
  - \$ cp -r Vahl\_Davis\_hexa\_ICoCo\_exchange
  - Vahl\_Davis\_hexa\_ICoCo\_para
  - \$ cd Vahl\_Davis\_hexa\_ICoCo\_para
- Clean the repository and rename the data file:
  - \$ trust -clean
  - \$ rm main.cpp
  - \$ mv Vahl\_Davis\_hexa\_ICoCo\_exchange.data
  - Vahl\_Davis\_hexa\_ICoCo\_para.data
  - \$ trust -partition Vahl\_Davis\_hexa\_ICoCo\_para
- Copy the following file in your folder:
  - \$ cp \$TRUST\_ROOT/doc/TRUST/exercices/ICoCo/main3.cpp main.cpp

### Adjusting the main.cpp file

- Open the main.cpp file and search for the MPI command lines.
- Open this file and look where:
  - o the processors are added: search for "dom ids"
  - o the names of the data files: search for "data file"
- Compile your new file:
  - \$ make
- To run parallel, you have to use the following mpirun command:
  - \$ mpirun -np 2 ./couplage
- Compare your results with the sequential ones:
  - \$ compare\_lata PAR\_Vahl\_Davis\_hexa\_ICoCo\_para.lml
  - ../Vahl\_Davis\_hexa\_trust/Vahl\_Davis\_hexa\_trust.lml



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# First with TRUST

#### Launch calculation with TRUST

- Copy a TRUST test case:
  - \$ cd ICoCo\_exercises
  - \$ trust -copy docond\_VEF\_3D
  - \$ mv docond\_VEF\_3D docond\_VEF\_3D\_trust
- Launch calculation:
  - \$ cd docond\_VEF\_3D\_trust
  - \$ trust docond\_VEF\_3D
- Let do it in parallel also:
  - \$ trust -partition docond\_VEF\_3D
  - \$ trust PAR\_docond\_VEF\_3D 2
- Compare the two results:
  - \$ compare\_lata docond\_VEF\_3D.lml PAR\_docond\_VEF\_3D.lml
- The results are the same. Differences are below the threshold:  $10^{-5}$ !

## Separate the meshes

- Create your ICoCo test case:
  - \$ cd ICoCo\_exercises
  - \$ trust -copy docond\_VEF\_3D
  - \$ mv docond\_VEF\_3D docond\_VEF\_3D\_ICoCo
  - \$ cd docond\_VEF\_3D\_ICoCo
- Separate mesh and calculation datas in two data files:
  - \$ cp docond\_VEF\_3D.data docond\_VEF\_3D\_mesh1.data
- Edit the file docond VEF 3D mesh1.data and remove:
  - the time scheme
  - the problem definitions
  - the 'scatter' block
  - the discretization, medium, gravity definition blocks
  - the associations
  - the definition and "Read pb1"/"Read pb2" block
  - the "fichier pb1"/"fichier pb2" lines
  - the "solve" keyword

#### Separate the meshes

- Keep only the domain definitions, meshes and cutting steps.
- Then create a data file for each domain:
- \$ cp docond\_VEF\_3D\_mesh1.data docond\_VEF\_3D\_mesh2.data
- In the file docond VEF 3D mesh1.data, keep only the informations of the solide domain.
- In the file docond VEF 3D mesh2.data, keep only the informations of the fluide domain.
- Uncomment the 'partition' step for each domain.
- Run these data files:
  - \$ trust docond\_VEF\_3D\_mesh1
  - \$ trust docond\_VEF\_3D\_mesh2
- You must have now the four files: DOM1 0000.Zones DOM2 0001.Zones DOM1 0001.Zones DOM2 0000.Zones

### Run with separated meshes

- In docond VEF 3D.data file:
  - o remove the mesh and cutting command (which are already in the mesh data file)
  - keep the 'scatter' command and uncomment it
- Launch the calculation:
  - \$ trust docond VEF 3D 2
- Compare results with previous parallel results:
  - \$ compare\_lata docond\_VEF\_3D.lml
  - ../docond\_VEF\_3D\_trust/PAR\_docond\_VEF\_3D.lml
- The results are the sames.
- Separate results in two lml files:
  - o add "fichier pb1.lml" in the "Post processing" block of the solid domain,
  - o add "fichier pb2.lml" in the "Post processing" block of the fluid domain,
- Run calculation to create this two files:
  - \$ trust docond\_VEF\_3D 2

