Online Training – Advanced session February 2021

Multiphase flows modeling in OpenFOAM®: Guided Tutorials



From this point on and unless it is strictly necessary, or we introduce new features, we will not explain every single step or comment on the directories and files to be used.

- Guided tutorial 1. Wave impact in a column. VOF Large scale.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT1/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.



- This is the big picture.
- For this case we are going to simulate only one support column (the column in the red square).

Guided tutorial 1 – Wave impact in a column









• And in case you were wondering. Yes, we do have a mesh and a solution using OpenFOAM.

Guided tutorial 1 – Wave impact in a column



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Guided tutorial 1 – Wave impact in a column









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Guided tutorial 1 – Wave impact in a column



Hexahedral mesh.

Left figure: free surface interaction with solid structure. Right figure: forces acting on the column. http://www.wolfdynamics.com/training/mphase/gt1/hexa/ani2.gif

• The incoming wave is a solitary wave.

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- We are not going to address the theory or how to setup the wave*.
- We will address the influence of the cell type on the initialization and solution accuracy and stability, basic case setup, and postprocessing using paraview.



- When conducting free surface simulations, it is extremely important to generate a mesh that is aligned with the free surface level (at least at the free surface level).
- If the cells are not aligned, this may introduce spurious oscillations which might or might not be significant.
- In this case, the polyhedral cells are not aligned with the free surface.
 - This does not mean that the mesh is wrong, we need to assess the final solution.



- When conducting free surface simulations, it is extremely important to generate a mesh that is aligned with the free surface level (at least at the free surface level).
- If the cells are not aligned, this may introduce spurious oscillations which might or might not be significant.
- In this case, the hexahedral cells are aligned with the free surface.
 - The quality of the solution at the free surface is much better than the solution with the polyhedral mesh.
 - Also, the error associated with this mesh is much lower that the error associated with the polyhedral mesh.



- When conducting free surface simulations, it is extremely important to generate a mesh that is aligned with the free surface level (at least at the free surface level).
- If the cells are not aligned, this may introduce spurious oscillations which might or might not be significant.
- Hybrid meshes can be used with no problem.
 - It is important to have in mind that at the free surface level, the cells should be aligned with the free surface.

Guided tutorial 1 – Wave impact in a column

Solution comparison – Hexahedral mesh against polyhedral mesh









Hexahedral mesh http://www.wolfdynamics.com/training/mphase/gt1/hexa/ani1.gif

Polyhedral mesh http://www.wolfdynamics.com/training/mphase/gt1/poly/ani1.gif GT-12

Guided tutorial 1 – Wave impact in a column

Solution comparison – Hexahedral mesh against polyhedral mesh



http://www.wolfdynamics.com/training/mphase/gt1/hexa/ani4.gif



http://www.wolfdynamics.com/training/mphase/gt1/hexa/ani5.gif

Hexahedral mesh



http://www.wolfdynamics.com/training/mphase/gt1/poly/ani4.gif



http://www.wolfdynamics.com/training/mphase/gt1/poly/ani5.gif

Polyhedral mesh

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Solution comparison – Hexahedral mesh against tetrahedral mesh



http://www.wolfdynamics.com/training/mphase/gt1/hexa/ani3.gif



http://www.wolfdynamics.com/training/mphase/gt1/hexa/ani5.gif

http://www.wolfdynamics.com/training/mphase/gt1/tetra/ani3.gif



http://www.wolfdynamics.com/training/mphase/gt1/tetra/ani5.gif

Hexahedral mesh

Guided tutorial 1 – Wave impact in a column

- Turbulence kinetic energy can be over-predicted in incompressible VOF solvers at the phase interface.
- This well know deficiency of the turbulence models can be corrected using multiphase stabilization techniques [1,2].
- If you are using the ESI OpenOFAM version, you use the fvOption multiphaseStabilizedTurbulence that applies corrections to the turbulence kinetic energy equation and turbulence viscosity field.
- At the following link, you can find the release notes addressing this correction,
 - <u>https://www.openfoam.com/news/main-news/openfoam-v1912/solver-and-physics</u>
- Unfortunately, and official version of this correction is not available in the foundation version (as far as we know).
- It is worth mentioning that there are many methods to correct this spurious behavior.

[1] Devolder, B., Rauwoens, P., and Troch, P. (2017). Application of a buoyancy-modified k-w SST turbulence model to simulate wave run-up around a monopile subjected to regular waves using OpenFOAM. Coastal Engineering, 125, 81-94.

[2] Larsen, B.E. and Fuhrman, D.R. (2018). On the over-production of turbulence beneath surface waves in Reynolds-averaged Navier-Stokes models J. Fluid Mech, 853, 419-460

- In this case, we are going to focus our attention on the influence of the cell type on the free surface resolution.
- We are also going to illustrate how to do advanced post-processing in paraview.
- Additionally, we will see how to do basic field initialization.
- We will address details of the numerics in the next tutorials.
- At this point, we are ready to run the simulation.
- To run the turbulent case, go to the **\$TM/multiphase/guided_tutorials/GT1/** directory.
- After running the simulation, launch paraFoam/paraview and load the predefined paraview states located in the directory **paraview**.
 - When asked for the Load State Data File Options, select Choose File Names, and load the file *GT1.OpenFOAM* located in the case directory.
 - In the state file, all the steps to conduct the post-processing have been saved.

- At this point, we are ready to run the simulation.
- To run the turbulent case, go to the **\$TM/multiphase/guided_tutorials/GT1/** directory.
- You can use the script run_all.sh to run the case automatically.
- Type in the terminal:
 - \$> sh run_all.sh
- Feel free to open the file run_all.sh to see all the commands used.
- Or if you prefer, you can insert the commands manually in the terminal window.

Guided tutorial 1 – Wave impact in a column

- To do field initialization, we use the command setFields.
- This command, will overwrite the input dictionaries that we want to initialize, *e.g.*, water volume fraction (*alpha.water*) or velocity field (*U*).
- It is always recommended to keep a backup of the un-initialized fields.
- To initialize the solution, you need to follow these steps,

1. \$> sh run mesh.sh

2. \$> cp 0/alpha.water.org 0/alpha.water

```
3. $> setFields
```

- In step 1, we generate the mesh using the script run_mesh.sh.
 - Remember, in order to initialize the solution, you need to have a valid mesh.
- In step 2, we copy the original files (un-initialized fields or *O/alpha.water.org*) to the field input file to be initialize (*alpha.water* in this case).
 - We do this to keep a backup of the original files, as the file *O/alpha.water* will be overwritten when using setFields.
- In step 3, we initialize the solution using setFields.

Guided tutorial 1 – Wave impact in a column

The setFieldsDict dictionary

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- This dictionary file is located in the directory system.
- The field to be initialized can be a scalar or a vector (or any of the fields supported in OpenFOAM).
- You can also initialize many fields at the same time. In this case, we are only initializing the field *alpha.water*, but if you want, you can also initialize the velocity field (have in mind that this is a vector).
- In lines 17-20 we set the default value of the field in the whole domain (*alpha.water* in this case).
- In lines 22-42, we initialize regions. You can initialize as many regions as you like.
- In lines 24-31 we initialize a rectangular region (cell values).
 - In this case, we are using a box. The input values are the absolute minimum and maximum coordinates of the region.
- In lines 33-40 we apply the same initialization to the face boundaries.
- There are many initialization types implemented in OpenFOAM.
 - Use the banana trick to find the options available or read the source code.



