#### Dam break free surface flow



# Physical and numerical side of the problem:

- In this case we are going to use the volume of fluid (VOF) method.
- This method solves the incompressible Navier-Stokes equations plus an additional equation to track the phases (free surface location).
- As this is a multiphase case, we need to define the physical properties for each phase involved (viscosity, density and surface tension).
- The working fluids are water and air.
- Additionally, we need to define the gravity vector and initialize the two flows.
- This is a three-dimensional and unsteady case.
- The details of the case setup can be found in the following reference:

A Volume-of-Fluid Based Simulation Method for Wave Impact Problems. Journal of Computational Physics 206(1):363-393. June, 2005.

#### Workflow of the case



#### At the end of the day, you should get something like this





Initial conditions – Coarse mesh

Solution at Time = 1 second – Coarse mesh

#### **VOF Fraction (Free surface tracking) – Very fine mesh**

http://www.wolfdynamics.com/validations/3d\_db/dbreak.gif



3D dam-break simulation using OpenFOAM 4.x • Let us run this case. Go to the directory:

# \$PTOFC/3d\_damBreak

- \$PTOFC is pointing to the directory where you extracted the training material.
- In the case directory, you will find the README.FIRST file. In this file, you will find the general instructions of how to run the case. In this file, you might also find some additional comments.
- You will also find a few additional files (or scripts) with the extension .sh, namely, run\_all.sh, run\_mesh.sh, run\_sampling.sh, run\_solver.sh, and so on. These files can be used to run the case automatically by typing in the terminal, for example, sh run\_solver.
- We highly recommend you to open the README.FIRST file and type the commands in the terminal, in this way, you will get used with the command line interface and OpenFOAM® commands.
- If you are already comfortable with OpenFOAM®, use the automatic scripts to run the cases.

#### What are we going to do?

- We will use this case to introduce the multiphase solver interFoam.
- interFoam is a solver for 2 incompressible, isothermal immiscible fluids using a VOF (volume of fluid) phase-fraction based interface capturing approach
- We will define the physical properties of two phases and we are going to initialize these phases.
- We will define the gravity vector in the dictionary g.
- After finding the solution, we will visualize the results. This is an unsteady case so now we are going to see things moving.
- We are going to briefly address how to post-process multiphase flows.
- We are going to generate the mesh using snappyHexMesh, but for the purpose of this tutorial we are not going to discuss the dictionaries.
- Remember, different solvers have different input dictionaries.

The constant directory

- In this directory, we will find the following compulsory dictionary files:
  - g
  - transportProperties
  - turbulenceProperties
- g contains the definition of the gravity vector.
- *transportProperties* contains the definition of the physical properties of each phase.
- turbulenceProperties contains the definition of the turbulence model to use.

#### The g dictionary file

8	FoamFile	
9	{	
10	version	2.0;
11	format	ascii;
12	class	uniformDimensionedVectorField;
13	location	"constant";
14	object	g;
15	}	
17		
18	dimensions	[0 1 -2 0 0 0 0];
19	value	(0 0 -9.81);

- This dictionary file is located in the directory constant.
- For multiphase flows, this dictionary is compulsory.
- In this dictionary we define the gravity vector (line 19).
- Pay attention to the **class** type (line 12).

```
The transportProperties dictionary file
                 |≡]
           Primary phase
17
     phases (water air);
18
19
     water
20
     {
21
         transportModel Newtonian;
22
         nu
                        [0 2 -1 0 0 0 0] 1e-06;
23
         rho
                        [1 -3 0 0 0 0 0] 1000;
24
     }
25
26
     air
27
     {
28
         transportModel Newtonian;
29
         nu
                        [0 2 -1 0 0 0 0] 1.48e-05;
30
         rho
                        [1 -3 0 0 0 0 0] 1;
31
     ł
32
33
                    [1 0 -2 0 0 0 0] 0.07;
     sigma
```

- This dictionary file is located in the directory constant.
- We first define the name of the phases (line 17). In this case we are defining the names water and **air**. The first entry in this list is the primary phase (water).
- The name of the primary phase is the one you will use to initialize the solution.
- The name of the phases is given by the user.
- In this file we set the kinematic viscosity (**nu**), density (**rho**) and transport model (transportModel) of the phases.
- We also define the surface tension (**sigma**).