### Separate data files

- Create a data file for the solid domain:
  - \$ cp docond\_VEF\_3D.data docond\_VEF\_3D\_dom1.data
- In docond VEF 3D dom1.datafile, remove the lines:
  - o Probleme Couple pbc
  - Associate pbc pb1
  - Associate pbc pb2
- Change the following lines:
  - Associate pbc sch → Associate pb sch
  - Discretize pbc dis → Discretize pb dis
  - $\circ$  Solve pbc  $\rightarrow$  Solve pb
- Create a data file for the fluide domain:
  - \$ cp docond\_VEF\_3D\_dom1.data docond\_VEF\_3D\_dom2.data
- In docond VEF 3D dom1.data, keep only the informations about the solid domain (pb1, dom solide).
- Subtitute pb1  $\rightarrow$  pb

### Separate data files

- In docond VEF 3D dom2.data, keep only the informations about the fluide domain (pb2, dom fluide).
- Subtitute pb2  $\rightarrow$  pb
- You can see that the heat exchange is made on the "Paroi echange1" boundary for solid domain and "Paroi echange2" boundary for the fluid domain.
- Modify docond VEF 3D dom1.data file to have: Paroi echange1 paroi contact pb Paroi echange2
  - Paroi echange1 paroi temperature imposee Champ Front Uniforme 1 50.
- Modify docond VEF 3D dom2.data file to have: Paroi echange2 paroi contact pb Paroi echange1
  - Paroi echange2 paroi temperature imposee Champ Front Uniforme 1 50.

### Separate data files

- Run docond\_VEF\_3D\_dom1.data and docond\_VEF\_3D\_dom2.data in parallel:
  - \$ trust docond\_VEF\_3D\_dom1 2
  - \$ trust docond\_VEF\_3D\_dom2 2
- The two problems must run.
- Notice that there is no coupling at all for the moment.

#### Adjusting data files

- To create the ICoCo problem, just after 'dimension 3', add in docond\_VEF\_3D\_dom1.data and docond\_VEF\_3D\_dom2.data: 'Nom ICoCoProblemName Lire ICoCoProblemName pb'
- Remove 'Solve pb' because the solving step will be made by ICoCo.



## Creation of the main.cpp file

- We have to create a new executable which will use our data files.
- Copy the following main.cpp file in your repository:
  - \$ cp \$TRUST\_ROOT/doc/TRUST/exercices/ICoCo/main3.cpp main.cpp
- Open this file and look where:
  - the processors are added: search for "dom1 ids"
  - o the names of the data files: search for "data file"
  - o the loop to iterate on time steps: search for "while"

#### Compiling and launching

- Create a makefile to compile your main.cpp file:
  - \$ sh \$project\_directory/share/bin/create\_Makefile 4
- Compile the main.cpp file:
  - \$ make
- Launch calculation:
  - \$ mpirun -np 4 ./couplage
- Compare the results to the results of the coupled problem:
  - \$ compare\_lata pb1.lml docond\_VEF\_3D\_dom1.lml
  - \$ compare\_lata pb2.lml docond\_VEF\_3D\_dom2.lml
- As expected, there are differences between the results because there is no coupling here, we impose the temperature in the data files.

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## Adjusting data file

- Now we want to send the temperature from the Paroi echange2 boundary of the domain dom2 to the Paroi echange1 boundary of the domain dom1.
- Create a new directory:

```
$ cd ICoCo_exercises
$ cp -r docond_VEF_3D_ICoCo docond_VEF_3D_ICoCo_coupling1
```

\$ cd docond\_VEF\_3D\_ICoCo\_coupling1

• In the docond VEF 3D dom2.data data file, add in the "Definition champs" block of the post-processings:

```
TEMPERATURE_OUT_DOM2 Interpolation {
 localisation elem
 domaine dom_fluide_boundaries_Paroi_echange2
 source refChamp { Pb_Champ pb temperature }
```

where "dom fluide boundaries Paroi echange2" is a predefined name for the boundary Paroi echange2 of the domain dom2.