- Guided tutorial 2. Wigley hull Towing tank. VOF Medium scale.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT2/

- 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 that you open the README.FIRST, file and typing the commands in the terminal; in this way, you will get used with the command line interface and OpenFOAM commands.
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Guided tutorial 2 – Wigley Hull – Towing tank



Unsteady simulation Free surface colored by height – VOF CFL = 1 http://www.wolfdynamics.com/training/mphase/gt2/uns1.gif Unsteady simulation Free surface colored by height – VOF CFL = 10 http://www.wolfdynamics.com/training/mphase/gt2/uns2.gif

Guided tutorial 2 – Wigley Hull – Towing tank



Comparison of water level on hull surface

Drag coefficient monitor

Unsteady simulation - VOF CFL =1

Guided tutorial 2 – Wigley Hull – Towing tank





Local-time stepping (LTS) simulation Free surface colored by height – VOF CFL = 0.9 http://www.wolfdynamics.com/training/mphase/gt2/LTS2.gif

Local-time stepping (LTS) simulation Free surface colored by height – VOF CFL = 4 http://www.wolfdynamics.com/training/mphase/gt2/LTS1.gif

Guided tutorial 2 – Wigley Hull – Towing tank





Unsteady simulation – Global time-stepping Free surface colored by height – VOF CFL = 1 http://www.wolfdynamics.com/training/mphase/gt2/ani_sts.gif Unsteady simulation – Global time-stepping Free surface colored by height – VOF CFL = 10 http://www.wolfdynamics.com/training/mphase/gt2/ani_lts.gif

Guided tutorial 2 – Wigley Hull – Towing tank



Unsteady simulation Free surface colored by height – VOF CFL = 1 http://www.wolfdynamics.com/training/mphase/gt2/uns1.gif Local-time stepping (LTS) simulation Free surface colored by height – VOF CFL = 4 http://www.wolfdynamics.com/training/mphase/gt2/LTS1.gif



Case	Time marching method	Max. VOF CFL	Max. flow CFL	LTS smoothing	Max. time step (s)	Clock time - 4 cores (s)
LTS1	Local time-stepping	4	10	0.1	1	11853
LTS2	Local time-stepping	0.9	0.9	0.1	1	9547
UNS1	Global time-stepping - Unsteady	1	10	-	0.1	51961
UNS2	Global time-stepping - Unsteady	10	20	-	0.1	13161



Case	Time marching method	Max. VOF CFL	Max. flow CFL	Max. time step	Clock time - 4 cores (s)
UNS1	Global time-stepping - Unsteady	1	10	0.1	51961
UNS2	Global time-stepping - Unsteady	10	20	0.1	13161

- We are going to use the following solver: interFoam
- The first step is to set the physical properties. In the dictionary constant/transportProperties we defined the phases.
- Go to the directory constant and open the dictionary *transportProperties*.



- The next step is to set the boundary conditions and initial conditions.
- Therefore, in the directory 0 we define the dictionary *alpha.water* that will take the values of the phase water.
- In this case, you will find the directory O_org, here is where we keep a backup of the original files as we are doing field initialization using setFields.
- In the directory 0, you will find the dictionary *p_rgh*, in this dictionary we set the boundary and initial conditions for the pressure field, and the dimensions are in Pascals.
- The turbulence variables values were calculated using an eddy viscosity ratio equal to 1, turbulence intensity equal 5%, and the water properties.
- If you are simulating numerical towing tanks, the setup of the boundary conditions is always the same.
- Feel free to reuse this setup.
- The dictionaries used in this case are standard for the VOF solvers (interFoam family solvers).
- Remember, you should always conduct production runs using a second order discretization scheme



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Patch name	Pressure	Velocity	Turbulence fields	alpha.water
inflow	fixedFluxPressure	fixedValue	fixedValue calculated (nut)	fixedValue
outflow	inletOutlet or zeroGradient	outletPhaseMeanVelocity	inletOutlet or zeroGradient calculated (nut)	variableHeightFlowRate
bottom	symmetry	symmetry	symmetry	symmetry
midplane	symmetry	symmetry	symmetry	symmetry
side	symmetry	symmetry	symmetry	symmetry
top	totalPressure	pressureInletOutletVelocity	inletOutlet or zeroGradient calculated (nut)	inletOutlet
ship	fixedFluxPressure	fixedValue	kqRWallFunction (k) omegaFunction (omega) nutkWallFunction (nut)	zeroGradient

Typical setup of boundary conditions for numerical towing tank simulations

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• OpenFOAM solves the following modified volume fraction convective equation to track the interface between the phases,



- Where a value of $c_{\alpha} = 1$ (cAlpha), is recommended to accurately resolve the sharp interface.
- To solve this equation, OpenFOAM uses the semi-implicit MULES* method.
- The MULES options can be controlled in the *fvSolution* dictionary.

- For interface capturing, OpenFOAM uses,
 - The MULES algorithm (semi-implicit and second order in time) to ensure that the volume fraction (alpha) remains between strict bounds of 0 and 1.
 - The interface compression scheme, based on counter-gradient transport, to maintain sharp interfaces during a simulation.
- At the following link, you can find the release notes related latest developments related to the interface capturing in OpenFOAM 8,
 - <u>https://cfd.direct/openfoam/free-software/multiphase-interface-capturing/</u>
- Among the latest improvements and developments, was the addition of the Piecewise-linear interface calculation (PLIC) family of interpolation schemes.
 - PLIC represents an interface by surface-cuts which split each cell to match the volume fraction of the phase in that cell.
 - The surface-cuts are oriented according to the point field of the local phase fraction.
 - The phase fraction on each cell face (the interpolated value), is then calculated from the amount submerged below the surface-cut.

- In addition, the MPLIC or multicut PLIC method was also introduced in OpenFOAM 8.
- Where a single cut is insufficient, MPLIC performs a topological face-edge-face walk to produce multiple splits of a cell.
- If that is still insufficient, MPLIC decomposes the cell into tetrahedrons on which the cuts are applied.
- The extra cutting carries an additional computational cost but requires no fallback to the interface compression method.
- The PLIC and MPLIC methods are more precise than interface compression for meshes with refinement patterns.
- These methods have the potential of removing or reducing the numerical oscillations that appear on the interface due to refinement patterns.
- These oscillations may cause excessive entrainment of air beneath the hull in naval applications, leading to an inaccurate prediction of forces on the hull.
- (M)PLIC reduces these oscillations for a better prediction of the forces.

- MULES options in the *fvSolution* dictionary.
- The semi-implicit MULES offers significant speed-up and stability over the explicit MULES.



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• Additional notes on the *fvSolution* dictionary.



If you are planning to use large time-steps (CFL number larger than 1), it is recommended to do at least 2 nCorrector and 2 nOuterCorrectors correctors.
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• Additional notes on the *fvSolution* dictionary.



Guided tutorial 2 – Wigley Hull – Towing tank

- Finally, we need to set the discretization schemes
- This is done in the dictionary *fvSchemes*.
- In this dictionary, we set the discretization method for every term appearing in the governing equations.
- Convective terms discretization is set as follows:



- Notice that we are using a high-resolution scheme for the surface tracking (div(phi,alpha)).
- The new notation if OpenFOAM 8, is meant to improve usability with a simple selection of interpolation scheme in the *fvSchemes* file, among many other things.

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• In OpenFOAM 8, the PLIC method can be enabled as follows,



- The basic PLIC method generates a single cut so cannot handle cells in which there are multiple interfaces or where the interface is not fully resolved.
- In those cells, the interpolation reverts to an alternative scheme, typically standard interface compression.
- The PLIC method, with a fallback to interface compression, produces robust solutions and it can run with large time steps.
- The PLIC potentially provides extra accuracy at the cost of an additional computational cost. GT-39

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- In addition, the MPLIC or multicut PLIC method was also introduced in OpenFOAM 8.
- The MPLIC method can be enabled as follows,



- The PLIC and MPLIC methods are more precise than interface compression for meshes with refinement patterns.
- However, the extra cutting carries an additional computational cost.
- Regarding stability, still is a little bit early to give you our opinion.