The turbulenceProperties dictionary file

- In this dictionary file we select what model we would like to use (laminar or turbulent).
- This dictionary is compulsory.
- In this case we use a RANS turbulence model (kEpsilon).

```
17
      simulationType
                          RAS;
18
19
      RAS
20
      {
21
           RASModel kEpsilon;
22
23
           turbulence on;
24
25
          printCoeffs on;
26
      }
```

#### The 0 directory

- In this directory, we will find the dictionary files that contain the boundary and initial conditions for all the primitive variables.
- As we are solving the incompressible RANS Navier-Stokes equations using the VOF method, we will find the following field files:
  - *alpha.water* (volume fraction of water phase)
  - p\_rgh (pressure field minus hydrostatic component)
  - U (velocity field)

nut

•

- k (turbulent kinetic energy field)
- epsilon (rate of dissipation of turbulence energy field)
  - (turbulence viscosity field)

#### The file 0/alpha.water

17	dimensions	[0 0 0 0 0 0];
18		
19	internalField	uniform 0;
20		
21	boundaryField	
22	{	
23	front	
24	{	
25	type	<pre>zeroGradient;</pre>
26	}	
27	back	
28	{	
29	type	zeroGradient;
30	}	
31	leit	
32	1	
33	туре	zeroGradient;
34	} 	
35	right	
36	1	
20	туре	zerogradient;
20	} bottom	
40	1 DOLLOIN	
40	1	zeroGradient.
42	ı cype	Zerogradient,
43	top	
44	1	
45		inletOutlet:
46	inletV	alue uniform 0:
47	value	uniform 0:
48	}	······································
49	stlSurface	
50	{	
51	type	wall;
52	}	
53		
54	}	

- This file contains the boundary and initial conditions for the non-dimensional scalar field **alpha.water**
- This file is named *alpha.water*, because the primary phase is water (we defined the primary phase in the *transportProperties* dictionary).
- Initially, this field is initialize as 0 in the whole domain (line 19). This means that there is no water in the domain at time 0. Later, we will initialize the water column and this file will be overwritten with a non-uniform field for the **internalField**.
- For the front, back, left, right, bottom and stlSurface patches we are using a zeroGradient boundary condition (we are just extrapolating the internal values to the boundary face).
- For the top patch we are using an inletOutlet boundary condition. This boundary condition avoids backflow into the domain. If the flow is going out it will use zeroGradient and if the flow is coming back it will assign the value set in the keyword inletValue (line 46).

#### **The file** 0/p\_rgh

17	dimensions	[1 -1 -2 0 0 0 0];
18		
19	internalField	uniform 0;
20		
21	boundaryField	
22	{	
23	front	
24	۲. ۱	
25	type	fixedFluxPressure;
26	value	uniform 0;
27	}	
28	back	
33	left	
38	right	
43	bottom	
48	top	
49	{	
50	type	totalPressure;
51	p0	uniform 0;
52	υ	U;
53	phi	phi;
54	rho	rho;
55	psi	none;
56	gamma	1;
57	value	uniform 0;
58	}	
59	stlSurface	
60	{	
61	type	fixedFluxPressure;
62	value	uniform 0;
63	}	
64		
65	}	

- This file contains the boundary and initial conditions for the dimensional scalar field p\_rgh. The dimensions of this field are given in Pascal (line 17)
- This scalar field contains the value of the static pressure field minus the hydrostatic component.
- This field is initialize as 0 in the whole domain (line 19).
- For the front, back, left, right, bottom and stlSurface patches we are using a fixedFluxPressure boundary condition (refer to the source code or doxygen documentation to know more about this boundary condition).
- For the **top** patch we are using the **totalPressure** boundary condition (refer to the source code or doxygen documentation to know more about this boundary condition).

