# Supplement

High level programming in OpenFOAM® Building blocks

## Roadmap

- 1. Programming in OpenFOAM®. Building blocks.
- 2. <u>Implementing</u> boundary conditions using high level programming
- 3. **Modifying applications** Highlights
- 4. Implementing an application from scratch
- 5. Adding the scalar transport equation to icoFoam

- In the directory \$WM\_PROJECT\_DIR/applications/test, you will find the source code of several test cases that show the usage of most of the OpenFOAM® classes.
- We highly encourage you to take a look at these test cases and try to understand how to use the classes.
- We will use these basic test cases to understand the following base classes: tensors, fields, mesh, and basic discretization.
- For your convenience, we already copied the directory
   \$WM\_PROJECT\_DIR/applications/test into the directory
   \$PTOFC/programming playground/test

- During this session we will study the building blocks to write basic programs in OpenFOAM®:
  - First, we will start by taking a look at the <u>algebra of tensors</u> in OpenFOAM®.
  - Then, we will take a look at how to generate tensor fields from tensors.
  - Next, we will learn how to access mesh information.
  - Finally we will see how to discretize a model equation and solve the linear system of equations using OpenFOAM® classes and templates.
  - And of course, we are going to program a little bit in C++. But do not be afraid, after all this is not a C++ course.
- Remember, all OpenFOAM® components are implemented in library form for easy re-use.
- OpenFOAM® encourage code re-use. So basically, we are going to take something that already exist, and we are going to modify it to fix our needs.
- We like to call this method CPAC (copy-paste-adapt-compile).

#### Basic tensor classes in OpenFOAM®

- OpenFOAM® represents scalars, vectors and matrices as tensor fields. A zero rank tensor is a scalar, a first rank tensor is a vector and a second rank tensor is a matrix.
- OpenFOAM® contains a C++ class library named **primitive** (\$FOAM\_SRC/OpenFOAM/primitives/). In this library, you will find the classes for the tensor mathematics.
- In the following table, we show the <u>basic tensor classes available in OpenFOAM®</u>, with their respective access functions.

Tensor Rank	Common name	Basic class	Access function
0	Scalar	scalar	
1	Vector	vector	x(), y(), z()
2	Tensor	tensor	xx(), xy(), xz()

#### **Basic tensor classes in OpenFOAM®**

In OpenFOAM®, the second rank tensor (or matrix)

$$\mathbf{T} = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{pmatrix}$$

can be declared in the following way

• We can access the component  $\,T_{13}\,$  or  $\,T_{xz}\,$  using the  ${\bf xz}$  ( ) access function,

#### **Basic tensor classes in OpenFOAM®**

For instance, the following statement,

Info << "Txz = " << T.xz ( ) << endl; 
$$\mathbf{T} = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{pmatrix}$$

Will generate the following screen output,

$$\Rightarrow Txz = 3$$

- Notice that to output information to the screen in OpenFOAM®, we use the <u>function Info</u> instead of the function cout (used in standard C++).
- The function **cout** will work fine, but it will give you problems when running in parallel.

#### Algebraic tensor operations in OpenFOAM®

- Tensor operations operate on the entire tensor entity.
- OpenFOAM® syntax closely mimics the syntax used in written mathematics, using descriptive functions (e.g. mag) or symbolic operators (e.g. +).
- OpenFOAM® also follow the standard rules of linear algebra when working with tensors.
- Some of the algebraic tensor operations are listed in the following table (where a and b are vectors, s is a scalar, and T is a tensor).

Operation	Remarks	Mathematical description	OpenFOAM® description
Addition		a + b	a + b
Scalar multiplication		s <b>a</b>	s * a
Outer product	rank a, b >=1	ab	a * b
Inner product	rank a, b >=1	a.b	a & b
Double inner product	rank a, b >=2	a:b	a && b
Magnitude		a	mag(a)
Determinant		det <b>T</b>	det(T)

#### **Dimensional units in OpenFOAM®**

- As we already know, OpenFOAM® is fully dimensional.
- Dimensional checking is implemented as a safeguard against implementing a meaningless operation.
- OpenFOAM® encourages the user to attach dimensional units to any tensor and it will perform dimension checking of any tensor operation.
- You can find the dimensional classes in the directory \$FOAM\_SRC/OpenFOAM/dimensionedTypes/
- The dimensions can be hardwired directly in the source code or can be defined in the input dictionaries.
- From this point on, we will be attaching dimensions to all the tensors.

#### **Dimensional units in OpenFOAM®**

- Units are defined using the dimensionSet class tensor, with its units defined using the dimensioned<Type> template class, the <Type> being scalar, vector, tensor, etc. The dimensioned<Type> stores the variable name, the dimensions and the tensor values.
- For example, a tensor with dimensions is declare in the following way:

- In line 1 we create the object sigma.
- In line 4, we use the class dimensonSet to attach units to the object sigma.
- In line 5, we set the input values of the tensor **sigma**.

#### Units correspondence in dimensionSet

The units of the class dimensionSet are defined as follows

```
dimensionSet (kg, m, s, K, mol, A, cd)
```

Therefore, the tensor sigma,

 $\,\cdot\,\,$  Has pressure units or  $\,\,\,kg\,\,m^{-1}\,s^{-2}$ 

#### **Dimensional units examples**

- To attach dimensions to any tensor, you need to access dimensional units class.
- To do so, just add the header file dimensionedTensor.H to your program.

```
#include "dimensionedTensor.H"

...
dimensionedTensor sigma
(
    "sigma",
    dimensionSet(1, -1, -2, 0, 0, 0, 0),
    tensor(1e6,0,0,0,1e6,0,0,0,1e6)
);
Info<< "Sigma: " << sigma << endl;
...
...
...</pre>
```

The output of the previous program should look like this:

```
sigma sigma [1 -1 -2 0 0 0 0] (1e+06 0 0 0 1e+06 0 0 0 1e+06)
```

#### **Dimensional units examples**

- As for base tensors, you can access the information of dimensioned tensors.
- For example, to access the name, dimensions, and values of a dimensioned tensor, you can proceed as follows:

```
Info << "Sigma name: " << sigma.name ( ) << endl;
Info << "Sigma dimensions: " << sigma.dimensions ( ) << endl;
Info << "Sigma value: " << sigma.value ( ) << endl;</pre>
```

To extract a value of a dimensioned tensor, you can proceed as follows:

```
Info<< "Sigma yy (22) value: " << sigma.value().yy() << endl;</pre>
```

- Note that the value() member function first converts the expression to a tensor, which has a yy()
  member function.
- The **dimensionedTensor** class does not have a **yy()** member function, so it is not possible to directly get its value by using **sigma.yy()**.

#### OpenFOAM® lists and fields

- OpenFOAM® frequently needs to store sets of data and perform mathematical operations.
- OpenFOAM® provides an array template class List<Type>, making it possible to create a list of any object of class Type that inherits the functions of the Type. For example a List of vector is List<vector>.
- Lists of the tensor classes are defined in OpenFOAM® by the template class Field<Type>.
- For better code legibility, all instances of Field<Type>, e.g. Field<vector>, are renamed using typedef declarations as scalarField, vectorField, tensorField, symmTensorField, tensorThirdField and symmTensorThirdField.
- You can find the field classes in the directory \$FOAM\_SRC/OpenFOAM/fields/Fields.
- Algebraic operations can be performed between fields, subject to obvious restrictions such as the fields having the same number of elements.
- OpenFOAM® also supports operations between a field and a zero-rank tensor, e.g. all values of a **Field U** can be multiplied by the **scalar** 2 by simple coding the following line, **U = 2.0 \* U**.