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 In OpenFOAM 7 and earlier, the discretization of the volume fraction convective term was defined as follows,



• The new notation if OpenFOAM 8, is meant to improve usability with a simple selection of interpolation scheme in the fvSchemes file, among many other things.

div(phi,alpha) Gauss interfaceCompression vanLeer 1;

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- For time discretization we can use an unsteady formulation (Euler in this case).
- This scheme requires setting the time-step, and it should be choosing in such a way that it resolves the mean physics.
- Remember, as the free surface is a strong discontinuity, for stability and good resolution we need to use a CFL less than one for the interface courant.



- Hereafter, we are using what is know as global time stepping, that is, the CFL number is limited by the smallest cell.
- The simulation is time-accurate, but it requires a lot of CPU time to reach a steady state (if it reaches one).

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- A way to accelerate the convergence to steady state, is by using local time stepping* (LTS).
- In LTS, the time-step is manipulated for each individual cell in the mesh, making it as high as possible to enable the simulation to reach steady-state quickly.
- When we use LTS, the transient solution is no longer time accurate.
- The stability and accuracy of the method are driven by the local CFL number of each cell.
- To avoid instabilities caused by sudden changes in the time-step of each cell, the local timestep can be smoothed and damped across the domain.
- Try to avoid having local time-steps that differ by several order of magnitudes.
- To enable LTS, we use the localEuler method.



• LTS in OpenFOAM can be used with any solver that supports the **PISO** or **PIMPLE** loop.

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• In the LTS method, the maximum flow CFL number, maximum interface CFL number, and the smoothing and damping of the solution across the cells, can be controlled in the dictionary *fvSolution*, in the sub-dictionary **PIMPLE**.



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- At this point, we are ready to run the simulation.
- Remember to adjust the numerics according to your physics.
- You can choose between running using global time stepping or unsteady (directory uns) or local time stepping (directory LTS).
- You will find the instructions of how to run the cases in the file *README*.FIRST located in the case directory.

- Guided tutorial 3. Syphon effect (Flushing a tank). VOF Small scale.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT3/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.

Guided tutorial 3 – Syphon effect (flushing a tank)

What will happen in this case?



- In this guided tutorial, we model a tank emptying due to syphon effect.
- This problem is transient in nature.
- We will use the VOF approach to resolve the interface between the two phases.



- In reality, nothing should happen.
- Maybe some wiggles due to numerical instabilities and mesh alignment.

Guided tutorial 3 – Syphon effect (flushing a tank)



http://www.wolfdynamics.com/training/mphase/image34d.gif

- In this case, there are some small wiggles at the surface.
- These wiggles are due to the effect of the surface tension model and spurious velocity currents at the interface.
- At small scales, these effects can become significant and influence your results.

Guided tutorial 3 – Syphon effect (flushing a tank)

What will happen in this case?



Guided tutorial 3 – Syphon effect (flushing a tank)



http://www.wolfdynamics.com/training/mphase/image34c.gif

- In this case and due to the initialization used, the tank will be emptied thanks to the syphon effect.
- This problem appears to be easy but there are many things going on.
 - A fast transient, small bubbles, maybe some cavitation, effect of surface tension, turbulence, spurious velocity currents, back flow at the outlet, mesh alignment, and so on.
- All these factors can have a strong effect on the solution stability and accuracy.

Guided tutorial 3 – Syphon effect (flushing a tank)



• When generating the mesh, try to do your best work so the cells are aligned with the free surface.

- We are going to use the following solver: **interFoam**
- Let us explore every dictionary in the case directory.
- The first step is to set the physical properties.
- Go to the directory **constant** and open the following dictionaries:
 - *g*: in this dictionary we set the gravity.
 - *transportProperties*: this dictionary contains the material properties for each phase, separated into two blocks (one for water and the other for air).
 - *momentumTransport*: in this dictionary we select the turbulence model to use.
- Remember, you will need to set the boundary conditions, initial conditions, discretization method and solution method for the turbulent variables related to the turbulence model.

- The next step is to set the boundary and initial conditions.
- Go to the directory 0 (or 0_org) and open the following dictionaries:
 - *U*: in this dictionary we set the boundary and initial conditions for the velocity vector field.
 - p_rgh: in this dictionary we set the boundary and initial conditions for the pressure scalar field.
 - *alpha.water.org*: in this dictionary we set the boundary and initial conditions for the alpha scalar field (volume of fraction).
- In the dictionary *constant/transportProperties* we defined the phases water and air, hence in the directory 0 we define the dictionary alpha.water.org that will take the values of the phase water.
- The other phase will take the remaining values.
- We set the boundary and initial conditions for the alpha scalar field (volume of fraction) in the dictionary *alpha.water*.
- The dictionary *alpha.water.org* is a backup of the original dictionary as it will be overwritten when we initialize the fields.

- Finally, we set the run-time parameter, numerical method and linear solvers.
- Go to the directory **system** and open the following dictionaries:
 - *controlDict*: in this dictionary we set general run-time parameters and function objects.
 - *fvSchemes*: in this dictionary we set the discretization method.
 - *fvSolution*: in this dictionary we set the linear solvers and parameters specific of the pressure-velocity coupling method.
- Do not worry, later on we are going to study in more details the dictionaries *fvSchemes* and *fvSolution*.

- Before proceeding to the simulation stage, we need to do a custom initialization of the fields.
- Go to the directory **system** and open the following dictionary:
 - *setFieldsDict*: we use this dictionary for custom initialization of the fields



Guided tutorial 3 – Syphon effect (flushing a tank)

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.

Important note on the VOF method and turbulent flows

- Dealing with turbulent flows using the VOF method is exactly the same as dealing with turbulence in single phase flows.
- When setting boundary and initial conditions for turbulent flows using the VOF method, it is recommended to use the primary phase properties to compute the turbulent quantities.
- If you are simulating a quiescent process, set the turbulent quantities to a small value, e.g., 10e8.

- Guided tutorial 4. Surface tension driven flow Bubble in zero gravity. VOF Small scale.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT4/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF



• According to you, what will happen to a perfect circular bubble, in perfect equilibrium, in zero gravity, with no perturbations and with given surface tension?

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF



• In reality, nothing should happen but due to the mesh, numerical diffusion, and discretization errors the bubble wobbles a little bit.

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF



• According to you, what will happen to this elliptical bubble in zero gravity, with no perturbations and no surface tension?

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF



• According to you, what will happen to this elliptical bubble in zero gravity, with no perturbations and with given surface tension?

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF



http://www.wolfdynamics.com/training/mphase/image37.gif

- In this guided tutorial we model an elliptical bubble in a zero-gravity field.
- The problem is transient in nature.
- The wobbling is due to surface tension.
- We will use the VOF approach to resolve the interface between the two phases.
- When dealing with surface tension driven flows, numerical accuracy is extremely important as the surface tension forces depend on the curvature of the surface.

Guided tutorial 4

- We are going to use the following solver: **interFoam**
- Let us explore every dictionary in the case directory.
- The first step is to set the physical properties.
- Go to the directory **constant** and open the following dictionaries:
 - *g*: in this dictionary we set the gravity.
 - *transportProperties*: this dictionary contains the material properties for each phase, separated into two blocks (one for water and the other for air).
 - *momentumTransport*: in this dictionary we select the turbulence model to use.
- In this case we are not using turbulence model (laminar).
- If you want, you can use a turbulence model. Remember, you will need to set the boundary conditions, initial conditions, discretization method and solution method for the new turbulent variables.