The file 0/U

17	dimensions	[0 -1 -1 0 0 0 0];
18		
19	internalField	uniform (0 0 0);
20		
21	boundaryField	
22	{	
23	front	
24	{	
25	type	fixedValue;
26	value	uniform (0 0 0);
27	}	
28	back	
33	left	
38	right	
43	bottom	
48	top	
49	{	
50	type	<pre>pressureInletOutletVelocity;</pre>
51	value	uniform (0 0 0);
52	}	
53	stlSurface	
54	{	
55	type	fixedValue;
56	value	uniform (0 0 0);
57	}	
58		
59	}	
	,	

**|**≞

- This file contains the boundary and initial conditions for the dimensional vector field **U**.
- We are using uniform initial conditions and the numerical value is (0 0 0) (keyword internalField in line 19).
- The front, back, left, right, bottom and stlSurface patches are no-slip walls, therefore we impose a fixedValue boundary condition with a value of (0 0 0) at the wall.
- For the **top** patch we are using the **pressureInIterOutletVelocity** boundary condition (refer to the source code or doxygen documentation to know more about this boundary condition).

The file 0/k

17	dimensions	[0 2 -2 0 0 0 0];
18		
19	internalField	uniform 0.1;
20		
21	boundaryField	
22	{	
23	"(front ba	ck left right bottom stlSurface)"
24	{	
25	type	kqRWallFunction;
26	value	<pre>\$internalField;</pre>
27	}	
28		
29	top	
30	{	
31	type	inletOutlet;
32	inlet	Value \$internalField;
33	value	<pre>\$internalField;</pre>
34	}	
35		
36	}	

**|**≞

- This file contains the boundary and initial conditions for the dimensional scalar field **k**.
- This scalar (turbulent kinetic energy), is related to the turbulence model.
- This field is initialize as 0.1 in the whole domain, and all the boundary patches take the same value (**\$internalField**).
- For the front, back, left, right, bottom and stlSurface patches we are using a kqRWallFunction boundary condition, which applies a wall function at the walls (refer to the source code or doxygen documentation to know more about this boundary condition).
- For the **top** patch we are using the **inletOutlet** boundary condition, this boundary condition handles backflow (refer to the source code or doxygen documentation to know more about this boundary condition).
- We will deal with turbulence modeling later.

#### The file 0/epsilon

17	dimensions	[0 2 -3 0 0 0 0];
18		
19	internalField	uniform 0.1;
20		
21	boundaryField	
22	{	
23	"(front bac	k left right bottom stlSurface)"
24	{	
25	type	<pre>epsilonWallFunction;</pre>
26	value	<pre>\$internalField;</pre>
27	}	
28		
29	top	
30	{	
31	type	inletOutlet;
32	inletVa	lue \$internalField;
33	value	<pre>\$internalField;</pre>
34	}	
35		
36	}	

- This file contains the boundary and initial conditions for the dimensional scalar field **epsilon**.
- This scalar (rate of dissipation of turbulence energy), is related to the turbulence model.
- This field is initialize as 0.1 in the whole domain, and all the boundary patches take the same value (\$internalField).
- For the front, back, left, right, bottom and stlSurface patches we are using a epsilonWallFunction boundary condition, which applies a wall function at the walls (refer to the source code or doxygen documentation to know more about this boundary condition).
- For the **top** patch we are using the **inletOutlet** boundary condition, this boundary condition handles backflow (refer to the source code or doxygen documentation to know more about this boundary condition).
- We will deal with turbulence modeling later.

#### The file 0/nut

17	dimensions	[0 2 -1 0 0 0 0];
18		
19	internalField	uniform 0;
20		
21	boundaryField	
22	{	
23	"(front bac	<pre>k left right bottom stlSurface)</pre>
24	{	
25	type	<pre>nutkWallFunction;</pre>
26	value	<pre>\$internalField;</pre>
27	}	
28		
29	top	
30	ł	
31	type	calculated;
32	value	<pre>\$internalField;;</pre>
33	}	
34		
35	}	

**|**≞

- This file contains the boundary and initial conditions for the dimensional scalar field **nut**.
- This scalar (turbulent viscosity), is related to the turbulence model.
- This field is initialize as 0 in the whole domain, and all the boundary patches take the same value (\$internalField).
- For the front, back, left, right, bottom and stlSurface patches we are using a nutkWallFunction boundary condition, which applies a wall function at the walls (refer to the source code or doxygen documentation to know more about this boundary condition).
- For the **top** patch we are using the **calculated** boundary condition, this boundary condition computes the value of nut from k and epsilon (refer to the source code or doxygen documentation to know more about this boundary condition).
- We will deal with turbulence modeling later.