#### Construction of a tensor field in OpenFOAM®

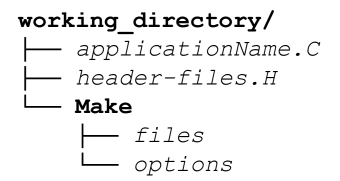
- To create fields, you need to access the tensor class.
- To do so, just add the header file tensorField.H to your program. This class inherit all the tensor algebra.

```
#include "tensorField.H"
...
tensorField tf1(2, tensor::one);
Info<< "tf1: " << tf1 << endl;
tf1[0] = tensor(1, 2, 3, 4, 5, 6, 7, 8, 9);
Info<< "tf1: " << tf1 << endl;
Info<< "tf1: " << tf1 << endl;
...
...
...
...
...</pre>
```

- In this example, we created a list of two tensor fields (tf1), and both tensors are initialized to
  one.
- We can access components on the list using the access operator [].

#### **Example of use of tensor and field classes**

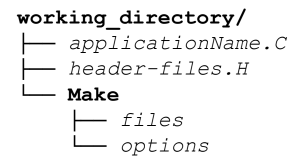
- In the directory \$PTOFC/programming\_playground/my\_tensor you will find a tensor class example.
- The original example is located in the directory \$HOME/OpenFOAM/OpenFOAM-8/applications/test. Feel free to compare the files to spot the differences.
- Before compiling the file, let us recall how applications are structure,



- applicationName.C: is the actual source code of the application.
- header\_files.H: header files required to compile the application.

#### Example of use of tensor and field classes

Before compiling the file, let us recall how applications are structure.



- The Make directory contains compilation instructions.
  - files: names all the source files (.C), it specifies the name of the new application and the location of the output file.
  - options: specifies directories to search for include files and libraries to link the solver against.
- At the end of the file files, you will find the following line of code,
   EXE = \$(FOAM\_USER\_APPBIN)/my\_Test-tensor
- This is telling the compiler to name your application my\_Test-tensor and to copy the executable in the directory \$FOAM\_USER\_APPBIN.
- To avoid conflicts between applications, always remember to give a proper name and a location to your programs and libraries.

#### **Example of use of tensor and field classes**

• Let us now compile the tensor class example. Type in the terminal:

```
1. $\$ cd $PTOFC/programming_playground/my_tensor
```

- 2. | \$> wmake
- 3. | \$> my\_Test-tensor
- In step 2, we used wmake (distributed with OpenFOAM®) to compile the source code.
- The name of the executable will be my\_Test-tensor and it will be located in the directory
   \$FOAM\_USER\_APPBIN (as specified in the file Make/files)
- At this point, take a look at the output and study the file <code>Test-tensor.C</code>. Try to understand what we have done.
- After all, is not that difficult. Right?

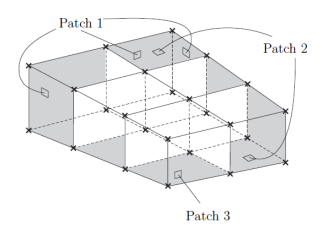
- At this point, we are a little bit familiar with tensor, fields, and lists in OpenFOAM®.
- They are the base to building applications in OpenFOAM®.
- Let us now take a look at the whole solution process:
  - Creation of the tensors.
  - Mesh assembly.
  - Fields creation.
  - Equation discretization.
- All by using OpenFOAM® classes and template classes

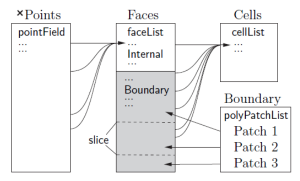
#### Discretization of a tensor field in OpenFOAM®

- The discretization is done using the FVM (Finite Volume Method).
- The cells are contiguous, *i.e.*, they do not overlap and completely fill the domain.
- Dependent variables and other properties are stored at the cell centroid.
- No limitations on the number of faces bounding each cell.
- No restriction on the alignment of each face.
- The mesh class polyMesh is used to construct the polyhedral mesh using the minimum information required.
- You can find the polyMesh classes in the directory \$FOAM\_SRC/OpenFOAM/meshes
- The fvMesh class extends the polyMesh class to include additional data needed for the FVM discretization.
- You can find the fvMesh classes in the directory \$FOAM\_SRC/src/finiteVolume/fvMesh

#### Discretization of a tensor field in OpenFOAM®

- The template class geometricField relates a tensor field to a fvMesh
- Using typedef declarations geometricField is renamed to volField (cell center), surfaceField (cell faces), and pointField (cell vertices).
- You can find the geometricField classes in the directory \$FOAM\_SRC/OpenFOAM/fields/GeometricFields.
- The template class geometricField stores internal fields, boundary fields, mesh information, dimensions, old values and previous iteration values.
- A geometricField inherits all the tensor algebra of its corresponding field, has dimension checking, and can be subjected to specific discretization procedures.
- Let us now access the mesh information of a simple case.



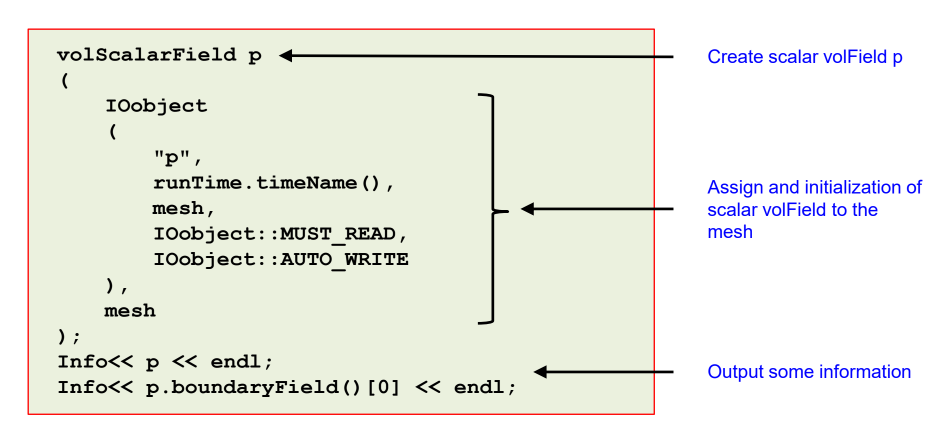


#### Data stored in the fvMesh class

Class	Description	Symbol	Access function
volScalarField	Cell volumes	V	V()
surfaceVectorField	Face area vector	$\mathbf{S}_f$	Sf()
surfaceScalarField	Face area magnitude	$ \mathbf{S}_f $	magSf()
volVectorField	Cell centres	${f C}$	C()
surfaceVectorField	Face centres	$\mathbf{C}_f$	Cf()
surfaceScalarField	Face fluxes	$\phi_g$	Phi()

### Accessing fields defined in a mesh

- To access fields defined at cell centers of the mesh you need to use the class volField.
- The class volField can be accessed by adding the header volFields.H to your program.