Guided tutorial 4

- The next step is to set the boundary and initial conditions.
- Go to the directory 0 (or 0_org) and open the following dictionaries:
 - U: in this dictionary we set the boundary and initial conditions for the velocity vector field.
 - p_rgh: in this dictionary we set the boundary and initial conditions for the pressure scalar field.
 - *alpha.phase1.org*: in this dictionary we set the boundary and initial conditions for the alpha scalar field (volume of fraction).
- In the dictionary *constant/transportProperties* we defined the phases phase1 and phase2, hence in the directory 0 we define the dictionary *alpha.phase1.org* that will take the values of the phase phase1.
- The other phase will take the remaining values.

Guided tutorial 4

- We set the boundary and initial conditions for the alpha scalar field (volume of fraction) in the dictionary *alpha.water*.
- The dictionary *alpha.water.org* is a backup of the original dictionary as it will be overwritten when we initialize the fields.
- Remember, the name of the **base type** boundary condition (dictionary *constant/polyMesh/boundary*) and the name of the **numerical type** boundary condition (dictionaries located in the directory 0) need to be the same, if not, OpenFOAM will complain.
- Also, the **base type** and **numerical type** boundary conditions need to be consistent.

Guided tutorial 4

- Finally, we set the run-time parameter, numerical method and linear solvers.
- Go to the directory **system** and open the following dictionaries:
 - *controlDict*: in this dictionary we set general run-time parameters and function objects.
 - *fvSchemes*: in this dictionary we set the discretization method.
 - *fvSolution*: in this dictionary we set the linear solvers and parameters specific of the pressure-velocity coupling method.
- Do not worry, later on we are going to study in more details the dictionaries *fvSchemes* and *fvSolution*.

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF

- Before proceeding to the simulation stage, we need to do a custom initialization of the fields.
- Go to the directory **system** and open the following dictionary:

Z ¥

• setFieldsDict.ellipse we use this dictionary for custom initialization of the
fields using a STL file



- 1.0e+00 - 0.8

- -1.3e-08

Guided tutorial 4

Surface tension driven flow or bubble in zero gravity – VOF

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.

- Guided tutorial 5. Rising bubble. VOF Small scale.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT5/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.

Guided tutorial 5

Rising bubble – VOF

- In this guided tutorial we model a rising bubble.
- The problem is transient in nature.
- The bubble will rise due to buoyancy forces.
- We will use the VOF approach to resolve the interface between the two phases.
- We will use this case to study the numerics.





References:

- http://www.featflow.de/en/benchmarks/cfdbenchmarking/bubble.html
- Klostermann, J.; Schaake, K.; Schwarze, R.: Numerical simulation of a single rising bubble by VOF with surface compression, International Journal for Numerical Methods in Fluids, DOI: 10.1002/fld.3692
Guided tutorial 5

Rising bubble – VOF



http://www.wolfdynamics.com/training/mphase/image38.gif

References:

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- http://www.featflow.de/en/benchmarks/cfdbenchmarking/bubble.html
- Klostermann, J.; Schaake, K.; Schwarze, R.: Numerical simulation of a single rising bubble by VOF with surface compression, International Journal for Numerical Methods in Fluids, DOI: 10.1002/fld.3692

Guided tutorial 5

Rising bubble – VOF



References:

- http://www.featflow.de/en/benchmarks/cfdbenchmarking/bubble.html
- Klostermann, J.; Schaake, K.; Schwarze, R.: Numerical simulation of a single rising bubble by VOF with surface compression, International Journal for Numerical Methods in Fluids, DOI: 10.1002/fld.3692

Guided tutorial 5

- We are going to use the following solver: **interFoam**
- Let us explore the dictionaries in the system directory.
- Remember:
 - The discretization method is set in the dictionary *fvSchemes*.
 - The linear solvers and parameters specific of the pressure-velocity coupling method is set in the dictionary *fvSolution*.
 - Runtime parameters such as time-step, CFL number, saving frequency, and so on, are set in the dictionary *controlDict*.
- In the folder **system/default**, you will find the case default setup.
- All the cases were run in parallel using 4 processors.

Guided tutorial 5

- Let us study the dictionary *fvSchemes*.
- In this dictionary we set the discretization method for every term appearing in the governing equations.
- Time discretization is set as follows:



- Currently the backward method is not supported for multiphase flows.
- The keyword default means that we use the same method for every requested term.
- Setting the blending factor to 0 is equivalent to the Euler method (first order accurate). Setting the blending factor to 1 is equivalent to use the pure CrankNicolson method (second order accurate).

Guided tutorial 5

- Let us study the dictionary *fvSchemes*.
- In this dictionary we set the discretization method for every term appearing in the governing equations.
- You can set the time discretization in a term-by-term basis as follows:

Guided tutorial 5

- Let us study the dictionary *fvSchemes*.
- In this dictionary we set the discretization method for every term appearing in the governing equations.
- Gradient discretization is set as follows:



Guided tutorial 5

- Let us study the dictionary *fvSchemes*.
- In this dictionary we set the discretization method for every term appearing in the governing equations.
- You can set the gradient discretization in a term-by-term basis as follows:

```
gradSchemes
{
    default Gauss linear;
    grad(U) cellMDLimited Gauss linear 0.5;
    grad(alpha.phase1) cellLimited Gauss linear 1
    grad(alpha.phase2) faceLimited Gauss linear 1
}
```

Guided tutorial 5

- If you set all the grad terms using the default option, you might get strange interface resolution.
- The difference in the results, is related to the term nHat (face unit interface normal), which should be discretize with no slope limiters.



Guided tutorial 5

Rising bubble – VOF

• The difference in the results is related to the term nHat (face unit interface normal), which should be discretize with no slope limiters if you want to get a smooth interface.



Guided tutorial 5

- Let us study the dictionary *fvSchemes*.
- In this dictionary we set the discretization method for every term appearing in the governing equations.
- Convective terms discretization is set as follows:

```
divSchemes
{
    div(rhoPhi,U)
    div(phi,alpha)
    div((((rho*nuEff)*dev2(T(grad(U))))))
    Gauss linear;
}
Gauss linear;
```

Guided tutorial 5

- Let us study the dictionary *fvSchemes*.
- In this dictionary we set the discretization method for every term appearing in the governing equations.
- Laplacian terms discretization is set as follows:



- You can also set the discretization in a term-by-term basis.
- Remember, the choice of the blending factor is related to the quality of the mesh.
- The entry snGradSchemes should use the same method as in laplacianSchemes (limited 1 in this case).

Guided tutorial 5

Rising bubble – VOF

• Comparison of different numerical schemes.





div(phirb,alpha) Gauss linear;

Z X

div(phirb,alpha) Gauss vanLeer;

Guided tutorial 5

Rising bubble – VOF

• Comparison of different numerical schemes.





div(phirb,alpha) Gauss vanLeer;

Z X

div(phirb,alpha) Gauss upwind;

Guided tutorial 5

- Let us study the dictionary *fvSolution*.
- In this dictionary we set the linear solvers and parameters specific of the pressure-velocity coupling method.
- The solution of the phasic volume of fraction is set as follows:



Guided tutorial 5

Rising bubble – VOF

• Comparison of different solution methods for phasic phases.



cAlpha 1;

, ,

cAlpha 0;



- 0.8

-0.6

- 0.4

Guided tutorial 5

Rising bubble – VOF

• Comparison of different solution methods for phasic phases.



cAlpha 1; MULESCorr yes; Execution time = 75 seconds cAlpha 1; MULESCorr no; Execution time = 65 seconds

Guided tutorial 5

Rising bubble – VOF

- Let us study the dictionary *fvSolution*.
- In this dictionary we set the linear solvers and parameters specific of the pressure-velocity coupling method.
- The solution of p_rghFinal is set as follows:



• The entry p_rghFinal is related to the last iteration of p_rgh. Remember, is a good idea to put more computational effort on this iteration, and advance fast the inner iterations.