The system directory

- The **system** directory consists of the following compulsory dictionary files:
  - controlDict
  - fvSchemes
  - fvSolution
- *controlDict* contains general instructions on how to run the case.
- *fvSchemes* contains instructions for the discretization schemes that will be used for the different terms in the equations.
- fvSolution contains instructions on how to solve each discretized linear equation system.

#### The controlDict dictionary

17 application interFoam; 18 19 startFrom startTime; 20 21 startTime 0: 22 23 stopAt endTime; 24 25 endTime 8: 26 27 deltaT 0.0001; 28 29 writeControl adjustableRunTime; 30 31 writeInterval 0.02; 32 33 purgeWrite 0; 34 35 writeFormat ascii; 36 37 writePrecision 8; 38 39 writeCompression uncompressed; 40 41 timeFormat general; 42 43 timePrecision 8: 44 45 runTimeModifiable yes; 46 47 adjustTimeStep yes; 48 49 0.5; maxCo 50 maxAlphaCo 0.5; 51 maxDeltaT 0.01;

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- This case starts from time 0 (**startTime**), and it will run up to 8 seconds (**endTime**).
- The initial time step of the simulation is 0.0001 seconds (deltaT).
- It will write the solution every 0.02 seconds (writeInterval) of simulation time (runTime). It will automatically adjust the time step (adjustableRunTime), in order to save the solution at the precise write interval.
- It will keep all the solution directories (purgeWrite).
- It will save the solution in ascii format (writeFormat).
- The write precision is 8 digits (**writePrecision**). It will only save eight digits in the output files.
- And as the option **runTimeModifiable** is on, we can modify all these entries while we are running the simulation.
- In line 47 we turn on the option **adjustTimeStep**. This option will automatically adjust the time step to achieve the maximum desired courant number (lines 49-50). We also set a maximum time step in line 51.
- Remember, the first time step of the simulation is done using the value set in line 27 and then it is automatically scaled to achieve the desired maximum values (lines 49-51).

#### The controlDict dictionary



- Let us take a look at the functionObjects definitions.
- In lines 60-76 we define the fieldMinMax functionObject which computes the minimum and maximum values of the field variables (p p\_rgh U alpha.water k epsilon).

#### The controlDict dictionary

55	functions
56	{
81	water_in_domain
82	t in the second s
83	type volkegion;
84	functionObjectLibs ("libiteldFunctionObjects.so"),
85	enabled true;
07	anahlad turus.
87	enabled true;
88	
00	writeControl timeSter:
90	writeInterval 1:
92	WIICEINCEIVAI I,
93	log true:
94	109 0140,
95	regionType all:
96	
97	operation volIntegrate:
98	fields
99	(
100	alpha.water
101	);
102	}
144	};

- Let us take a look at the **functionObjects** definitions.
- In lines 81-102 we define the **volRegion functionObject** which computes the volume integral (**volIntegrate**) of the field variable **alpha.water** in all the domain.
- Basically, we are monitoring the quantity of water in the domain.

#### The controlDict dictionary

```
55
        functions
56
        ł
107
         probes1
108
         ſ
109
              type
                                probes;
110
              functionObjectLibs ("libsampling.so");
111
112
             pobeLocations
113
114
                 (0.82450002 \ 0 \ 0.021)
115
                 (0.82450002 \ 0 \ 0.061)
116
                 (0.82450002 \ 0 \ 0.101)
117
                 (0.82450002 \ 0 \ 0.141)
118
                 (0.8035 \ 0 \ 0.161)
119
                 (0.7635 \ 0 \ 0.161)
120
                 (0.7235 \ 0 \ 0.161)
121
                 (0.6835 \ 0 \ 0.161)
122
             );
123
124
            fields
125
             (
126
                 p p_rgh
127
            );
128
129
              writeControl
                               timeStep;
130
              writeInterval 1;
131
        ł
144
        };
```

- Let us take a look at the functionObjects definitions.
- In lines 107-131 we define the **probes functionObject** which sample the selected fields (lines 124-127) at the selected locations (lines 112-122).
- This sampling is done on-the-fly. All the information sample by this functionObject is saved in the directory ./postProcessing/probes1
- As we are sampling starting from time 0, the sampled data will be located in the directory:

#### postProcessing/probes1/0

 Feel free to open the files located in the directory postProcessing/probes1/0 using your favorite text editor.



#### 

#### The controlDict dictionary

55	functions
56	{
135	yplus
136	{
137	type yPlus;
138	<pre>functionObjectLibs ("libutilityFunctionObjects.so ");</pre>
139	enabled true;
140	<pre>writeControl outputTime;</pre>
141	}
144	};

- Let us take a look at the **functionObjects** definitions.
- In lines 135-141 we define the **yplus functionObject** which computes the yplus value.
- This quantity is related to the turbulence modeling.
- This **functionObject** will save the yplus field in the solution directories with the same saving frequency as the solution (line 140).
- It will also save the minimum, maximum and mean values of yplus in the directory:

#### postProcessing/yplus

#### The fvSchemes dictionary

```
17
      ddtSchemes
18
      {
19
          default
                           Euler:
21
      }
22
23
      gradSchemes
24
      {
25
          default.
                           Gauss linear;
26
          grad(U)
                           cellLimited Gauss linear 1;
27
      }
28
29
      divSchemes
30
      {
31
          div(rhoPhi,U) Gauss linearUpwindV grad(U);
32
          div(phi,alpha) Gauss vanLeer;
33
          div(phirb,alpha) Gauss linear;
35
          div(phi,k) Gauss upwind;
36
          div(phi,epsilon) Gauss upwind;
          div(((rho*nuEff)*dev2(T(grad(U))))) Gauss linear;
37
38
      ł
39
40
      laplacianSchemes
41
      {
42
          default
                           Gauss linear corrected;
43
      ł
44
45
      interpolationSchemes
46
      {
47
          default
                           linear:
48
      }
49
50
      snGradSchemes
51
      {
52
          default
                           corrected;
53
      ı
```

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- In this case, for time discretization (**ddtSchemes**) we are using the **Euler** method.
- For gradient discretization (gradSchemes) we are using the Gauss linear as the default method and slope limiters (cellLimited) for the velocity gradient or grad(U).
- For the discretization of the convective terms (divSchemes) we are using linearUpwindV interpolation method for the term div(rhoPhi,U).
- For the term **div(phi,alpha)** we are using **vanLeer** interpolation. For the term **div(phirb,alpha)** we are using **linear** interpolation. These terms are related to the volume fraction equation.
- For the terms **div(phi,alpha)** and **div(phi,alpha)** we are using upwind (these terms are related to the turbulence modeling).
- For the term div(((rho\*nuEff)\*dev2(T(grad(U))))) we are using linear interpolation (this term is related to the turbulence modeling).
- For the discretization of the Laplacian (laplacianSchemes and snGradSchemes) we are using the Gauss linear corrected method
- In overall, this method is second order accurate but a little bit diffusive. Remember, at the end of the day we want a solution that is second order accurate.

#### The fvSolution dictionary

```
17
      solvers
18
      ſ
19
           "alpha.water.*"
20
          {
21
               nAlphaCorr
                                3;
22
               nAlphaSubCycles 1;
23
               cAlpha
                                1;
24
25
               MULESCorr
                                yes;
26
               nLimiterIter
                                10;
27
28
               solver
                                smoothSolver;
29
               smoother
                                symGaussSeidel;
30
               tolerance
                                1e-8;
31
               relTol
                                0;
32
          }
33
34
          "(pcorr|pcorrFinal)"
35
           {
36
               solver
                                PCG;
37
               preconditioner DIC;
38
               tolerance
                                1e-8;
39
               relTol
                                0;
40
          }
41
42
          p_rgh
43
           {
44
                                PCG;
               solver
45
               preconditioner
                               DIC;
46
               tolerance
                                1e-06;
47
               relTol
                                0.01;
48
               minIter
                                1;
49
          }
```

- To solve the volume fraction or **alpha.water** (lines 19-32) we are using the **smoothSolver** method.
- In line 25 we turn on the semi-implicit method MULES. The keyword nLimiterIter controls the number of MULES iterations over the limiter.
- To have more stability it is possible to increase the number of loops and corrections used to solve **alpha.water** (lines 21-22).
- The keyword **cAlpha** (line 23) controls the sharpness of the interface (1 is usually fine for most cases).
- In lines 34-40 we setup the solver for **pcorr** and **pcorrFinal** (pressure correction).
- In this case **pcorr** is solved only one time at the beginning of the computation.
- In lines 42-49 we setup the solver for **p\_rgh**.
- The keyword **miniter** (line 48), means that the linear solver will do at least one iteration.