#### Accessing fields using for loops

To access fields using for loops, we can use OpenFOAM® macro forAll, as follows,

```
Outputs name of patch

forAll(mesh.boundaryMesh(), patchI)

Info << "Patch " << patchI << ": " << mesh.boundary()[patchI].name() << " with "

<< mesh.boundary()[patchI].Cf().size() << " faces. Starts at total face "

<< mesh.boundary()[patchI].start() << endl;

Outputs size of patch (number of faces)
```

- In the previous statement **mesh.boundaryMesh()** is the size of the loop, and **patchI** is the iterator. The iterator always starts from zero.
- The forAll loop is equivalent to the standard for loop in C++.

• Notice that we used as iterator i instead of patchI, this does not make any difference.

#### **Equation discretization in OpenFOAM®**

- At this stage, OpenFOAM® converts the PDEs into a set of linear algebraic equations, A x = b, where x and b are volFields (geometricField).
- A is a fvMatrix, which is created by the discretization of a geometricField and inherits the
  algebra of its corresponding field, and it supports many of the standard algebraic matrix
  operations.
- The fvm (finiteVolumeMethod) and fvc (finiteVolumeCalculus) classes contain static functions for the differential operators and discretize any geometricField.
- fvm returns a fvMatrix, and fvc returns a geometricField.
- In the directories \$FOAM\_SRC/finiteVolume/finiteVolume/fvc and \$FOAM\_SRC/finiteVolume/finiteVolume/fvm you will find the respective classes.
- Remember, the PDEs or ODEs we want to solve involve derivatives of tensor fields with respect
  to time and space. What we re doing at this point, is applying the finite volume classes to the
  fields, and assembling a linear system.

#### Discretization of the basic PDE terms in OpenFOAM®

The list is not complete

Term description	Mathematical expression	fvm:: fvc::
Laplacian	$ abla^2\phi$ , $ abla\cdot\Gamma abla\phi$	laplacian (phi) laplacian (Gamma, phi)
Time derivative	$rac{\partial \phi}{\partial t}$ , $rac{\partial  ho \phi}{\partial t}$	ddt(phi) ddt(rho,phi)
Convection	$ abla \cdot (\psi)$ , $ abla \cdot (\psi \phi)$	div(psi,scheme) div(psi,phi)
Source	$ ho\phi$	Sp(rho,phi) SuSp(rho,phi)
$\phi$ vol <type>Field <math> ho</math></type>	scalar, volScalarField	$\psi$ surfaceScalarField

#### Discretization of the basic PDE terms in OpenFOAM®

- To discretize the fields in a valid mesh, we need to access the finite volume class. This class can be accessed by adding the header  $f_{VCFD.H}$  to your program.
- To discretize the scalar transport equation in a mesh, we can proceed as follows,

```
Assemble and solve linear system arising form the discretization

fvm::ddt(T)

+ fvm::div(phi,T)

- fvm::laplacian(DT,T)

);
```

- Remember, you will need to first create the mesh, and initialize the variables and constants.
   That is, all the previous steps.
- Finally, everything we have done so far inherits all parallel directives. There is no need for specific parallel programming.

#### Discretization of the basic PDE terms in OpenFOAM®

The previous discretization is equivalent to,

```
fvScalarMatrix TEqn
{
    fvm::ddt(T)
    + fvm::div(phi,T)
    - fvm::laplacian(DT,T)
);
Creates object TEqn that contains the coefficient matrix arising from the discretization

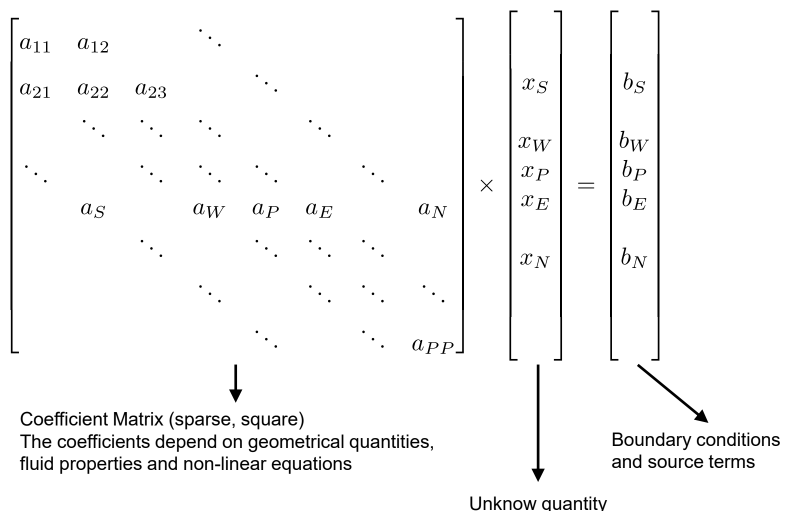
Discretize equations

Solve the linear system Teqn
```

- Here, fvScalarMatrix contains the matrix derived from the discretization of the model equation.
- fvScalarMatrix is used for scalar fields and fvVectorMatrix is used for vector fields.
- This syntax is more general, since it allows the easy addition of terms to the model equations.

#### Discretization of the basic PDE terms in OpenFOAM®

At this point, OpenFOAM® assembles and solves the following linear system,



#### **Example of use of tensor and field classes**

Let us study a **fvMesh** example. First let us compile the program my Test-mesh. Type in the terminal,

```
    $> cd $PTOFC/programming_playground/my_mesh/
    $> wmake
```

- To access the mesh information, we need to use this program in a valid mesh.
  - 1. | \$> cd \$PTOFC/programming\_playground/my\_mesh/cavity

  - 2. \$\\$ blockMesh3. \$\\$ my\_Test-mesh
- At this point, take a look at the output and study the file Test-mesh. C. Try to understand what we have done.
- FYI, the original example is located in the directory \$PTOFC/programming playground/test/mesh.

#### A few OpenFOAM® programming references

- You can access the API documentation in the following link, <a href="https://cpp.openfoam.org/v5/">https://cpp.openfoam.org/v5/</a>
- You can access the coding style guide in the following link, <a href="https://openfoam.org/dev/coding-style-guide/">https://openfoam.org/dev/coding-style-guide/</a>
- You can report programming issues in the following link, <a href="https://bugs.openfoam.org/rules.php">https://bugs.openfoam.org/rules.php</a>
- You can access openfoamwiki coding guide in the following link, <a href="http://openfoamwiki.net/index.php/OpenFOAM\_guide">http://openfoamwiki.net/index.php/OpenFOAM\_guide</a>
- You can access the user guide in the following link, https://cfd.direct/openfoam/user-guide/
- You can read the OpenFOAM® Programmer's guide in the following link (it seems that this guide is not supported anymore), <a href="http://foam.sourceforge.net/docs/Guides-a4/ProgrammersGuide.pdf">http://foam.sourceforge.net/docs/Guides-a4/ProgrammersGuide.pdf</a>

#### A few good C++ references

- The C++ Programming Language. B. Stroustrup. 2013, Addison-Wesley.
- The C++ Standard Library. N. Josuttis. 2012, Addison-Wesley.
- C++ for Engineers and Scientists. G. J. Bronson. 2012, Cengage Learning.
- Sams Teach Yourself C++ in One Hour a Day. J. Liberty, B. Jones. 2004, Sams Publishing.
- C++ Primer. S. Lippman, J. Lajoie, B. Moo. 2012, Addison-Wesley.
- http://www.cplusplus.com/
- http://www.learncpp.com/
- http://www.cprogramming.com/
- http://www.tutorialspoint.com/cplusplus/
- http://stackoverflow.com/