Guided tutorial 5

Rising bubble – VOF

• Comparison of different tolerance criterion for p_rgh and p_rghFinal.



Guided tutorial 5

Rising bubble – VOF

• Comparison of multigrid vs. Newton-Krylov solvers and preconditioners for p_rgh and p_rghFinal



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Guided tutorial 5

Rising bubble – VOF

• Comparison of multigrid vs. Newton-Krylov solvers and preconditioners for p_rgh and p_rghFinal



Z X

Guided tutorial 5

- Let us study some additional entries that appears in the dictionary files *fvSolution*, *fvSchemes* and *controlDict*.
- We will see the influence of many other parameters, such as:
 - momentumPredictor
 - maxCo
 - maxAlphaCo
 - maxDeltaT
 - writeControl
 - adjustTimeStep
 - deltaT

Guided tutorial 5

Rising bubble – VOF

- Parameter \rightarrow momentumPredictor
- Dictionary file → fvSolution

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Guided tutorial 5

Rising bubble – VOF

- Parameter \rightarrow maxCo, maxAlphaCo, maxDeltaT
- Dictionary file → controlDict



maxCo	
maxAlphaCo	
maxDeltaT	
MULESCorr	
Execution time =	

Z X

2; 2; 0.1; yes; 31 seconds









naxCo	0.1;
naxAlphaCo	0.1;
naxDeltaT	0.1;
MULESCorr	yes;
Execution time =	72 seconds

Guided tutorial 5

Rising bubble – VOF

• Parameter \rightarrow writeControl

AY

Z ¥

• Dictionary file → controlDict



- 1.0e+00

- 0.8

- 0.6

- 0.4

- 0.2

_ 0.0e+00

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alpha.phase

Guided tutorial 5

Rising bubble – VOF

• Parameter \rightarrow MULESCorr

Z X

• Dictionary file → fvSolution



maxCo	
maxAlphaCo	
maxDeltaT	
MULESCorr	
Execution time =	

2; 2; 0.1; yes; 31 seconds



Z X





maxCo2;maxAlphaCo2;maxDeltaT0.1;MULESCorrno;Execution time =25 seconds

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Guided tutorial 5

Rising bubble – VOF

- Parameter \rightarrow writeControl
- Dictionary file \rightarrow controlDict

DIVERGES

- Remember MULESCorr set to no is equivalent to use an explicit scheme.
- Explicit schemes are conditionally stable.

2;





maxCo maxAlphaCo maxDeltaT writeControl writeInterval MULESCorr Execution time =

2; 0.1; runTime; 0.01; no; 25 seconds maxCo maxAlphaCo maxDeltaT writeControl writeInterval MULESCorr Execution time =

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2; 2; 0.1; adjustableRunTime; 0.01; no; 25 seconds

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Guided tutorial 5

Rising bubble – VOF

- Parameter \rightarrow maxCo, maxAlphaCo, maxDeltaT, writeControl
- Dictionary file → controlDict



maxCo	0.5
maxAlphaCo	0.5
maxDeltaT	0.1
writeControl	run
writeInterval	0.0
MULESCorr	no;
Execution time =	20

Ž 🖌

0.5; 0.1; unTime; 0.01; 10; 20 seconds



Z X



0.5; 0.5; 0.1; adjustableRunTime; 0.01; no; 30 seconds



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Guided tutorial 5

Rising bubble – VOF

- Parameter \rightarrow adjustTimeStep
- Dictionary file → controlDict



adjustTimeStep deltaT Execution time =

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> no; 0.001; 50 seconds







adjustTimeStep deltaT Execution time =

yes; 0.001; 72 seconds

Guided tutorial 5

Additional notes on the *fvSchemes* and *fvSolution* dictionaries

• The following modified convective equation (alpha) is used to track the interface between the phases,



Use a TVD scheme with gradient limiters. Good choice is the vanLeer scheme.

(phirb, alpha) Use a high order scheme. The use of interfaceCompression or linear interpolation is fine for this term.

• Where a value of $c_{\alpha} = 1$ (cAlpha), is recommended to accurately resolve the sharp interface.

Guided tutorial 5

Additional notes on the *fvSchemes* dictionary

- For interface capturing, OpenFOAM uses,
 - The MULES algorithm (semi-implicit and second order in time) to ensure that the volume fraction (alpha) remains between strict bounds of 0 and 1.
 - The interface compression scheme, based on counter-gradient transport, to maintain sharp interfaces during a simulation.
- At the following link, you can find the release notes related latest developments related to the interface capturing in OpenFOAM 8,
 - <u>https://cfd.direct/openfoam/free-software/multiphase-interface-capturing/</u>
- Among the latest improvements and developments, was the addition of the Piecewise-linear interface calculation (PLIC) family of interpolation schemes.
- The PLIC and MPLIC methods are more precise than interface compression for meshes with refinement patterns.
- These methods carry an additional computational cost.
- Regarding stability, still is a little bit early to give you our opinion

Guided tutorial 5

Additional notes on the *fvSchemes* dictionary

• The new methods can be selected in the *fvSchemes* dictionary as follows,

div(phi,alpha)	Gauss PLIC interfaceCompression vanLeer 1;
	or
div(phi,alpha)	Gauss MPLIC;

Guided tutorial 5

Additional notes on the *fvSolution* dictionary



The semi-implicit MULES offers significant speed-up and stability over the explicit MULES GT-104

Guided tutorial 5

Additional notes on the *fvSolution* dictionary



Guided tutorial 5

Additional notes on the *fvSolution* dictionary



Guided tutorial 5

Rising bubble – VOF

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.

- Guided tutorial 6. Bubble column using an Eulerian-Eulerian approach.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT6/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.
Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



- In this guided tutorial we model a bubble column.
- The problem is transient in nature and is fully 3D.
- Initially, the domain is fill with water and air is injected at the bottom with a given velocity.
- We will use an Eulerian-Eulerian approach to solve the mixture and dynamics of the two phases.
- We are going to use this case to study the different interfacial models implemented in OpenFOAM.
- This is a validation case, we are going to use the same experimental setup as in reference [1].
- To validate the results, we sample the time averaged liquid vertical velocity.

References:

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[1] Vivek V. Buwa, Vivek V. Ranade, Dynamics of gas–liquid flow in a rectangular bubble column: experiments and single/multi-group CFD simulations. Chemical Engineering Science 57 (2002) 4715 – 4736 GT-109

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach

http://www.wolfdynamics.com/wiki/multiphase/ani15.gif



Bubble plume colored by air vertical velocity

- In this guided tutorial we model a bubble column.
- The problem is transient in nature and is fully 3D.
- Initially, the domain is fill with water and air is injected at the bottom with a given velocity.
- We will use an Eulerian-Eulerian approach to solve the mixture and dynamics of the two phases.
- We are going to use this case to study the different interfacial models implemented in OpenFOAM.
- This is a validation case, we are going to use the same experimental setup as in reference [1].
- To validate the results, we sample the time averaged liquid vertical velocity.

References:

[1] Vivek V. Buwa, Vivek V. Ranade, Dynamics of gas–liquid flow in a rectangular bubble column: experiments and single/multi-group CFD simulations. Chemical Engineering Science 57 (2002) 4715 – 4736

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



- In this guided tutorial we model a bubble column.
- The problem is transient in nature and is fully 3D.
- Initially, the domain is fill with water and air is injected at the bottom with a given velocity.
- We will use an Eulerian-Eulerian approach to solve the mixture and dynamics of the two phases.
- We are going to use this case to study the different interfacial models implemented in OpenFOAM.
- This is a validation case, we are going to use the same experimental setup as in reference [1].
- To validate the results, we sample the time averaged liquid vertical velocity.