#### The fvSolution dictionary

51		p_rghFinal	
52		{	
53		<pre>\$p_rgh;</pre>	
54		relTol	0;
55		minIter	1;
56		}	
57			
58		"(U UFinal)"	
59		{	
60		solver	PBiCGStab;
61		Preconditioner	DILU;
62		tolerance	1e-08;
63		relTol	0;
72		}	
73			
74		"(k epsilon).*"	
75		{	
76		solver	PBiCGStab;
77		Preconditioner	DILU;
78		tolerance	1e-08;
79		relTol	0;
80		}	
81	}		
82			

- In lines 51-56 we setup the solver for p\_rghFinal. This correspond to the last iteration in the loop (we can use a tighter convergence criteria to get more accuracy without increasing the computational cost)
- In lines 58-72 we setup the solvers for U and UFInal.
- In lines 74-80 we setup the solvers for the turbulent quantities, namely, **k** and **epsilon**.

#### The fvSolution dictionary

```
82
83
      PIMPLE
84
      {
85
          momentumPredictor
                               yes;
86
          nOuterCorrectors
                               1;
87
          nCorrectors
                               3;
88
          nNonOrthogonalCorrectors 1;
89
      }
90
91
      relaxationFactors
92
      ſ
93
          fields
94
          {
               ".*" 1;
95
96
          }
97
          equations
98
          {
              ".*" 1;
99
100
           }
101
       }
102
```

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- In lines 83-89 we setup the entries related to the pressurevelocity coupling method used (PIMPLE in this case). Setting the keyword nOuterCorrectors to 1 is equivalent to running using the PISO method.
- To gain more stability we can increase the number of correctors (lines 87-88), however this will increase the computational cost.
- In lines 91-101 we setup the under relaxation factors related to the **PIMPLE** method. By setting the coefficients to one we are not under-relaxing.
- The option **momentumPredictor** (line 85), is recommended for highly convective flows.

The system directory

- In the **system** directory you will find the following optional dictionary files:
  - decomposeParDict
  - setFieldsDict
- decomposeParDict is read by the utility decomposePar. This dictionary file contains information related to the mesh partitioning. This is used when running in parallel.
- *setFieldsDict* is read by the utility *setFields*. This utility set values on selected cells/faces.

#### The setFieldsDict dictionary

17	defaultFieldValues
18	(
19	volScalarFieldValue alpha.water 0
20	);
21	
22	regions
23	(
24	boxToCell
25	{
26	box (1.992 -10 0) (5 10 0.55);
27	fieldValues
28	(
29	volScalarFieldValue alpha.water 1
30	);
31	}
32	);

- This dictionary file is located in the directory **system**.
- In lines 17-20 we set the default value to be 0 in the whole domain (no water).
- In lines 22-32, we initialize a rectangular region (box) containing water (alpha.water 1).
- In this case, setFields will look for the dictionary file alpha.water and it will overwrite the original values according to the regions defined in setFieldsDict.
- We initialize the water phase because is the primary phase in the dictionary *transportProperties*.
- If you are interested in initializing the vector field U, you can proceed as follows volVectorFieldValue U (0 0 0)



boxToCell region

# The decomposeParDict dictionary

- This dictionary file is located in the directory **system**.
- This dictionary is used to decompose the domain in order to run in parallel.
- The keyword **numberOfSubdomains** (line 17) is used to set the number of cores we want to use in the parallel simulation.
- In this dictionary we also set the decomposition method (line 19).
- Most of the times the scotch method is fine.

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• In this case we set the **numberOfSubdomains** to 4, therefore we will run in parallel using 4 cores.

```
17 numberOfSubdomains 4;
18
19 method scotch;
20
```

When you run in parallel, the solution is saved in the directories **processorN**, where **N** stands for processor number. In this case you will find the following directories with the decomposed mesh and solution: **processor0**, **processor1**, **processor2**, and **processor3**.