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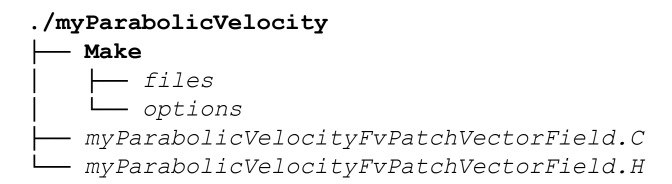
- Hereafter we will work with high level programming, this is the hard part of programming in OpenFOAM®.
- High level programming requires some knowledge on C++ and OpenFOAM® API library.
- Before doing high level programming, we highly recommend you to try with codeStream, most
  of the time it will work.
- We will implement the parabolic profile, so you can compare this implementation with codeStream ad codedFixedValue BCs.
- When we program boundary conditions, we are building a new library that can be linked with any solver. To compile the library, we use the command wmake (distributed with OpenFOAM®).
- At this point, you can work in any directory, but we recommend you to work in your OpenFOAM® user directory, type in the terminal,

Let us create the basic structure to write the new boundary condition, type in the terminal,

```
    $> foamNewBC -f -v myParabolicVelocity
    $> cd myParabolicVelocity
```

- The utility foamNewBC, will create the directory structure and all the files needed to write your own boundary conditions.
- We are setting the structure for a fixed (the option -f) velocity (the option -v), boundary condition, and we name our boundary condition Parabolic Velocity.
- If you want to get more information on how to use foamNewBC, type in the terminal,

#### Directory structure of the new boundary condition



The directory contains the source code of the boundary condition.

- myParabolicVelocityFvPatchVectorField. C: is the actual source code of the application. This file contains the definition of the classes and functions.
- myParabolicVelocityFvPatchVectorField. H: header files required to compile the application. This file contains variables, functions and classes declarations.
- The Make directory contains compilation instructions.
  - Make/files: names all the source files (.C), it specifies the boundary condition library name and location of the output file.
  - Make/options: specifies directories to search for include files and libraries to link the solver against.

#### The header file (.H)

Let us start to do some modifications. Open the header file using your favorite text editor (we use gedit).

```
//- Single valued scalar quantity, e.g. a coefficient
96
97
      scalar scalarData ;
98
      //- Single valued Type quantity, e.g. reference pressure pRefValue
99
      // Other options include vector, tensor
100
101
      vector data ;
102
103
      //- Field of Types, typically defined across patch faces
      // e.g. total pressure p0 . Other options include vectorField
104
      vectorField fieldData ;
105
106
      //- Type specified as a function of time for time-varying BCs
107
      autoPtr<Function1<vector>> timeVsData ;
108
109
110
      //- Word entry, e.g. pName for name of the pressure field on database
      word wordData ;
111
112
      //- Label, e.g. patch index, current time index
113
      label labelData ;
114
115
      //- Boolean for true/false, e.g. specify if flow rate is volumetric
116
      bool boolData ;
117
118
119
120
      // Private Member Functions
121
122
      //- Return current time
123
      scalar t() const;
```

- In lines 96-123 different types of private data are declared.
- These are the variables we will use for the implementation of the new BC.
- In our implementation we need to use vectors and scalars, therefore we can keep the lines 97 and 101.
- We can delete lines 103-117, as we do not need those datatypes.
- Also, as we will use two vectors in our implementation, we can duplicate line 101.
- You can leave the rest of the file as they are.

#### The header file (.H)

At this point, your header file should looks like this one,

```
//- Single valued scalar quantity, e.g. a coefficient
scalar scalarData_;

//- Single valued Type quantity, e.g. reference pressure pRefValue_
// Other options include vector, tensor
vector data_;
vector data_;
```

- Change the name of scalarData\_ to maxValue\_ (line 97).
- Change the names of the two vectors data\_ (lines 101-102). Name the first one n\_ and the last one y\_.

```
96
      //- Single valued scalar quantity, e.g. a coefficient
      scalar maxValue ;
97
98
                                                                                    It is recommended to initialize
      //- Single valued Type quantity, e.g. reference pressure pRefValue
99
                                                                                    them in the same order as you
      // Other options include vector, tensor
100
                                                                                    declare them in the header file
101
      vector n ;
102
      vector y ;
```

We just declared the variables that we will use. You can now save and close the file.

- Let us start to modify the source file. Open the source file with your favorite editor.
- Lines 34-37 refers to a private function definition. This function allows us to access simulation time. Since in our implementation we do not need to use time, we can safely remove these lines.

```
34    Foam::scalar Foam::myParabolicVelocityFvPatchVectorField::t() const
35    {
36        return db().time().timeOutputValue();
37    }
```

- Let us compile the library to see what errors we get. Type in the terminal,
  - 1. | \$> wmake
- You will get a lot of errors.
- Since we deleted the datatypes **fieldData**, **timeVsData**, **wordData**, **labelData** and **boolData** in the header file, we need to delete them as well in the C file. Otherwise the compiler complains.

- At this point, let us erase all the occurrences of the datatypes fieldData, timeVsData, wordData, labelData, and boolData.
- Locate line 38,

```
38 Foam::myParabolicVelocityFvPatchVectorField::
...
...
```

- Using this line as your reference location in the source code, follow these steps,
  - Erase the following lines in incremental order (be sure to erase only the lines that contain the words **fieldData**, **timeVsData**, **wordData**, **labelData** and **boolData**): 48-52, 63-67, 90-94, 102-106, 115-119, 172-175.
  - Erase the following lines (they contain the word fieldData), 126, 140, 151-153.
  - Replace all the occurrences of the word scalarData with maxValue (11 occurrences).

#### The source file (.C)

Duplicate all the lines where the word **data** appears (6 lines), change the word **data** to **n** in the first line, and to **y** in the second line, erase the comma in the last line. For example,

#### Original statements

```
45 fixedValueFvPatchVectorField(p, iF),
46 maxValue_(0.0),
47 data_(Zero),
48 data_(Zero),
```

#### Modified statements

```
45 fixedValueFvPatchVectorField(p, iF),
46 maxValue_(0.0),
47 n_(Zero),
48 y_(Zero) 		Remember to erase the comma
```

- We are almost done; we just defined all the datatypes. Now we need to implement the actual boundary condition.
- Look for line 147 ( updateCoeffs() member function), and add the following statements,

```
The actual
                                                                                       implementation of
         void Foam::myParabolicVelocityFvPatchVectorField::updateCoeffs()
 147
                                                                                       the BC is always
 148
         {
                                                                                       done in this class
             if (updated())
 149
 150
 151
                  return;
 152
                                                                               Find patch bounds (minimum
 153
                                                                               and maximum points)
 154
             boundBox bb(patch().patch().localPoints(), true);
      lines
 155
 156
             vector ctr = 0.5*(bb.max() + bb.min()); ← Coordinates of patch midpoint
 157
 158
             const vectorField& c = patch().Cf();
Access patch face centers
      Add
 159
             scalarField\ coord = 2*((c - ctr) & y)/((bb.max() - bb.min()) & y);
 160
Computes scalar field to be used for defining the parabolic profile
```

#### The source file (.C)

Add the following statement in line 164,

- The last step before compiling the new BC is to erase a few commas.
- Look for lines 48, 64, 92, 105, 119, and erase the comma at the end of each line.
- At this point we have a valid library where we implemented a new BC.
- Finally, you can go back to the header file (\*.H) and document your boundary condition implementation.
  - You can add the comments in the header of the file (lines 1-73).

- At this point we have a valid library where we have implemented a new BC.
- Try to compile it, we should not get any error (maybe one warning). Type in the terminal,
  - 1. | \$> wmake
- If you are feeling lazy, or if you can not fix the compilation errors, you will find the source code in the directory,
  - \$PTOFC/101programming/src/myParabolicVelocity

- Before moving forward, let us comment a little bit the source file.
- First at all, there are five classes constructors and each of them have a specific task.
- In our implementation we did not use all the classes, we only use the first two classes.
- The first class is related to the initialization of the variables.
- The second class is related to reading the input dictionaries.
- We will not comment on the other classes as it is out of the scope of this example (they deal
  with input tables, mapping, and things like that).
- The implementation of the boundary condition is always done using the updateCoeffs()
  member function.
- When we compile the source code, it will compile a library with the name specified in the file
   Make/file. In this case, the name of the library is libmyParabolicVelocity.
- The library will be located in the directory \$(FOAM\_USER\_LIBBIN), as specified in the file
   Make/file.

- The first class is related to the initialization of the variables declared in the header file.
- In line 47 we initialize **maxValue** with the value of zero. The vectors **n** and **y** are initialized as a zero vector by default or (0, 0, 0).
- It is not a good idea to initialize these vectors as zero vectors by default. Let us use as default initialization (1, 0, 0) for vector **n** and (0,1,0) for vector **y**.

```
38
     Foam::myParabolicVelocityFvPatchVectorField::
39
     myParabolicVelocityFvPatchVectorField
40
41
         const fvPatch& p,
42
         const DimensionedField<vector, volMesh>& iF
43
     )
44
         fixedValueFvPatchVectorField(p, iF),
45
         maxValue (0.0),
46
                              Change to n_(1,0,0)
         n (Zero), ◀
47
         y (Zero)
48
49
50
                        Change to y (0,1,0)
```

- The second class is used to read the input dictionary.
- Here we are reading the values defined by the user in the dictionary U.
- The function lookup will search the specific keyword in the input file.

```
53
     Foam::myParabolicVelocityFvPatchVectorField::
54
     myParabolicVelocityFvPatchVectorField
55
56
         const fvPatch& p,
57
         const DimensionedField<vector, volMesh>& iF,
58
         const dictionary& dict
59
60
61
         fixedValueFvPatchVectorField(p, iF),
                                                                          dict.lookup will look for
         maxValue (readScalar(dict.lookup("maxValue"))),
62
                                                                          these keywords in the
63
         n (pTraits<vector>(dict.lookup("n"))),
                                                                          input dictionary
64
         y (pTraits<vector>(dict.lookup("y")))
65
66
67
68
         fixedValueFvPatchVectorField::evaluate();
77
```

- Since we do not want the vectors **n** and **y** to be zero vectors, we add the following sanity check starting form line 67.
- These statements check if the given **n** and **y** vectors in the input dictionary is zero or not.
- If any of the vectors are zero it gives the fatal error and terminate the program.
- On the other hand, if everything is ok it will normalize **n** and **y** (since in our implementation they are direction vectors).

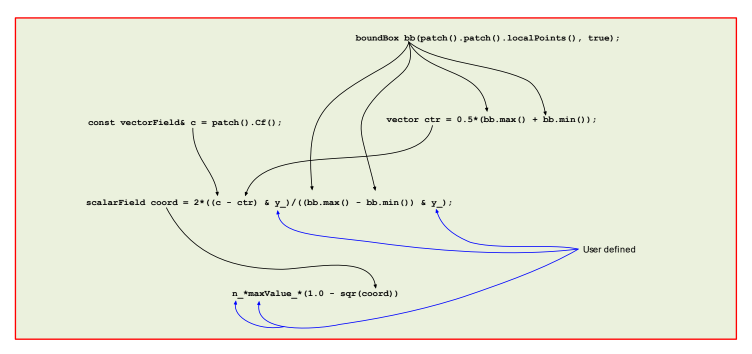
```
66
         if (mag(n) < SMALL \mid \mid mag(y) < SMALL)
                                                                                     Add these statements
67
68
69
               FatalErrorIn("parabolicVelocityFvPatchVectorField(dict)")
                   << "n or y given with zero size not correct"
70
71
                   << abort(FatalError);
72
73
                         //This is equivalent to n = n / mag(n)
74
         n /= mag(n);
                              //This is equivalent to y = y / (mag(y))
         y /= mag(y);
75
76
         fixedValueFvPatchVectorField::evaluate();
77
78
```

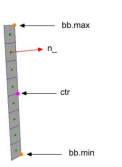
- At this point, we are ready to go.
- Save the files and recompile. Type in the terminal,

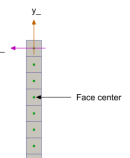
- We should not get any error (maybe one warning).
- At this point we have a valid library that can be linked with any solver.
- If you get compilation errors, read the screen and try to sort it out, the compiler is always telling
  you what is the problem.
- If you are feeling lazy, or if you can not fix the compilation errors, you will find the source code in the directory,
  - \$PTOFC/101programming/src/myParabolicVelocity

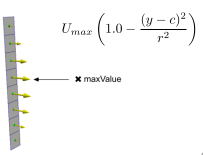
#### The source file (.C)

Before using the new BC, let us take a look at the logic behind the implementation.









#### Running the case

- This case is ready to run, the input files are located in the directory \$PTOFC/101programming/src/case elbow2d
- Go to the case directory,

```
1. | $> cd $PTOFC/101programming/src/case_elbow2d
```

Open the file O/U, and look for the definition of the new BC velocity-inlet-5,

#### Running the case

- We also need to tell the application that we want to use the library we just compiled.
- To do so, we need to add the new library in the dictionary file controlDict,

- The solver will dynamically link the library.
- At this point, we are ready to launch the simulation.

#### Running the case

- This case is ready to run, the input files are located in the directory \$PTOFC/101programming/src/case\_elbow2d
- To run the case, type in the terminal,

```
    $> foamCleanTutorials
    $> fluentMeshToFoam ../../meshes_and_geometries/fluent_elbow2d_1/ascii.msh
    $> icoFoam | tee log.solver
    $> paraFoam
```

- At this point, you can compare the three implementations (codeStream, codedFixedValue and high-level programming).
- All of them will give the same outcome.

#### Adding some verbosity to the BC implementation

- Let us add some outputs to the BC.
- After the member function updateCoeffs (line 177), add the following lines,

```
177
      fixedValueFvPatchVectorField::updateCoeffs();
178
179
      Info << endl << "Face centers (c):" << endl;</pre>
180
      Info << c << endl;</pre>
181
      Info << endl << "Patch center (ctr):" << endl;</pre>
182
      Info << ctr << endl;</pre>
183
      Info << endl << "Patch (c - ctr):" << endl;</pre>
184
      Info << c - ctr << endl;</pre>
185
      Info << endl << "Patch max bound (bb.max):" << endl;</pre>
      Info << bb.max() << endl;</pre>
186
187
      Info << endl << "Patch min bound (bb.max):" << endl;</pre>
188
      Info << bb.min() << endl;</pre>
      Info << endl << "Patch coord ( 2*((c - ctr) & y )/((bb.max() - bb.min()) & y ) ):" << endl;</pre>
189
190
      Info << coord << endl;</pre>
      Info << endl << "Patch ( 1.0 - sqr(coord)) :" << endl;</pre>
191
      Info << n *maxValue *(1.0 - sqr(coord))<< endl;</pre>
192
193
      Info << endl << "Loop for c, BC assignment << endl;</pre>
194
      forAll(c, faceI)
195
196
           Info << c[faceI] << " << n *maxValue *(1.0 - sqr(coord[faceI])) << endl;
197
198
199
```

Recompile, rerun the simulation, look at the output, and do the math.

- In the directory \$PTOFC/101programming/src/myParabolicVelocityMod, you will find
  an implementation of this boundary condition with conditional switches and screen output
  information.
- Try to figure out how this BC works.

#### Do you take the challenge?

- Starting from this boundary condition, try to implement a paraboloid BC.
- If you are feeling lazy or at any point do you get lost, in the directory \$PTOFC/101programming/src/myParaboloidVelocity you will find a working implementation of the paraboloid profile.
- Open the source code and try to understand what we did (pretty much similar to the previous case).
- In the directory \$PTOFC/101programming/src/case\_elbow3d you will find a case ready to use.

# Roadmap

- 1. Programming in OpenFOAM®. Building blocks.
- 2. Implementing boundary conditions using high level programming
- 3. Modifying applications Highlights
- 4. Implementing an application from scratch
- 5. Adding the scalar transport equation to icoFoam

## **Modifying applications – Highlights**

- Implementing a new application from scratch in OpenFOAM® (or any other high-level programming library), can be an incredible daunting task.
- OpenFOAM® comes with many solvers, and as it is today, you do not need to implement new solvers from scratch.
- Of course, if your goal is to write a new solver, you will need to deal with programming. What
  you usually do, is take an existing solver and modify it.
- But in case that you would like to take the road of implementing new applications from scratch, we are going to give you the basic building blocks.
- We are also going to show how to add basic modifications to existing solvers.
- We want to remind you that this requires some knowledge on C++ and OpenFOAM® API library.
- Also, you need to understand the FVM, and be familiar with the basic algebra of tensors.
- Some common sense is also helpful.

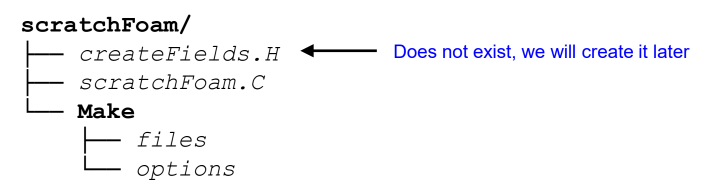
# Roadmap

- 1. Programming in OpenFOAM®. Building blocks.
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- Let us do a little bit of high-level programming, this is the hard part of working with OpenFOAM®.
- At this point, you can work in any directory. But we recommend you to work in your OpenFOAM® user directory, type in the terminal,

- To create the basic structure of a new application, type in the terminal,
  - \$> foamNewApp scratchFoam
     \$> cd scratchFoam
- The utility foamNewApp, will create the directory structure and all the files needed to create the new application from scratch. The name of the application is **scratchFoam**.
- If you want to get more information on how to use foamNewApp, type in the terminal,

#### Directory structure of the new boundary condition



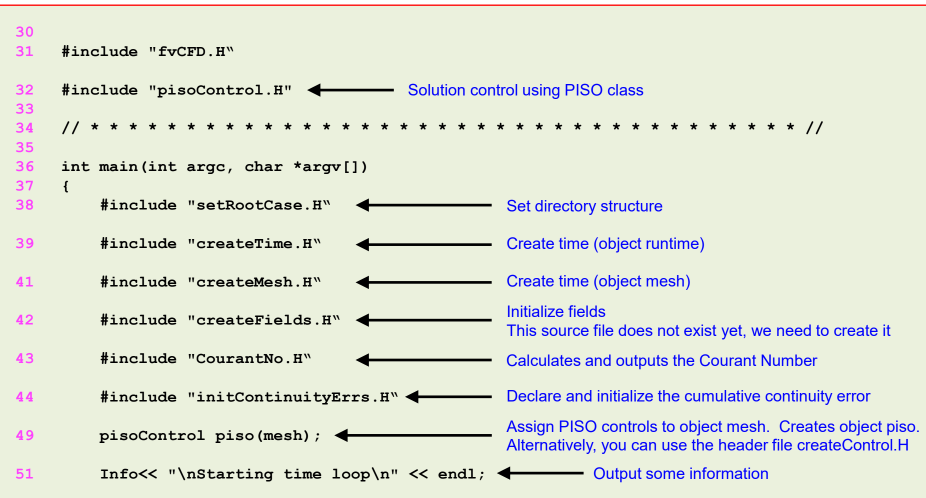
The scratchFoam directory contains the source code of the solver.

- scratchFoam. C: contains the starting point to implement the new application.
- createFields. H: in this file we declare all the field variables and initializes the solution. This file does not exist at this point, we will create it later.
- The Make directory contains compilation instructions.
  - Make/files: names all the source files (.C), it specifies the name of the solver and location of the output file.
  - Make/options: specifies directories to search for include files and libraries to link the solver against.
- To compile the new application, we use the command wmake.

- Open the file scratchFoam.C using your favorite text editor, we will use gedit.
- At this point you should have this file that does not do anything. We need to add the statements
  to create a working applications.
- This is the starting point for new applications.

```
This header is extremely important, it will add all the class
30
                                        declarations needed to access mesh, fields, tensor algebra, fvm/fvc
     #include "fvCFD.H"
31
                                        operators, time, parallel communication, linear algebra, and so on.
32
33
34
35
     int main(int argc, char *argv[])
36
37
         #include "setRootCase.H"
         #include "createTime.H"
38
39
40
41
42
         Info<< nl << "ExecutionTime = " << runTime.elapsedCpuTime() << " s"</pre>
              << " ClockTime = " << runTime.elapsedClockTime() << " s"</pre>
43
44
              << nl << endl;
45
46
         Info<< "End\n" << endl;</pre>
47
         return 0;
48
49
50
```

- Stating from line 31, add the following statements.
- We are going to use the PISO control options, even if we do not have to deal with velocitypressure coupling.



 We are going to use the PISO control options, even if we do not have to deal with velocitypressure coupling.

```
53
            55
                Info<< "Time = " << runTime.timeName() << n1 << endl;</pre>
56
57
                #include "CourantNo.H" ◀

    Calculates and outputs the Courant Number

58
59
                while (piso.correct()) ←
                                                           PISO options (correct loop)
60
                                                                              PISO options (non orthogonal corrections
                   while (piso.correctNonOrthogonal()) 
61
                                                                              loop)
62
                                                                              Create object TEqn.
63
                         fvScalarMatrix Tegn <
                                                                              fvScalarMatrix is a scalar instance of fvMatrix
64
                                                                              Model equation (convection-diffusion)
                                fvm::ddt(T)
65
                                                                              We need to create the scalar field T, vector
                                + fvm::div(phi, T)
66
                                                                              field U (used in phi or face fluxes), and the
                                - fvm::laplacian(DT)
67
                                                                              constant DT.
68
                         );
                                                                              We will declare these variables in the
                                                                              createFields.H header file.
                                                                              In the dictionary fvSchemes, you will need to
                             \frac{\partial T}{\partial t} + \nabla \cdot (\phi T) - \nabla \cdot (\Gamma \nabla T) = 0
                                                                              define how to compute the differential
                                                                              operators, that is,
                                                                                     ddt(T)
70
                         TEqn.solve();
                                                                                     div(phi, T)
                                                                                     laplacian(DT, T)
                                   Solve TEan
72
                                                                              You will need to define the linear solver for T
                                   At this point the object
                                                                              in the dictionary fvSolution
                                   TEgn holds the solution.
```

 We are going to use the PISO control options, even if we do not have to deal with velocitypressure coupling.

```
compute the CPU time of each iteration,
                 #include "continuityErrs.H" Computes continuity errors
74
                                                                                                                     Write CPU time at the end of the time loop.
76
                 runTime.write();  Write the solution in the runtime folder
                                                   It will write the data requested in the file createFields.H
78
                          At this point we are outside of the time loop
79
80
81
           Info<< nl << "ExecutionTime = " << runTime.elapsedCpuTime() << " s"</pre>
82
83
                 << " ClockTime = " << runTime.elapsedClockTime() << " s"</pre>
84
                 << nl << endl;
85
                                                                                                                          add the same
86
           Info<< "End\n" << endl; ← Output this message
87
88
                                End of the program (exit status).
            return 0;
89
      }
                                    If everything went fine, the program should return 0.
90
                                    To now the return value, type in the terminal,
                                     $> echo $?
91
```

• Let us create the file createFields.H, type in the terminal,

```
1. $> touch createFields.H
```

Now open the file with your favorite editor, and start to add the following information,

```
Info<< "Reading field T\n" << endl;</pre>
                                    Create scalar field T
          volScalarField T
               IOobject
                                    Create object for input/output operations
                                     Name of the dictionary file to read/write
                   runTime.timeName(), runtime directory
9
                                                                                 Object registry
                   mesh,
10
                   IOobject::MUST READ,
                                                             Read the dictionary in the runtime directory
                    IOobject::AUTO WRITE
11
                                                             (MUST READ, and write the value in the runtime
12
               ),
                                                             directory (AUTO WRITE).
13
               mesh Link object to mesh
                                                             If you do not want to write the value, use the option
14
          );
                                                             NO WRITE
```

- Remember, in the file <code>createFields.H</code>, we declare all the variables (or fields) that we will use (U and T in this case).
- The dimensions of the fields are defined in the input dictionaries, you also have the option to define the dimensions in the source code.
- You can also define the fields directly in the source file <code>scratchFoam.C</code>, but it is good practice to do it in the header. This improves code readability.

```
17
      Info<< "Reading field U\n" << endl;</pre>
18
          volVectorField U

    Create vector field U

19
20
21
               IOobject
22
                                                                         Name of the dictionary file to read/write
23
                    "U".
24
                   runTime.timeName(),
25
                   mesh,
26
                   IOobject::MUST READ,
27
                   IOobject::AUTO WRITE
28
               ),
29
               mesh
30
          );
31
```

- We also need to declare the constant DT, that is read from the dictionary transportProperties.
- The dimensions are defined in the input dictionary.

```
33
       Info<< "Reading transportProperties\n" << endl;</pre>
34
35
          IOdictionary transportProperties ◀
                                                                       Create object transportProperties used to
                                                                       read data
36
37
               IOobject
                                                                       Name of the input dictionary
38
                    "transportProperties",
39
                                                                       Location of the input dictionary, in this case
                    runTime.constant(),
40
                                                                       is located in the directory constant
41
                    mesh,
                    IOobject::MUST READ IF MODIFIED, 	←
                                                                       Re-read data if it is modified
42
43
                    IOobject::NO WRITE
44
                                                                       Do not write anything in the dictionary
          );
45
46
47
48
          Info<< "Reading diffusivity DT\n" << endl;</pre>
49
                                                                       Create scalar DT (diffusion coefficient)
          dimensionedScalar DT <
50
51
                                                                       Access value of DT in the object
52
               transportProperties.lookup("DT")
                                                                       transportProperties
53
          );
54
                                                                       Creates and initializes the relative face-
55
          #include "createPhi.H"
                                                                       flux field phi.
56
```

At this point, we are ready to compile. Type in the terminal,

- If everything went fine, you should have a working solver named scratchFoam.
- If you are feeling lazy or you can not fix the compilation errors, you will find the source code in the directory,
  - \$PTOFC/101programming/applications/solvers/scratchFoam
- You will find a case ready to run in the directory,

\$PTOFC/101programming/applications/solvers/scratchFoam/test\_case

At this point, we are all familiar with the convection-diffusion equation and OpenFOAM®, so you know how to run the case. Do your magic.

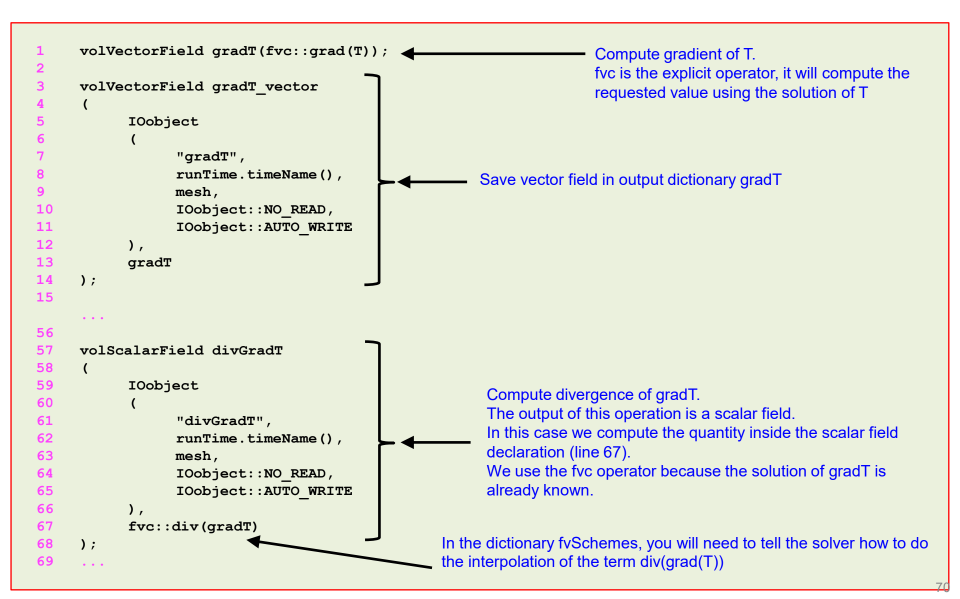
- Let us now add a little bit more complexity, a non-uniform initialization of the scalar field T.
- Remember codeStream? Well, we just need to proceed in a similar way.
- As you will see, initializing directly in the source code of the solver is more intrusive than using codeStream in the input dicitionaries.
- It also requires recompiling the application.
- Add the following statements to the createFields.H file, recompile and run again the test case.

```
16
17
         18
             const scalar x = mesh.C()[i][0];
19
                                                             Access cell center coordinates.
             const scalar y = mesh.C()[i][1];
20
                                                             In this case y and z coordinates are not used.
21
             const scalar z = mesh.C()[i][2];
22
23
             if (0.3 < x \&\& x < 0.7)
                                                 Conditional structure
24
25
                     T[i] = 1.;
26
27
                                                Write field T. As the file createFields.H is outside the time loop
         T.write();
28
                                                the value is saved in the time directory 0
```

- Let us compute a few extra fields. We are going to compute the gradient, divergence, and Laplacian of T.
- We are going to compute these fields in an explicit way, that is, after finding the solution of T.
- Therefore we are going to use the operator fvc.
- Add the following statements to the source code of the solver (scratchFoam.C),

- Recompile the solver and rerun the test case.
- The solver will complain, try to figure out what is the problem (you are missing some information in the fvSchemes dictionary).

Let us talk about the file write.H,



# Roadmap

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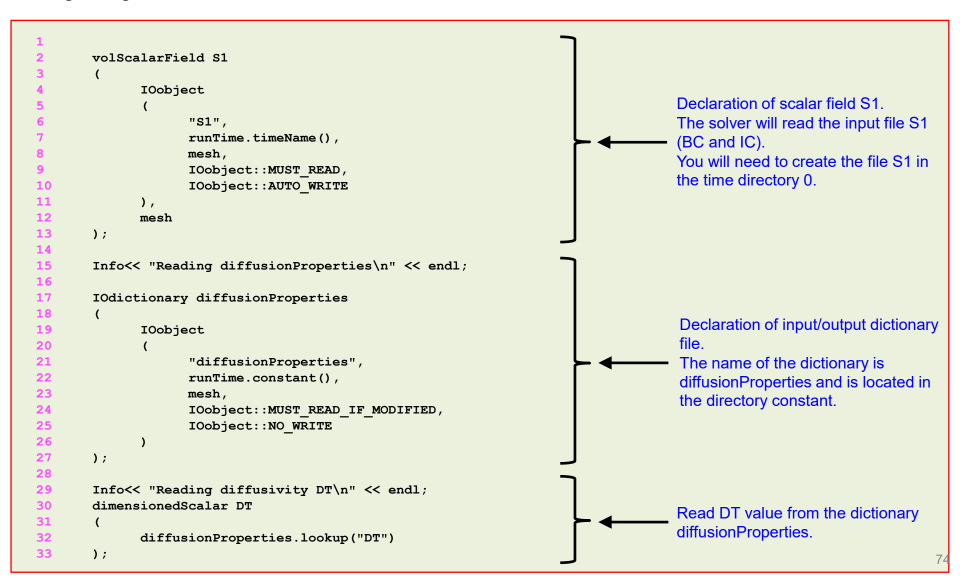
- Let us modify a solver, we will work with **icoFoam**.
- We will add a passive scalar (convection-diffusion equation).
- At this point, you can work in any directory. But we recommend you to work in your OpenFOAM® user directory, type in the terminal,
  - 1. | \$> cd \$WM\_PROJECT\_USER\_DIR/run
- Let us clone the original solver, type in the terminal,
  - \$> cp -r \$FOAM\_APP/solvers/incompressible/icoFoam/ my\_icoFoam
     \$> cd my\_icoFoam
- At this point, we are ready to modify the solver.

Open the file icoFoam. C using your favorite editor and add the new equation in lines 115-120,

```
111
                       U = HbyA - rAU*fvc::grad(p);
112
                       U.correctBoundaryConditions();
                                                                         Scalar transport equation.
113
                  }
                                                                         The name of the scalar is S1.
114
                                                                         We need to declare it in the
                                                                         createFields.H file.
116
                  solve
                                                                         We also need to read the coefficient DT.
117
                                                                         In the dictionary fvSchemes, you will need
118
                       fvm::ddt(S1)
                                                                         to define how to compute the differential
119
                       + fvm::div(phi, S1)
                                                                         operators, that is,
                        - fvm::laplacian(DT, S1)
120
                                                                               ddt(S1)
121
                  );
                                                                               div(phi, S1)
                                                                               laplacian(DT, S1)
122
                                                                         You will need to define the linear solver
123
                                                                         for S1 in the dictionary fvSolution
124
                  runTime.write();
```

 As the passive scalar equation depends on the vector field U, we need to add this equation after solving U.

 Open the file createFields. H using your favorite editor and add the following lines at the beginning of the file,



- Those are all the modifications we need to do.
- But before compiling the new solver, we need to modify the compilation instructions.



• Using your favorite editor, open the file Make/files,

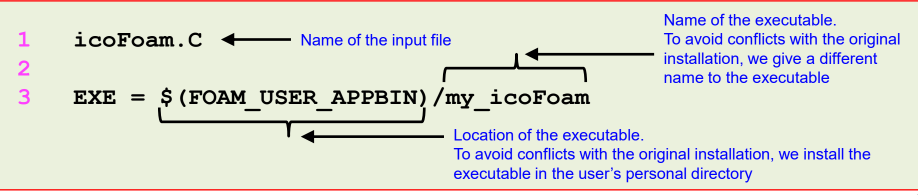
#### **Original file**

1 icoFoam.C

2

3 EXE = \$(FOAM APPBIN)/icoFoam

#### **Modified file**



At this point we are ready to compile, type in the terminal,

- If everything went fine, you should have a working solver named my\_icoFoam.
- If you are feeling lazy or you can not fix the compilation errors, you will find the source code in the directory,
  - \$PTOFC/101programming/applications/solvers/my\_icoFoam

You will find a case ready to run in the directory,

\$PTOFC/101programming/applications/solvers/my\_icoFoam/test\_case

#### Running the case

- This case is ready to run, the input files are located in the directory
   \$PTOFC/101programming/applications/solvers/my\_icoFoam/test\_case
- To run the case, type in the terminal,

\$> paraFoam

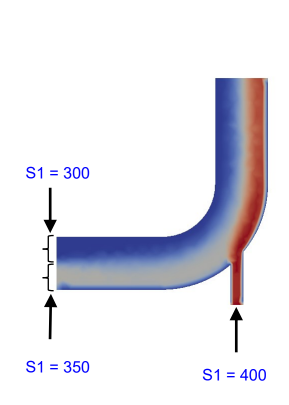
```
    $> foamCleanTutorials
    $> fluentMeshToFoam ../../../meshes_and_geometries/fluent_elbow2d_1/ascii.msh
    $> my_icoFoam | tee log
```

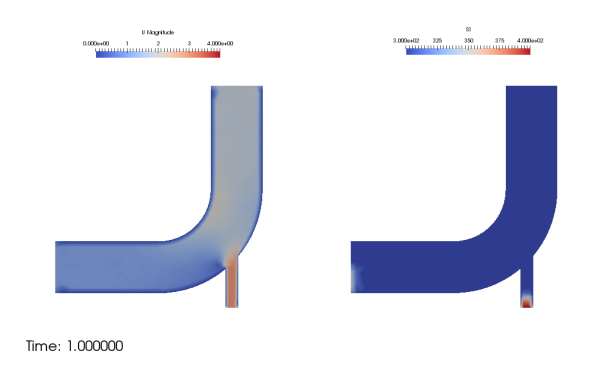
- Remember, you will need to create the file 0/S1 (boundary conditions and initial conditions for the new scalar).
- You will also need to create the input dictionary constant/diffusionProperties, from this dictionary we will read the diffusion coefficient value.
- Finally, remember to update the files <code>system/fvSchemes</code> and <code>system/fvSolution</code> to take into account the new equation.



#### Running the case

If everything went fine, you should get something like this





S1 inlet values



Visualization of velocity magnitude and passive scalar S1 www.wolfdynamics.com/wiki/BCIC/2delbow S1