References:

[1] Vivek V. Buwa, Vivek V. Ranade, Dynamics of gas-liquid flow in a rectangular bubble column: experiments and single/multi-group CFD simulations. Chemical Engineering Science 57 (2002) 4715 – 4736

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Guided tutorial 6

Bubble column – Eulerian-Eulerian approach

• Cases performed for this validation study:

C1	DM1-VM-LM1-NFI	NTM	TPF	
C2	DM1-VM-NFI	NTM	MPF	
C3	DM1-VM-LM1-WL-NFI	NTM	TPF	
C4	DM1-VM-NFI	NTM	TPF	
C5	DM2-VM-NFI	NTM	TPF	
C6	DM1-VM-FI	NTM	TPF	
C7	DM1-VM-LM1-FI	NTM	TPF	
C8	DM1-VM-LM2-FI	NTM	TPF	
C9	DM1-VM-FI	TM	TPF	RANS1
C10	DM1-VM-FI-TD1	TM	TPF	RANS1
C11	DM1-VM-LM3-FI-TD2	TM	TPF	RANS1
C12	DM1-VM-LM4-FI	NTM	TPF	
C13	DM3-VM-LM2-FI	NTM	TPF	
C14	DM1-VM-FI-TD3	TM	TPF	RANS2
C15	DM1-VM-FI	TM	TPF	LES1
C16	DM1-VM-FI	TM	TPF	LES2
	C1 C2 C3 C4 C5 C6 C7 C8 C9 C10 C11 C12 C13 C14 C15 C16	C1 DM1-VM-LM1-NFI C2 DM1-VM-NFI C3 DM1-VM-LM1-WL-NFI C4 DM1-VM-NFI C5 DM2-VM-NFI C6 DM1-VM-FI C7 DM1-VM-LM1-FI C8 DM1-VM-LM2-FI C9 DM1-VM-FI C10 DM1-VM-FI C11 DM1-VM-FI C12 DM1-VM-LM3-FI-TD2 C13 DM3-VM-LM2-FI C14 DM1-VM-FI-TD3 C15 DM1-VM-FI	C1 DM1-VM-LM1-NFI NTM C2 DM1-VM-NFI NTM C3 DM1-VM-LM1-WL-NFI NTM C4 DM1-VM-NFI NTM C5 DM2-VM-NFI NTM C6 DM1-VM-FI NTM C7 DM1-VM-FI NTM C8 DM1-VM-LM1-FI NTM C9 DM1-VM-EM2-FI NTM C10 DM1-VM-FI-TD1 TM C11 DM1-VM-EM3-FI-TD2 TM C12 DM1-VM-LM4-FI NTM C13 DM3-VM-LM2-FI NTM C14 DM1-VM-FI-TD3 TM C15 DM1-VM-FI TM C16 DM1-VM-FI TM	C1 DM1-VM-LM1-NFI NTM TPF C2 DM1-VM-NFI NTM MPF C3 DM1-VM-LM1-WL-NFI NTM TPF C4 DM1-VM-NFI NTM TPF C5 DM2-VM-NFI NTM TPF C6 DM1-VM-FI NTM TPF C6 DM1-VM-FI NTM TPF C7 DM1-VM-LM1-FI NTM TPF C8 DM1-VM-LM2-FI NTM TPF C9 DM1-VM-FI TM TPF C10 DM1-VM-FI-TD1 TM TPF C11 DM1-VM-LM3-FI-TD2 TM TPF C12 DM1-VM-LM3-FI-TD2 TM TPF C13 DM3-VM-LM2-FI NTM TPF C14 DM1-VM-FI-TD3 TM TPF C15 DM1-VM-FI TM TPF C16 DM1-VM-FI TM TPF

DM = drag model, VM = virtual mass, LM = lift model, TD = turbulent dispersion, WL = wall lubrication, NFI = no phase inversion, FI = phase inversion, NTM = no turbulence model, TM = turbulence model, TPF = twoPhaseEulerFoam, MPF = multiphaseEulerFoam

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



Lift models were used

No lift models were used

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



http://www.wolfdynamics.com/wiki/multiphase/ani6.gif







http://www.wolfdynamics.com/wiki/multiphase/ani7.gif





http://www.wolfdynamics.com/wiki/multiphase/ani8.gif



Note: the results presented were obtained using OpenFOAM 3.x

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach



Note: the results presented were obtained using OpenFOAM 3.x

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach

- This case is ready to run.
- Despite the fact that the mesh is small the cases are time consuming.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT6/validation_case

- In the case directory you will find the following sub-directories
 - validation_case/laminar: laminar bubble column using multiphaseEulerFoam
 - validation_case/turbulent: turbulent bubble column using multiphaseEulerFoam
- Hereafter we report the execution times for each case (serial runs):
 - validation_case/laminar : 77801 seconds
 - validation_case/turbulent : 84643 seconds

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach

- We are going to use the following solver: **multiphaseEulerFoam**
- Let us explore the dictionaries in the constant directory.
 - g: in this dictionary you set the gravity field.
 - *phaseProperties*: in this dictionary you set how phases interacts and the physical and interfacial models.
 - thermophysicalProperties.air: in this dictionary you set the thermo physical properties for the phase air.
 - thermophysicalProperties.water: in this dictionary you set the thermo physical properties for the phase water.
 - momentumTransport.air: in this dictionary you set the turbulence model for the phase air.
 - *momentumTransport.water*: in this dictionary you set the turbulence model for the phase water.

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach

- We are going to explore as well the dictionaries in the system directory.
- The discretization method is set in the dictionary *fvSchemes*.
- The linear solvers and parameters specific of the pressure-velocity coupling method is set in the dictionary *fvSolution*.
- Runtime parameters such as time-step, CFL number, saving frequency, and so on, are set in the dictionary *controlDict*.
- Remember, as we are solving different equations, the dictionaries will be slightly different from the dictionaries we have used so far.

Guided tutorial 6

Bubble column – Eulerian-Eulerian approach

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.
- If you are interested, the experimental results (time averaged liquid vertical velocity) are located in the case directory
 - \$TM/multiphase/guided_tutorials/GT6/validation_case/exp

you can plot the experimental results and the simulation results using gnuplot, Python, or your favorite plotting application.

- Guided tutorial 7. Fluidized bed using kinetic theory of granular flows (KTGF)
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT7/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.

Guided tutorial 7

Fluidized bed – Eulerian-Eulerian approach using kinetic theory of granular flows

- In this guided tutorial we model a fluidized bed.
- The problem is transient in nature.
- Air is injected at the bottom with a given velocity.
- The particles bed has an initial height.
- We will use an Eulerian-Eulerian approach with kinetic theory of granular flows.
- We are going to use this case to study how to setup a granular flow and the different interfacial models implemented in OpenFOAM.
- This is a validation case, we are going to use an experimental setup similar to the one found in reference [1].
- To validate the results, we compare the results qualitatively

Bed initial height Gas inlet

1.0e+00

alpha.air

- 0.9

- 0.8

- 0.7

- 0.6

- 0.5 - 4.5e-0



[1] Goldschmidt, M.J.V. and Hoomans, B.P.B. and Kuipers, J.A.M. (2002) *Detailed comparison of Euler-Lagrange and Euler-Euler models for simulation of dense gas fluidised beds.* In: 10th Workshop on Two-phase Flow Predictions, April 9-12, 2002, Merseburg, Germany (pp. pp. 285-299)

Guided tutorial 7



Guided tutorial 7



Guided tutorial 7



Guided tutorial 7



Guided tutorial 7

Fluidized bed – Eulerian-Eulerian approach using kinetic theory of granular flows



References:

[1] Goldschmidt, M.J.V. and Hoomans, B.P.B. and Kuipers, J.A.M. (2002) *Detailed comparison of Euler-Lagrange and Euler-Euler models for simulation of dense gas fluidised beds.* In: 10th Workshop on Two-phase Flow Predictions, April 9-12, 2002, Merseburg, Germany (pp. pp. 285-299)

Guided tutorial 7

Fluidized bed – Eulerian-Eulerian approach using kinetic theory of granular flows

- For completeness, we present the results of particle-particle interactions (lagrangian solvers).
- The eulerian-eulerian solver does not provide this information.

http://www.wolfdynamics.com/training/mphase/image43.gif

http://www.wolfdynamics.com/training/mphase/image44.gif



Guided tutorial 7

- In the case directory \$TM/multiphase/guided_tutorials/GT7/KTGF, you will find the following sub-directories:
 - Goldschmidt/C1: laminar fluidized bed using multiphaseEulerFoam with phasePressure model for the solid phase (particles).
 - Goldschmidt/C2: turbulent fluidized bed using multiphaseEulerFoam with kineticTheory model for the solid phase (particles).
- At this point, choose with case do you want to run.
- The computing time and run instructions are similar.