#### **Running the case**

- Let us first generate the mesh.
- To generate the mesh will use snappyHexMesh (sHM), do not worry we will talk about sHM tomorrow.
  - 1. \$> foamCleanTutorials
  - **2.** \$> rm -rf 0
  - 3. \$> blockMesh
  - 4. \$> surfaceFeatureExtract
  - 5. \$> snappyHexMesh -overwrite
  - 6. \$> createPatch -dict system/createPatchDict.0 -overwrite
  - 7. \$> createPatch -dict system/createPatchDict.1 -overwrite
  - 8. \$> checkMesh
  - 9. \$> paraFoam

#### Running the case

- Let us run the simulation in parallel using the solver interFoam.
- We will talk more about running in parallel tomorrow
- To run the case, type in the terminal:

Т

- 3. | \$> setFields
- 4. \$> paraFoam
- 5. \$> decomposePar
- 6. \$> mpirun -np 4 interFoam -parallel | tee log.interFoam
- 7. \$> reconstructPar
- 8. \$> paraFoam

#### Running the case

- In steps 1-2 we copy the information of the backup directory 0\_org into the directory
   0. We do this because in the next step the utility setFields will overwrite the file
   0/alpha.water, so it is a good idea to keep a backup.
- In step 3 we initialize the solution using the utility setFields. This utility reads the dictionary *setFieldsDict* located in the system directory.
- In step 4 we visualize the initialization using paraFoam.
- In step 5 we use the utility decomposePar to do the domain decomposition needed to run in parallel.
- In step 6 we run the simulation in parallel. Notice that np means number of
  processors and the value used should be the same number as the one you set in the
  dictionary decomposeParDict.
- If you want to run in serial, type in the terminal: interFoam | tee log
- In step 7 we reconstruct the parallel solution. This step is only needed if you are running in parallel.
- Finally, in step 8 we visualize the solution.

• To plot the sampled data using gnuplot you can proceed as follows. To enter to the gnuplot prompt type in the terminal:

```
1. $> gnuplot
```

• Now that we are inside the gnuplot prompt, we can type,

```
1. set xlabel 'Time (seconds)'
```

- 2. set ylabel 'Water volume integral'
- 3. gnuplot> plot 'postProcessing/water\_in\_domain/0/volRegion.dat' u 1:2 w l title 'Water in domain'
- 4. | set xlabel 'Time (seconds)'

```
5. | set ylabel 'Pressure'
```

6. plot 'SPHERIC\_Test2/case.txt' u 1:2 w l title 'Experiment', 'postProcessing/probes1/0/p' u 1:2 w l title 'Numerical simulation'

```
7. gnuplot> exit
To exit gnuplot
```

• The output of steps 3 and 6 is the following:





alpha.water vs. time

p vs. time (at probe 0)

#### The output screen



volIntegrate() of alpha.water = 0.66459985

Volume integral functionObject

Post-processing multiphase flows in paraFoam

• To visualize the volume fraction, proceed as follows,



Post-processing multiphase flows in paraFoam

• To visualize a surface representing the interface, proceed as follows,



#### Post-processing multiphase flows in paraFoam

• To visualize all the cells representing the water fraction, proceed as follows,



#### Exercises

- Instead of using the boundary condition totalPressure and pressureInletOutletVelocity for the patch top, try to use zeroGradient. Do you get the same results? Any comments?
   (Hint: this combination of boundary conditions will give you an error, read carefully the screen, you will need to add a fix in the file fvSolution)
- Instead of using the boundary condition **fixedFluxPressure** for the walls, try to use **zeroGradient**. Do you get the same results? Any comments?
- Run the simulation in a close domain. Does the volume integral of **alpha.water** remains the same? Why the value is not constant when the domain is open?
- Use a **functionObject** to measure the average pressure at the obstacle.
- How many initialization methods are there available in the dictionary *setFieldsDict*? (Hint: use the banana method)
- Run the simulation using Gauss upwind instead of Gauss vanLeer for the term div(phi,alpha) (fvSchemes).
   Do you get the same quantitative results?
- Run a numerical experiment for **cAlpha** equal to **0**, **1**, and **2**. Do you see any difference in the solution? What about computing time?
- Use the solver GAMG instead of using the solver PCG for the variable p\_rgh. Do you see any difference on the solution or computing time?
- Increase the number of **nOuterCorrector** to 2 and study the output screen. What difference do you see?
- Turn off the MULES corrector (**MULESCorr**). Do you see any difference on the solution or computing time?
- If you set the gravity vector to (0 0 0), what do you think will happen?
- Try to break the solver and identify the cause of the error. You are free to try any kind of setup.