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## Adjusting data file

- In the docond VEF 3D dom1.data data file, change the boundary condition on the 'Paroi echange1' boundary to: Paroi echange1 paroi temperature imposee ch front input { nb comp 1 nom TEMPERATURE IN DOM1 probleme pb }
- Copy the main.cpp file for this exchange: \$ cp \$TRUST\_ROOT/doc/TRUST/exercices/ICoCo/main4.cpp main.cpp
- Compare the new main.cpp file to the previous one: \$ tkdiff main.cpp ../docond\_VEF\_3D\_ICoCo/main.cpp
- You can see where the new fields TEMPERATURE IN DOM1 and TEMPERATURE OUT DOM2 added to the data files, are used in a new part which makes exchanges.
- Notice that we use two new objects: one TrioDEC object and one TrioField object.
- Some comments are written to help you.

## Adjusting data file

- Compile the main.cpp file:
  - \$ make
- Launch calculation:
  - \$ mpirun -np 4 ./couplage
- Compare results with those without coupling:
  - \$ compare\_lata docond\_VEF\_3D\_dom1.lml
  - ../docond\_VEF\_3D\_ICoCo/docond\_VEF\_3D\_dom1.lml
  - \$ compare\_lata docond\_VEF\_3D\_dom2.lml
  - ../docond\_VEF\_3D\_ICoCo/docond\_VEF\_3D\_dom2.lml
- The results on the domain dom2 are the same as this calculation made only one more post-traitment.
- But we can see that the coupling works well because the results on the domain dom1 changes.

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## Adjusting data file

- Two way coupling for thermal problems should use Dirichlet and Neumann boundary conditions (using only Dirichlet boundary conditions for coupling both sides would not work).
- So we want to send the heat flux from the Paroi\_echange1 boundary of the domain dom1 to the Paroi\_echange2 boundary of the domain dom2.
- Create a new directory:
  - \$ cd ICoCo\_exercises
  - \$ cp -r docond\_VEF\_3D\_ICoCo\_coupling1
  - ${\tt docond\_VEF\_3D\_ICoCo\_coupling2}$
  - \$ cd docond\_VEF\_3D\_ICoCo\_coupling2
- Inspire you of the previous part to make a Neumann boundary condition (heat flux imposed):

## Adjusting data file

• In the docond VEF 3D dom1.data data file, add in the "Definition champs" block of the post-processings: FLUX\_SURFACIQUE\_OUT\_DOM1 Interpolation { localisation elem domaine dom\_fluide\_boundaries\_Paroi\_echange1 source Morceau\_equation { type operateur numero 0 option flux\_surfacique\_bords source refChamp { Pb\_Champ pb temperature } where "dom solide boundaries Paroi echange1" is a predefined name for

• In the docond VEF 3D dom2.data data file, change the boundary condition on the 'Paroi echange2' boundary to: Paroi echange2 paroi flux impose ch front input { nb comp 1 nom

the boundary Paroi echange1 of the domain dom1.

FLUX SURFACIQUE IN DOM2 probleme pb }

### Adjusting data file

- Modify the main.cpp file to add a new exchange:
  - Create a new TrioDEC object to made exchange from domain dom2 to domain dom1:

```
TrioDEC dec flux surfacique2(dom2 ids, dom1 ids);
```

- Create a new TrioField object: TrioField field flux surfacique2;
- Add code lines into the while loop to made this exchange of informations.
- You can have a look at the main5.cpp file for this exchange: \$ cp \$TRUST\_ROOT/doc/TRUST/exercices/ICoCo/main5.cpp main5.cpp
- Compare it to the previous one: \$ tkdiff main5.cpp ../docond\_VEF\_3D\_ICoCo\_coupling1/main.cpp
- You can see the use of the new fields FLUX SURFACIQUE IN DOM2 and FLUX SURFACIQUE OUT DOM1 added to the data files.

### Adjusting data file

- Compile your main.cpp file:
  - \$ make
- Launch calculation:
  - \$ mpirun -np 4 ./couplage
- Compare results with the first ones:
  - \$ compare\_lata docond\_VEF\_3D\_dom1.lml
  - ../docond\_VEF\_3D\_ICoCo/pb1.lml
  - \$ compare\_lata docond\_VEF\_3D\_dom2.lml
  - ../docond\_VEF\_3D\_ICoCo/docond\_VEF\_3D\_dom2.lml