Guided tutorial 7

- In the case directory **\$TM/multiphase/guided_tutorials/GT7**, you will find the following additional sub-directories:
 - DPM: fluidized bed using DPMFoam
 - MPPIC: fluidized bed using MPPICFoam
- For completeness, hereafter we report the execution times for each solver (serial runs):
 - DPM: 166500 seconds (about two days)
 - MPPIC: 6810 seconds
 - KTGP/Goldschmidt/C2: 250 seconds

Guided tutorial 7

- We are going to use the following solver: **multiphaseEulerFoam**
- Let us explore the dictionaries in the constant directory.
 - g: in this dictionary you set the gravity field.
 - *phaseProperties*: in this dictionary you set how phases interacts and the physical and interfacial models.
 - thermophysicalProperties.air: in this dictionary you set the thermo physical properties for the phase air.
 - thermophysicalProperties.particles: in this dictionary you set the thermo physical properties for the particles.
 - momentumTransport.air: in this dictionary you set the turbulence model for the phase air.
 - *momentumTransport.particles*: in this dictionary you set the turbulence model for the particles.

Guided tutorial 7

- We are going to explore as well the dictionaries in the system directory.
- The discretization method is set in the dictionary *fvSchemes*.
- The linear solvers and parameters specific of the pressure-velocity coupling method is set in the dictionary *fvSolution*.
- Runtime parameters such as time-step, CFL number, saving frequency, and so on, are set in the dictionary *controlDict*.
- Remember, as we are solving different equations, the dictionaries will be slightly different from the dictionaries we have used so far.

Guided tutorial 7

Fluidized bed – Eulerian-Eulerian approach using kinetic theory of granular flows

- When setting the continuous phase name using the solver MPPICFoam, DPMFoam or multiphaseEulerFoam, remember to use the same phase name when assigning the boundary conditions.
- There are a few boundary conditions that require the name of the continuous phase field, e.g.,



 Check the online documentation or the source code to see what are the default values of the boundary conditions and update them accordingly

Guided tutorial 7

Fluidized bed – Eulerian-Eulerian approach using kinetic theory of granular flows

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.
- Remember to choose among the DPM, MPPIC and KTGF cases.

- Guided tutorial 8. Particle injection in a mixing tank Lagrangian approaches.
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT8/

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.

- The particles tracking is performed using a Lagrangian approach.
- In the Eulerian-Lagrangian approach the solution of the continuous phase and the particles (discrete phase) is performed at the same time or imported from a previous case.
- This tutorial focuses on three different models available in OpenFOAM:
 - particleFoam: an uncoupled lagrangian tracking model.
 - MPPICFoam: the MPPIC or Multiphase Particle-in-Cell method for collisional exchange. In this approach, particle-particle interactions are represented by models which utilize mean values calculated on the Eulerian mesh. The MPPIC is suitable for dense to dilute regimes.

- The following tutorials focus on the tracking of water droplets in a static mixing tank by using the following solvers:
 - particleFoam
 - MPPICFoam
- To perform the tutorial, we will use the following mixing tank geometry.



- 1. Case1: Lagrangian tracking tutorial without hydrodynamic coupling
- 2. Case2: Lagrangian tracking tutorial with hydrodynamic coupling

- Case 1: Lagrangian tracking tutorial without hydrodynamic coupling
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT8/icouncoupled

- 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 that you open the README.FIRST, file and typing 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.
- From this point on, please follow me. We are all going to work at the same pace.

- In this tutorial we will track water droplets in a static mixer by using a one-way coupling strategy.
- We will consider the gravity and the drag forces.



- The one-way coupled approach is based on the availability of an airflow solution previously computed using any solver.
- The solution is then used to compute the position of the particles using a Lagrangian model.
- This tutorial case is organized as follows:
 - **GT8/icouncoupled/c1_airflow**: in this directory you will find the case setup of the single-phase case (precursor simulation). We will run this case to obtain the flow solution that will be used to run the uncoupled Lagrangian solver.
 - **GT8/icouncoupled/c2_particles**: in this directory you will find Lagrangian particles tracking case setup. This case uses the solution obtained in the previous step. The solution can come from the same mesh or a finer or coarser mesh.

Guided tutorial 8 – Particle injection in a mixing tank

- In the c1_airflow directory you will find the precursor simulation case setup.
- You will also find a precomputed solution (laminar), so you do not need to run the solution.



http://www.wolfdynamics.com/training/mphase/image50.gif

Guided tutorial 8 – Particle injection in a mixing tank

- In the c1_airflow directory you will find the precursor simulation case setup.
- You will also find a precomputed solution (laminar), so you do not need to run the solution.



http://www.wolfdynamics.com/training/mphase/image51.gif

http://www.wolfdynamics.com/training/mphase/image52.gif
Guided tutorial 8

Particle injection in a mixing tank

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.
- Remember, you can start this case from scratch or from the pre-computed solution located in the directory c1_airflow.

Guided tutorial 8 – Particle injection in a mixing tank

• Go to the constant/ directory and explore the *kinematicCloudProperties* dictionary. In this input file, you will find the core options of the uncoupled lagrangian setup.

	solution					
	{ active true;					
	coupled false; transient yes;					
cellValueSourceCorrection on:						
interpolationSchemes						
	rho.air cell;					
	mu.air cell;					
	} integrationSchemes					
	{					
	U Euler;					
	constantProperties					
	{ rho0 100;					
	alphaMax 0.9;					
I						

Guided tutorial 8 – Particle injection in a mixing tank

• Go to the constant/ and explore the *kinematicCloudProperties* dictionary, here we find the core of our uncoupled lagrangian setup.

In the subModels section, you define the models to be used with the lagrangian particles. Such as the forces applied to the particles, the injection models, the particle-particle treatment, the wall interaction, collision, damping and dispersion models, etc.	<pre>subModels { particleForces { sphereDrag; gravity; } injectionModels { model1 { type patchInjection; parcelBasisType fixed; patchName mixer_inlet1; U0 (0 -8 0); rParticle 1; U0 (0 -8 0); reactive 1; reactive 1;</pre>
	patchName mixer_inlet1; U0 (0 -8 0); nParticle 1; parcelsPerSecond 1000; flowRateProfile constant 1; massTotal 0; SOI 0; duration 5;

• Go to the constant/ and explore the *kinematicCloudProperties* dictionary, here we find the core of our uncoupled lagrangian setup.

```
sizeDistribution
       type
                normal;
       normalDistribution
         expectation 100e-6;
         variance
                      25e-6;
         minValue
                    20e-6;
         maxValue 180e-6;
dispersionModel none;
patchInteractionModel localInteraction;
localInteractionCoeffs
  patches
    mixer_wall
       type rebound;
    mixer_inlet1
       type escape;
 . . .
```

- A detailed overview of the constant/kinematicCloudProperties dictionary is provided in the MPPIC tutorial.
- The icoUncoupledKinematicParcelFoam model is suitable when a low volume fraction of the discrete phase is expected and the influence of the particles momentum on the continuous phase is not relevant.
- The constant/g dictionary contains the specification of the gravity force



 Correction for non-spherical particles can be adopted by choosing a proper drag coefficient function.

Guided tutorial 8 – Particle injection in a mixing tank

Accessing the particle results

- The results of icoUncoupledKinematicParcelFoam are saved inside each time directory inside the <time_directory>/lagrangian/kinematicCloud sub-folder.
- The particle quantities can be stored as readable ASCII files and can be easily parsed by userdefined scripts.
- Each quantities of interest is stored in a dedicate file. *e.g.*, the particle *age* file will looks like the following:



Guided tutorial 8 – Particle injection in a mixing tank

Accessing the particle results

- Each particle is defined by its ID number that can be accessed in the origId file.
- At each time, the *origId* list is updated and the particle order may change. The user has to pay attention to the current *origID* when parsing the Lagrangian data set of a give time step.
- Common variables stored in the **lagrangian/kinematicCloud** sub-folder are the particle age, density, velocity, diameter, angular momentum, temperature, position, etc...



Guided tutorial 8 – Particle injection in a mixing tank

Visualizing Lagrangian data in Paraview



mesh part and the diameter fields

Guided tutorial 8 – Particle injection in a mixing tank

Visualizing Lagrangian data in Paraview

<u>File E</u> dit ⊻iew <u>S</u> ources	Filters Tools Catalyst Macros		AMR Contour		Generate Surface Normals
Search Ctrl+Space Becent AMR Annotation CTH		AMR CutPlane	0	Glyph	
		AMR Dual Clip		Glyph With Custom Source	
		AMR Fragment Integration		Gradient	
		AMR Fragments Filter		Gradient Of Unstructured Dat	
		Add Field Arrays		Grid Connectivity	
	<u>C</u> ommon		Angular Periodic Filter	90	Group Datasets
Data Analysis >		Annotate Time Filter	1	Histogram	
		Append Attributes		Image Data To AMR	
Miver OpenFOAM	Quadrature Points +		Append Datasets		Image Data to Point Set
B Mixel Open OAM	Statistics		Append Geometry		ImageResampling
	_Temporal →		Block Scalars		Integrate Variables
	<u>A</u> lphabetical +		Calculator		Interpolate to Quadrature Poi
			Cell Centers		Intersect Fragments
		Cell Data to Point Data		Iso Volume	
			Clean		K Means
			Clean Cells to Grid		Legacy Glyph
			Clean to Grid		Level Scalars(Non-Overlappin
		0	Clip		Level Scalars(Overlapping AM
			Clip Closed Surface		Linear Extrusion
			Clip Generic Dataset		Loop Subdivision
			Compute Derivatives		Mask Points
			Compute Quartiles		Material Interface Filter
L		_	Connectivity		Median
Properties Information		Contingency Statistics		Merge Blocks	
Properties contraction of the second se			S Contour		Mesh Quality
		_	Contour Generic Dataset		Multicorrelative Statistics
GP Apply	<u>Reset</u>		Convert AMR dataset to Multi-block		Normal Glyphs
			Curvature		Octree Depth Limit
Search (use Esc to clear text)		-	Decimate		Octree Depth Scalars
🔄 Include Zones	Patch Names		Delaunay 2D		Outline
		4	Delaunay 3D		Outline Corners
🗶 Interpolate volFields	Extrapolate Patches		Descriptive Statistics		Outline Curvilinear DataSet
· · · · · · · · · · · · · · · · · · ·		-	Elevation		ParticlePath
Update GUI			Environment Annotation		ParticleTracer
Lise VTKPolyhedron			Extract AMR Blocks		Pass Arrays
		18	Extract Bag Plots		Plot Data
Mesh Parts			Extract Block	0.0	Plot Global Variables Over Tir
🗶 internalMesh			Extract CTH Parts		Plot On Intersection Curves

Extracts the particle position bock

Guided tutorial 8 – Particle injection in a mixing tank

Visualizing Lagrangian data in Paraview



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Visualizing Lagrangian data in Paraview



Guided tutorial 8 – Particle injection in a mixing tank

Visualizing Lagrangian data in Paraview



Guided tutorial 8 – Particle injection in a mixing tank

- Particles trajectories evolution (injection ends after 10 seconds).
- Left image: particles colored by diameter distribution. The diameter of the particles is constant (0.02 m). Right image: particles colored by velocity.



http://www.wolfdynamics.com/training/mphase/image53.gif

Guided tutorial 8 – Particle injection in a mixing tank

• The particles stick at the wall due to large size of the boundary layer mesh (particles are not to scale, they have been exaggerated).



Guided tutorial 8 – Particle injection in a mixing tank

- 1. Lagrangian tracking tutorial without hydrodynamic coupling
- 2. Lagrangian tracking tutorial with hydrodynamic coupling

- Case 1: Lagrangian tracking tutorial with hydrodynamic coupling
- This case is ready to run.
- The case is located in the directory:

\$TM/multiphase/guided_tutorials/GT8/MPPICFoam

- 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 that you open the README.FIRST, file and typing the commands in the terminal; in this way, you will get used with the command line interface and OpenFOAM commands.
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- From this point on, please follow me. We are all going to work at the same pace.

Guided tutorial 8 – Particle injection in a mixing tank

- The MPPIC method maps the discrete phase particle properties on the Eulerian grid of the continuous phase. Then, after the solution of the fluid dynamics, the quantities of the discrete phase are updated by the model.
- The actual particle size distribution (a probabilistic function) is synthetized into a smaller number of *computational particles* that are directly solved by the Lagrangian method with a strong reduction of the computational requirements of the model.
- As for the other Lagrangian solvers implemented in OpenFOAM, non-spherical particles can be modelled by MPPICFoam by choosing a proper drag coefficient and aspect ratio model.

Guided tutorial 8

Particle injection in a mixing tank

- At this point, we are ready to run the simulation.
- To run the tutorial automatically, type in the terminal:

1. \$> run_all.sh

- Feel free to open the file run_all.sh to see all the commands used.
- If you prefer to run the case manually, type the commands in the terminal window.

• Go to the constant/ and explore the content of the *transportProperties* dictionary:



• The *turbulenceProperties.air* dictionary contains information on the turbulence model of the continuous phase (Eulerian approach).

















Guided tutorial 8 – Particle injection in a mixing tank

• You should get something like this:



Laminar solution – No packing model http://www.wolfdynamics.com/training/mphase/image54.gif

Note: particles scaled by a factor of 200 times.

Time: 0.050



Turbulent solution – No packing model

http://www.wolfdynamics.com/training/mphase/image55.gif

Guided tutorial 8 – Particle injection in a mixing tank

• You should get something like this:



Turbulent solution – No packing model http://www.wolfdynamics.com/training/mphase/image55.gif

Note: particles scaled by a factor of 200 times.

Turbulent solution – Implicit packing model

http://www.wolfdynamics.com/training/mphase/image56.gif

Guided tutorial 8 – Particle injection in a mixing tank

• You should get something like this:



Turbulent solution – No packing model http://www.wolfdynamics.com/training/mphase/image57.gif

Turbulent solution – Implicit packing model

http://www.wolfdynamics.com/training/mphase/image58.gif

Guided tutorial 8 – Particle injection in a mixing tank

With respect to a standard Eulerian simulation, the content of the log file has been modified with the addition of the following lines:



Guided tutorial 8 – Particle injection in a mixing tank

With respect to a standard Eulerian simulation, the content of the log file has been modified with the addition of the following lines:

