

# CFDEMcoupling Documentation

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## 1. Contents

The CFDEMcoupling documentation is organized into the following sections. If you find errors or omissions in this manual or have suggestions for useful information to add, please send an email to the developers so we can improve the CFDEMcoupling documentation.

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  - 1.2 [Installation](#)
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### 1.1 About CFDEMcoupling

CFDEM coupling provides an open source parallel coupled CFD-DEM framework combining the strengths of [LIGGGHTS](#) DEM code and the Open Source CFD package [OpenFOAM\(R\)\(\\*\)](#). The CFDEMcoupling toolbox allows to expand standard CFD solvers of [OpenFOAM\(R\)\(\\*\)](#) to include a coupling to the DEM code [LIGGGHTS](#). In this toolbox the particle representation within the CFD solver is organized by "cloud" classes. Key functionalities are organised in sub-models (e.g. force models, data exchange models, etc.) which can easily be selected and combined by dictionary settings.

The coupled solvers run fully parallel on distributed-memory clusters. Features are:

- its modular approach allows users to easily implement new models
- its MPI parallelization enables to use it for large scale problems
- the [forum](#) on CFD-DEM gives the possibility to exchange with other users / developers

- the use of GIT allows to easily update to the latest version
- basic documentation is provided

The file structure:

- *src* directory including the source files of the coupling toolbox and models
- *applications* directory including the solver files for coupled CFD-DEM simulations
- *doc* directory including the documentation of CFDEMcoupling
- *tutorials* directory including basic tutorial cases showing the functionality

Details on installation are given on the [CFDEMproject WWW Site](#) . The functionality of this CFD-DEM framework is described via [tutorial cases](#) showing how to use different solvers and models.

CFDEMcoupling stands for Computational Fluid Dynamics (CFD) -Discrete Element Method (DEM) coupling.

CFDEMcoupling is an open-source code, distributed freely under the terms of the GNU Public License (GPL).

Core development of CFDEMcoupling is done by Christoph Goniva and Christoph Kloss, both at DCS Computing GmbH, 2012

This documentation was written by Christoph Goniva, DCS Computing GmbH, 2012

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## 1.2 Installation

Please follow the installation routine provided at [www.cfdem.com](http://www.cfdem.com). In order to get the latest code version, please use the git repository at <http://github.com> ([githubAccess](#)).

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## 1.3 Tutorials

### General:

Each solver of the CFDEMcoupling is comes with at least one tutorial example, showing its functionality and correct usage. Provided that the installation is correct, the tutorials can be run via "Allrun.sh" shell scripts. These scripts perform all necessary steps (preprocessing, run, postprocessing, visualization).

### Location:

The tutorials can be found in the directory \$CFDEM\_PROJECT\_DIR/tutorials, which can be reached by typing "cfdemTut"

### Structure:

Each case is structured in a directory called "CFD" covering the CFD relevant settings and data, and a directory called "DEM" covering the DEM relevant settings and data. This allows to easily expand a pure CFD or DEM simulation case to a coupled case.

## Usage:

Provided that the installation is correct, the tutorials can be run via "Allrun.sh" shell script, executed by typing `./Allrun.sh`. The successful run of the script might need some third party software (e.g. octave, evince, etc.).

## Settings:

The main settings of a simulation are done via dictionaries:

The DEM setup of each case is defined by a [LIGGGHTS](#) input file located in `$caseDir/DEM` (e.g. `in.liggghts_init`). For details on the [LIGGGHTS](#) setup, please have a look in the [LIGGGHTS](#) manual.

Standard CFD settings are defined in `$caseDir/CFD/constant` (e.g. `transportProperties`, `RASproperties`, etc.) and `$caseDir/CFD/system` (e.g. `fvSchemes`, `controlDict`). You can find more information on that in [OpenFOAM\(R\)\(\\*\)](#) documentations ([www.openFoam.com](http://www.openFoam.com))(\*).

Settings of the coupling routines are defined in `$caseDir/CFD/constant/couplingProperties` (e.g. force models, data exchange model, etc.) and `$caseDir/CFD/constant/ligggghtsCommands` (allows to execute a LIGGGHTS command during a coupled simulation).

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## 1.4 "couplingProperties" dictionary

### General:

In the "couplingProperties" dictionary the setup of the coupling routines of the CFD-DEM simulation are defined.

**Location:** `$caseDir/CFD/constant`

### Structure:

The dictionary is divided into two parts, "sub-models & settings" and "sub-model properties".

In "sub-models & settings" the following routines must be specified:

- `modelType`
- `couplingInterval`
- `voidFractionModel`
- `locateModel`
- `meshMotionModel`
- `regionModel`
- `IOModel`
- `dataExchangeModel`
- `averagingModel`
- `forceModels`
- `momCoupleModels`
- `turbulenceModelType`

In "sub-model properties" sub-dictionaries might be defined to specify model specific parameters.

### Settings:

Reasonable example settings for the "couplingProperties" dictionary are given in the tutorial cases.

modelType

"modelType" refers to the formulation of the equations to be solved. Choose "A" or "B", according to Zhou et al. (2010): "Discrete particle simulation of particle-fluid flow: model formulations and their applicability", JFM. "A" requires the use of the force models gradPForce and viscForce, whereas "B" requires the force model "Archimedes".

couplingInterval

The coupling interval determines the time passing between two CFD-DEM data exchanges.

A useful procedure would be: 1) Set the DEM TS in the in.xxx according to the needs of the pure DEM problem. 2) Set the "couplingInterval", which refers to the DEM timesteps. Depending on the problem you will need to have a close (small couplingInterval) or loose coupling. 3) Choose the CFD timestep in the controlDict. it must be equal or smaller than the coupling time, otherwise you will get the error: "Error - TS bigger than coupling interval!".

Example: DEMts=0.00001s, couplingInterval=10 exchange data (=couple) will happen every 0.0001s.

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## 1.5 "liggghtsCommands" dictionary

### General:

In the "liggghtsCommands" dictionary liggghts commands being executed during a coupled CFD-DEM simulation are specified.

**Location:** \$caseDir/CFD/constant

### Structure:

The dictionary is divided into two parts, first a list of "liggghtsCommandModels" is defined, then the settings for each model must be specified.

### Settings:

Reasonable example settings for the "liggghtsCommands" dictionary are given in the tutorial cases.

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## 1.6 Models/Solvers

This section lists all CFDEMcoupling sub-models and solvers alphabetically, with a separate listing below of styles within certain commands.

<a href="#">IOModel</a>	<a href="#">IOModel_basicIO</a>
<a href="#">IOModel_noIO</a>	<a href="#">IOModel_trackIO</a>
<a href="#">averagingModel</a>	<a href="#">averagingModel_dilute</a>
<a href="#">cfdemSolverIB</a>	<a href="#">cfdemSolverPiso</a>
<a href="#">cfdemSolverPisoScalar</a>	<a href="#">clockModel</a>
<a href="#">clockModel_noClock</a>	<a href="#">clockModel_standardClock</a>
<a href="#">dataExchangeModel</a>	<a href="#">dataExchangeModel_noDataExchange</a>
<a href="#">dataExchangeModel_oneWayVTK</a>	<a href="#">dataExchangeModel_twoWayFiles</a>
<a href="#">dataExchangeModel_twoWayMPI</a>	<a href="#">forceModel</a>
<a href="#">forceModel_Archimedes</a>	<a href="#">forceModel_ArchimedesIB</a>

<a href="#"><u>forceModel_DiFeliceDrag</u></a>	<a href="#"><u>forceModel_GidaspowDrag</u></a>
<a href="#"><u>forceModel_KochHillDrag</u></a>	<a href="#"><u>forceModel_LaEuScalarTemp</u></a>
<a href="#"><u>forceModel_MeiLift</u></a>	<a href="#"><u>forceModel_SchillerNaumannDrag</u></a>
<a href="#"><u>forceModel_ShirgaonkarIB</u></a>	<a href="#"><u>forceModel_gradPForce</u></a>
<a href="#"><u>forceModel_noDrag</u></a>	<a href="#"><u>forceModel_virtualMassForce</u></a>
<a href="#"><u>forceModel_viscForce</u></a>	<a href="#"><u>liggghtsCommandModel</u></a>
<a href="#"><u>liggghtsCommandModel_execute</u></a>	<a href="#"><u>liggghtsCommandModel_readLiggghtsData</u></a>
<a href="#"><u>liggghtsCommandModel_runLiggghts</u></a>	<a href="#"><u>liggghtsCommandModel_writeLiggghts</u></a>
<a href="#"><u>locateModel</u></a>	<a href="#"><u>locateModel_engineSearch</u></a>
<a href="#"><u>locateModel_engineSearchIB</u></a>	<a href="#"><u>locateModel_standardSearch</u></a>
<a href="#"><u>locateModel_turboEngineSearch</u></a>	<a href="#"><u>meshMotionModel</u></a>
<a href="#"><u>meshMotionModel_noMeshMotion</u></a>	<a href="#"><u>momCoupleModel</u></a>
<a href="#"><u>momCoupleModel_explicitCouple</u></a>	<a href="#"><u>momCoupleModel_implicitCouple</u></a>
<a href="#"><u>momCoupleModel_noCouple</u></a>	<a href="#"><u>regionModel</u></a>
<a href="#"><u>regionModel_allRegion</u></a>	<a href="#"><u>voidfractionModel</u></a>
<a href="#"><u>voidfractionModel_GaussVoidFraction</u></a>	<a href="#"><u>voidfractionModel_IBVoidFraction</u></a>
<a href="#"><u>voidfractionModel_bigParticleVoidFraction</u></a>	<a href="#"><u>voidfractionModel_centreVoidFraction</u></a>
<a href="#"><u>voidfractionModel_dividedVoidFraction</u></a>	

## **averagingModel\_dense command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
averagingModel dense;
```

### **Examples:**

```
averagingModel dense;
```

### **Description:**

The averaging model performs the Lagrangian->Eulerian mapping of data (e.g. particle velocities). In the "cfDEMParticle cloud" this averaging model is used to calculate the average particle velocity inside a CFD cell. The "dense" model is supposed to be applied to cases where the granular regime is rather dense. The particle velocity inside a CFD cell is evaluated as an ensemble average of the particle velocities.

### **Restrictions:**

None.

### **Related commands:**

[averagingModel](#), [dilute](#)

**Default:** none

## **averagingModel\_dilute command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
averagingModel dilute;
```

### **Examples:**

```
averagingModel dilute;
```

### **Description:**

The averaging model performs the Lagrangian->Eulerian mapping of data (e.g. particle velocities). In the "cfDEMParticle cloud" this averaging model is used to calculate the average particle velocity inside a CFD cell. The "dilute" model is supposed to be applied to cases where the granular regime is rather dilute. The particle velocity inside a CFD cell is evaluated from a single particle in a cell (no averaging).

### **Restrictions:**

This model is computationally efficient, but should only be used when only one particle is inside one CFD cell.

### **Related commands:**

[averagingModel\\_dense](#)

## averagingModel command

### Syntax:

Defined in couplingProperties dictionary.

```
averagingModel model;
```

- model = name of averaging model to be applied

### Examples:

```
averagingModel dense;  
averagingModel dilute;
```

Note: This examples list might not be complete - please look for other averaging models (averagingModel\_XY) in this documentation.

### Description:

The averaging model performs the Lagrangian->Eulerian mapping of data (e.g. particle velocities).

### Restrictions:

None.

### Related commands:

[dense](#), [dilute](#)

**Default:** none



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### 1.1 About CFDEMcoupling

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## 1.3 Tutorials

### General:

Each solver of the CFDEMcoupling is comes with at least one tutorial example, showing its functionality and correct usage. Provided that the installation is correct, the tutorials can be run via "Allrun.sh" shell scripts. These scripts perform all necessary steps (preprocessing, run, postprocessing, visualization).

### Location:

The tutorials can be found in the directory \$CFDEM\_PROJECT\_DIR/tutorials, which can be reached by typing "cfdemTut"

## Structure:

Each case is structured in a directory called "CFD" covering the CFD relevant settings and data, and a directory called "DEM" covering the DEM relevant settings and data. This allows to easily expand a pure CFD or DEM simulation case to a coupled case.

## Usage:

Provided that the installation is correct, the tutorials can be run via "Allrun.sh" shell script, executed by typing "./Allrun.sh". The successful run of the script might need some third party software (e.g. octave, evince, etc.).

## Settings:

The main settings of a simulation are done via dictionaries:

The DEM setup of each case is defined by a [LIGGGHTS](#) input file located in \$caseDir/DEM (e.g. in.liggghts\_init). For details on the [LIGGGHTS](#) setup, please have a look in the [LIGGGHTS](#) manual.

Standard CFD settings are defined in \$caseDir/CFD/constant (e.g. transportProperties, RASproperties, etc.) and \$caseDir/CFD/system (e.g. fvSchemes, controlDict). You can find more information on that in [OpenFOAM\(R\)\(\\*\)](#) documentations ([www.openFoam.com](http://www.openFoam.com))(\*).

Settings of the coupling routines are defined in \$caseDir/CFD/constant/[couplingProperties](#) (e.g. force models, data exchange model, etc.) and \$caseDir/CFD/constant/[liggghtsCommands](#) (allows to execute a LIGGGHTS command during a coupled simulation).

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## 1.4 "couplingProperties" dictionary

### General:

In the "couplingProperties" dictionary the setup of the coupling routines of the CFD-DEM simulation are defined.

**Location:** \$caseDir/CFD/constant

### Structure:

The dictionary is divided into two parts, "sub-models & settings" and "sub-model properties".

In "sub-models & settings" the following routines must be specified:

- modelType
- couplingInterval
- voidFractionModel
- locateModel
- meshMotionModel
- regionModel
- IOModel
- dataExchangeModel
- averagingModel

- forceModels
- momCoupleModels
- turbulenceModelType

In "sub-model properties" sub-dictionaries might be defined to specify model specific parameters.

### Settings:

Reasonable example settings for the "couplingProperties" dictionary are given in the tutorial cases.

modelType

"modelType" refers to the formulation of the equations to be solved. Choose "A" or "B", according to Zhou et al. (2010): "Discrete particle simulation of particle-fluid flow: model formulations and their applicability", JFM. "A" requires the use of the force models gradPForce and viscForce, whereas "B" requires the force model "Archimedes".

couplingInterval

The coupling interval determines the time passing between two CFD-DEM data exchanges.

A useful procedure would be: 1) Set the DEM TS in the in.xxx according to the needs of the pure DEM problem. 2) Set the "couplingInterval", which refers to the DEM timesteps. Depending on the problem you will need to have a close (small couplingInterval) or loose coupling. 3) Choose the CFD timestep in the controlDict. it must be equal or smaller than the coupling time, otherwise you will get the error: "Error - TS bigger than coupling interval!".

Example: DEMts=0.00001s, couplingInterval=10 exchange data (=couple) will happen every 0.0001s.

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## 1.5 "liggghtsCommands" dictionary

### General:

In the "liggghtsCommands" dictionary liggghts commands being executed during a coupled CFD-DEM simulation are specified.

**Location:** \$caseDir/CFD/constant

### Structure:

The dictionary is divided into two parts, first a list of "liggghtsCommandModels" is defined, then the settings for each model must be specified.

### Settings:

Reasonable example settings for the "liggghtsCommands" dictionary are given in the tutorial cases.

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## 1.6 Models/Solvers

This section lists all CFDEMcoupling sub-models and solvers alphabetically, with a separate listing below of styles within certain commands.

<a href="#"><u>IOModel</u></a>	<a href="#"><u>IOModel basicIO</u></a>
<a href="#"><u>IOModel noIO</u></a>	<a href="#"><u>IOModel trackIO</u></a>
<a href="#"><u>averagingModel</u></a>	<a href="#"><u>averagingModel dilute</u></a>
<a href="#"><u>cfdemSolverIB</u></a>	<a href="#"><u>cfdemSolverPiso</u></a>
<a href="#"><u>cfdemSolverPisoScalar</u></a>	<a href="#"><u>clockModel</u></a>
<a href="#"><u>clockModel noClock</u></a>	<a href="#"><u>clockModel standardClock</u></a>
<a href="#"><u>dataExchangeModel</u></a>	<a href="#"><u>dataExchangeModel noDataExchange</u></a>
<a href="#"><u>dataExchangeModel oneWayVTK</u></a>	<a href="#"><u>dataExchangeModel twoWayFiles</u></a>
<a href="#"><u>dataExchangeModel twoWayMPI</u></a>	<a href="#"><u>forceModel</u></a>
<a href="#"><u>forceModel Archimedes</u></a>	<a href="#"><u>forceModel ArchimedesIB</u></a>
<a href="#"><u>forceModel DiFeliceDrag</u></a>	<a href="#"><u>forceModel GidaspowDrag</u></a>
<a href="#"><u>forceModel KochHillDrag</u></a>	<a href="#"><u>forceModel LaEuScalarTemp</u></a>
<a href="#"><u>forceModel MeiLift</u></a>	<a href="#"><u>forceModel SchillerNaumannDrag</u></a>
<a href="#"><u>forceModel ShirgaonkarIB</u></a>	<a href="#"><u>forceModel gradPForce</u></a>
<a href="#"><u>forceModel noDrag</u></a>	<a href="#"><u>forceModel virtualMassForce</u></a>
<a href="#"><u>forceModel viscForce</u></a>	<a href="#"><u>liggghtsCommandModel</u></a>
<a href="#"><u>liggghtsCommandModel execute</u></a>	<a href="#"><u>liggghtsCommandModel readLiggghtsData</u></a>
<a href="#"><u>liggghtsCommandModel runLiggghts</u></a>	<a href="#"><u>liggghtsCommandModel writeLiggghts</u></a>
<a href="#"><u>locateModel</u></a>	<a href="#"><u>locateModel engineSearch</u></a>
<a href="#"><u>locateModel engineSearchIB</u></a>	<a href="#"><u>locateModel standardSearch</u></a>
<a href="#"><u>locateModel turboEngineSearch</u></a>	<a href="#"><u>meshMotionModel</u></a>
<a href="#"><u>meshMotionModel noMeshMotion</u></a>	<a href="#"><u>momCoupleModel</u></a>
<a href="#"><u>momCoupleModel explicitCouple</u></a>	<a href="#"><u>momCoupleModel implicitCouple</u></a>
<a href="#"><u>momCoupleModel noCouple</u></a>	<a href="#"><u>regionModel</u></a>
<a href="#"><u>regionModel allRegion</u></a>	<a href="#"><u>voidfractionModel</u></a>
<a href="#"><u>voidfractionModel GaussVoidFraction</u></a>	<a href="#"><u>voidfractionModel IBVoidFraction</u></a>
<a href="#"><u>voidfractionModel bigParticleVoidFraction</u></a>	<a href="#"><u>voidfractionModel centreVoidFraction</u></a>
<a href="#"><u>voidfractionModel dividedVoidFraction</u></a>	

## cfdemSolverIB command

### Description:

"cfdemSolverIB" is a coupled CFD-DEM solver using CFDEMcoupling, an open source parallel coupled CFD-DEM framework, for calculating the dynamics between immersed bodies and the surrounding fluid. Being an implementation of an immersed boundary method it allows tackling problems where the body diameter exceeds the maximal size of a fluid cell. Using the toolbox of OpenFOAM(R)(\*) the governing equations of the fluid are computed and the corrections of velocity and pressure field with respect to the body-movement information, gained from LIGGGHTS, are incorporated.

Code of this solver contributions by Alice Hager, JKU.

### Algorithm:

For each time step ...

- the motion of the spheres is calculated (position, velocity, angular velocity, force...) with LIGGGHTS using the velocity and pressure-field from the previous time step (initial condition for  $t=0$ ).
- the Navier-Stokes equations are solved on the whole computational domain, disregarding the solid phase.
- the spheres are located within the mesh: each sphere is represented by a cluster of cells, which are either totally or partially covered by the body, depending on its exact position.
- the correction of the velocity and pressure field of the fluid phase takes place, using the information about the location of the spheres and their (angular) velocity.

### Use:

The solver is realized within the Open Source framework CFDEMcoupling. Just as for the unresolved CFD-DEM solver cfdemSolverPiso the file CFD/constant/couplingProperties contains information about the settings for the different models. While IOmodel, DataExchangeModel etc. are applicable for all CFDEMcoupling-solvers, special locate-, force- and void fraction models were designed for the present case:

[engineSearchIB](#), [ArchimedesIB](#), [ShirgaonkarIB](#), [IBVoidfraction](#)

### References:

GONIVA, C., KLOSS, C., HAGER, A., WIERINK, G. and PIRKER, S. (2011): "A MULTI-PURPOSE OPEN SOURCE CFD-DEM APPROACH", Proc. of the 8th Int. Conf. on CFD in Oil and Gas, Metallurgical and Process Industries, Trondheim, Norway

and

HAGER, A., KLOSS, C. and GONIVA, C. (2011): "TOWARDS AN EFFICIENT IMMERSSED BOUNDARY METHOD WITHIN AN OPEN SOURCE FRAMEWORK", Proc. of the 8th Int. Conf. on CFD in Oil and Gas, Metallurgical and Process Industries, Trondheim, Norway

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## cfdemSolverPiso command

### Description:

"cfdemSolverPiso" is a coupled CFD-DEM solver using CFDEMcoupling, an open source parallel coupled CFD-DEM framework. Based on pisoFoam(R)(\*), a finite volume based solver for turbulent Navier-Stokes equations applying PISO algorithm, "cfdemSolverPiso" has additional functionality for a coupling to the DEM code "LIGGGHTS". The volume averaged Navier-Stokes Equations are solved accounting for momentum exchange and volume displacement of discrete particles whose trajectories are calculated in the DEM code LIGGGHTS.

see:

GONIVA, C., KLOSS, C., HAGER, A. and PIRKER, S. (2010): "An Open Source CFD-DEM Perspective", Proc. of OpenFOAM Workshop, Göteborg, June 22.-24.

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## **cfdemSolverPisoScalar command**

### **Description:**

"cfdemSolverPisoScalar" is a coupled CFD-DEM solver using CFDEMcoupling, an open source parallel coupled CFD-DEM framework. Based on pisoFoam(R)(\*), a finite volume based solver for turbulent Navier-Stokes equations applying PISO algorithm, "cfdemSolverPisoScalar" has additional functionality for a coupling to the DEM code "LIGGGHTS" as well as a scalar transport equation. The volume averaged Navier-Stokes Equations are solved accounting for momentum exchange and volume displacement of discrete particles whose trajectories are calculated in the DEM code LIGGGHTS. The scalar transport equation is coupled to scalar properties of the particle phase, thus convective heat transfer in a fluid granular system can be modeled with "cfdemSolverPisoScalar".

see:

GONIVA, C., KLOSS, C., HAGER, A. and PIRKER, S. (2010): "An Open Source CFD-DEM Perspective", Proc. of OpenFOAM Workshop, Göteborg, June 22.-24.

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## clockModel command

### Syntax:

Defined in couplingProperties dictionary.

```
clockModel model;
```

- model = name of the clockModel to be applied

### Examples:

```
clockModel standardClock;
```

Note: This examples list might not be complete - please look for other models (clockModel\_XY) in this documentation.

### Description:

The clockModel is the base class for models to examine the code/algorithm with respect to run time.

Main parts of the clockModel classes are written by Josef Kerbl, JKU.

**Restrictions:** none.

**Default:** none.

## clockModel\_noClock command

### Syntax:

Defined in couplingProperties dictionary.

```
clockModel off;
```

### Examples:

```
clockModel off;
```

### Description:

The "noClock" model is a dummy clockModel model which does not measure/evaluate the run time.

**Restrictions:** none.

### Related commands:

[clockModel](#)

## clockModel\_standardClock command

### Syntax:

Defined in couplingProperties dictionary.

```
clockModel standardClock;
```

### Examples:

```
clockModel standardClock;
```

### Description:

The "standardClock" model is a basic clockModel model which measures the run time between every ".start(int arrayPos,string name)" and ".stop(string name)" statement placed in the code. If a ".start(name)" is called more than once (e.g. in a loop) the accumulated times are calculated. After the simulation has finished, the data is stored in \$caseDir/CFD/clockData/\$startTime/\*.txt . Since the measurements are stored in an array, it is necessary to put a variable *arrayPos* (type integer) at the start command. Those do not need to be in ascending order and positions may be omitted. The standard size of this array is 30 and can be changed at the initialization of the standardClock class. If *arrayPos* is out of bounds, the array size will be doubled. The stop command does not need *arrayPos*, since the class remembers the positions. The string name is for easier evaluation afterwards and may be omitted like ".start(int arrayPos)" and ".stop()". The command ".stop(string name)" is a safety feature, because if the name is not equal to the started name, output will be produced for information. After the case ran you may use the matPlot.py script located in \$CFDEM\_UT\_DIR/vizClock/ to produce a graphical output of your measurements. The usage is like 'python < matPlot.py' and you have to be in the directory of the desired time step, where there is a file called "timeEvalFull.txt", which contains averaged and maximum data with respect to the number of processes.

**Restrictions:** none.

**Related commands:**

[clockModel](#)

## **dataExchangeModel command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
dataExchangeModel model;
```

- model = name of data exchange model to be applied

### **Examples:**

```
dataExchangeModel twoWayFiles;  
dataExchangeModel twoWayMPI;
```

Note: This examples list might not be complete - please look for other models (dataExchangeModel\_XY) in this documentation.

### **Description:**

The data exchange model performs the data exchange between the DEM code and the CFD code.

### **Restrictions:**

None.

### **Related commands:**

[noDataExchange](#), [oneWayVTK](#), [twoWayFiles](#), [twoWayMPI](#)

**Default:** none

## **dataExchangeModel\_noDataExchange command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
dataExchangeModel noDataExchange;
```

### **Examples:**

```
dataExchangeModel noDataExchange;
```

### **Description:**

The data exchange model performs the data exchange between the DEM code and the CFD code. The noDataExchange model is a dummy model where no data is exchanged.

### **Restrictions:**

None.

### **Related commands:**

[dataExchangeModel](#)

## dataExchangeModel\_oneWayVTK command

### Syntax:

Defined in couplingProperties dictionary.

```
dataExchangeModel oneWayVTK;  
oneWayVTKProps  
{  
    DEMts timeStep;  
    relativePath "path";  
    couplingFilename "filename";  
    maxNumberOfParticles number;  
};
```

- *timeStep* = time step size of stored DEM data
- *path* = path to the VTK data files relative to simulation directory
- *filename* = filename of the VTK file series
- *number* = maximum number of particles in DEM simulation

### Examples:

```
dataExchangeModel oneWayVTK;  
oneWayVTKProps  
{  
    DEMts 0.0001;  
    relativePath "../DEM/post";  
    couplingFilename "vtk_out%4.4d.vtk";  
    maxNumberOfParticles 30000;  
}
```

### Description:

The data exchange model performs the data exchange between the DEM code and the CFD code. The oneWayVTK model is a model that can exchange particle properties from DEM to CFD based on previously stored VTK data.

### Restrictions:

None.

### Related commands:

[dataExchangeModel](#)

## dataExchangeModel\_twoWayFiles command

### Syntax:

Defined in couplingProperties dictionary.

```
dataExchangeModel twoWayFiles;  
twoWayFilesProps  
{  
    couplingFilename "filename";  
    maxNumberOfParticles number;  
};
```

- *filename* = filename of the VTK file series
- *number* = maximum number of particles in DEM simulation

### Examples:

```
dataExchangeModel twoWayFiles;  
twoWayFilesProps  
{  
    couplingFilename "vtk_out%4.4d.vtk";  
    maxNumberOfParticles 30000;  
}
```

### Description:

The data exchange model performs the data exchange between the DEM code and the CFD code. The twoWayFiles model is a model that can exchange particle properties from DEM to CFD and from CFD to DEM. Data is exchanged via files that are sequentially written/read by the codes.

### Restrictions:

Developed only for two processors, one for DEM and one for CFD run.

### Related commands:

[dataExchangeModel](#)



## dataExchangeModel\_twoWayMPI command

### Syntax:

Defined in couplingProperties dictionary.

```
dataExchangeModel twoWayMPI;  
twoWayMPIProps  
{  
    liggghtsPath "path";  
};
```

- *path* = path to the DEM simulation input file

### Examples:

```
dataExchangeModel twoWayMPI;  
twoWayMPIProps  
{  
    liggghtsPath "../DEM/in.liggghts_init";  
}
```

### Description:

The data exchange model performs the data exchange between the DEM code and the CFD code. The twoWayMPI model is a model that can exchange particle properties from DEM to CFD and from CFD to DEM. Data is exchanged via MPI technique. The DEM run is executed by the coupling model, via a liggghtsCommandModel object.

### Restrictions:

none.

### Related commands:

[dataExchangeModel](#)

## forceModel\_Archimedes command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    Archimedes
);
ArchimedesProps
{
    densityFieldName "density";
    gravityFieldName "gravity";
};
```

- *density* = name of the finite volume density field
- *gravity* = name of the finite volume gravity field

### Examples:

```
forceModels
(
    Archimedes
);
ArchimedesProps
{
    densityFieldName "rho";
    gravityFieldName "g";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The Archimedes model is a model that calculates the Archimedes' volumetric lift force stemming from density difference of fluid and particle.

### Restrictions:

none.

### Related commands:

[forceModel](#)

## forceModel\_ArchimedesIB command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    ArchimedesIB
);
ArchimedesIBProps
{
    densityFieldName "density";
    gravityFieldName "gravity";
    voidfractionFieldName "voidfraction";
};
```

- *density* = name of the finite volume density field
- *gravity* = name of the finite volume gravity field
- *voidfraction* = name of the finite volume voidfraction field

### Examples:

```
forceModels
(
    ArchimedesIB
);
ArchimedesIBProps
{
    densityFieldName "rho";
    gravityFieldName "g";
    voidfractionFieldName "voidfractionNext";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The ArchimedesIB model is a model that calculates the ArchimedesIB' volumetric lift force stemming from density difference of fluid and particle. This model is especially suited for resolved CFD-DEM simulations where the particle is represented by immersed boundary method.

### Restrictions:

Only for immersed boundary solvers.

### Related commands:

[forceModel](#)

## forceModel\_DiFeliceDrag command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    DiFeliceDrag
);
DiFeliceDragProps
{
    velFieldName "U";
    densityFieldName "density";
    interpolation;
};
```

- *U* = name of the finite volume fluid velocity field
- *density* = name of the finite volume gravity field
- *interpolation* = flag to use interpolate interpolated voidfraction and velocity values (normally off)

### Examples:

```
forceModels
(
    DiFeliceDrag
);
DiFeliceDragProps
{
    velFieldName "U";
    densityFieldName "rho";
    interpolation;
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The DiFeliceDrag model is a model that calculates the particle based drag force following the correlation of Di Felice (see Zhou et al. (2010), JFM).

### Restrictions:

none.

### Related commands:

[forceModel](#)

## forceModel\_GidaspowDrag command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    GidaspowDrag
);
GidaspowDragProps
{
    velFieldName "U";
    densityFieldName "density";
};
```

- $U$  = name of the finite volume fluid velocity field
- *density* = name of the finite volume gravity field

### Examples:

```
forceModels
(
    GidaspowDrag
);
GidaspowDragProps
{
    velFieldName "U";
    densityFieldName "rho";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The GidaspowDrag model is a model that calculates the particle based drag force following the correlation of Gidaspow which is a combination of Egrun (1952) and Wen & Yu (1966) (see Zhu et al. (2007): "Discrete particle simulation of particulate systems: Theoretical developments" ,ChemEngScience).

### Restrictions:

none.

### Related commands:

[forceModel](#)

## forceModel\_gradPForce command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    gradPForce;
);
gradPForceProps
{
    pFieldName "pressure";
    densityFieldName "density";
    velocityFieldName "U";
    interpolation;
};
```

- *pressure* = name of the finite volume fluid pressure field
- *density* = name of the finite volume gravity field
- *U* = name of the finite volume fluid velocity field
- *interpolation* = flag to use interpolate interpolated pressure values (normally off)

### Examples:

```
forceModels
(
    gradPForce;
);
gradPForceProps
{
    pFieldName "p";
    densityFieldName "rho";
    velocityFieldName "U";
    interpolation;
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The gradPForce model is a model that calculates the particle based pressure gradient force  $-(\text{grad}(p)) * V_{\text{particle}}$  (see Zhou et al. (2010): "Discrete particle simulation of particle-fluid flow: model formulations and their applicability" ,JFM).

### Restrictions:

none.

### Related commands:

[forceModel](#)

## forceModel command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    model_x
    model_y
);
```

- model = name of force model to be applied

### Examples:

```
forceModels
(
    Archimedes
    DiFeliceDrag
);
```

Note: This examples list might not be complete - please look for other models (forceModel\_XY) in this documentation.

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. All force models selected are executed sequentially and the forces on the particles are superposed.

### Restrictions:

None.

### Related commands:

[Archimedes](#), [DiFeliceDrag](#), [gradPForce](#), [viscForce](#)

Note: This examples list may be incomplete - please look for other models (forceModel\_XY) in this documentation.

**Default:** none.

## forceModel\_KochHillDrag command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    KochHillDrag
);
KochHillDragProps
{
    velFieldName "U";
    densityFieldName "density";
    voidfractionFieldName "voidfraction";
    interpolation;
};
```

- *U* = name of the finite volume fluid velocity field
- *density* = name of the finite volume gravity field
- *voidfraction* = name of the finite volume voidfraction field
- *interpolation* = flag to use interpolated voidfraction and fluid velocity values (normally off)

### Examples:

```
forceModels
(
    KochHillDrag
);
KochHillDragProps
{
    velFieldName "U";
    densityFieldName "rho";
    voidfractionFieldName "voidfraction";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The KochHillDrag model is a model that calculates the particle based drag force following the correlation of Koch & Hill (2001) (see van Buijtenen et al. (2011): "Numerical and experimental study on multiple-spout fluidized beds" ,ChemEngScience).

### Restrictions:

none.

### Related commands:

[forceModel](#)



## forceModel\_LaEuScalarTemp command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    LaEuScalarTemp
);
LaEuScalarTempProps
{
    velFieldName "U";
    tempFieldName "T";
    tempSourceFieldName "Tsource";
    voidfractionFieldName "voidfraction";
    partTempName "Temp";
    partHeatFluxName "convectiveHeatFlux";
    lambda value;
    Cp value1;
    densityFieldName "density";
};
```

- *U* = name of the finite volume fluid velocity field
- *T* = name of the finite volume scalar temperature field
- *Tsource* = name of the finite volume scalar temperature source field
- *voidfraction* = name of the finite volume voidfraction field
- *Temp* = name of the DEM data representing the particles temperature
- *convectiveHeatFlux* = name of the DEM data representing the particle-fluid convective heat flux
- *value* = fluid thermal conductivity [W/(m\*K)]
- *value1* = fluid specific heat capacity [W\*s/(kg\*K)]
- *density* = name of the finite volume fluid density field

### Examples:

```
forceModels
(
    LaEuScalarTemp
);
LaEuScalarTempProps
{
    velFieldName "U";
    tempFieldName "T";
    tempSourceFieldName "Tsource";
    voidfractionFieldName "voidfraction";
    partTempName "Temp";
    partHeatFluxName "convectiveHeatFlux";
    lambda 0.0256;
    Cp 1007;
    densityFieldName "rho";
}
```

### Description:

This "forceModel" does not influence the particles or the fluid flow! Using the particles' temperature a scalar field representing "particle-fluid heatflux" is calculated. The solver then uses this source field in the scalar transport equation for the temperature. The model for convective heat transfer is based on Li and Mason (2000), A computational investigation of transient heat transfer in pneumatic transport of granular particles, Pow.Tech 112

**Restrictions:**

Goes only with cfemSolverScalar.

**Related commands:**

[forceModel](#)

## forceModel\_MeiLift command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    MeiLift
);
MeiLiftProps
{
    velFieldName "U";
    densityFieldName "density";
};
```

- $U$  = name of the finite volume fluid velocity field
- *density* = name of the finite volume fluid density field

### Examples:

```
forceModels
(
    MeiLift
);
MeiLiftProps
{
    velFieldName "U";
    densityFieldName "rho";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The MeiLift model calculates the lift force for each particle based on Loth and Dorgan (2009)

### Restrictions:

None.

### Related commands:

[forceModel](#)

## **forceModel\_noDrag command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
forceModels
(
    off
);
```

### **Examples:**

```
forceModels
(
    off
);
```

### **Description:**

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The noDrag model sets the forces acting on the particle to zero. If several force models are selected and noDrag is the last model being executed, the fluid particle force will be set to zero.

### **Restrictions:**

None.

### **Related commands:**

[forceModel](#)

## forceModel\_SchillerNaumannDrag command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    SchillerNaumannDrag
);
SchillerNaumannDragProps
{
    velFieldName "U";
    densityFieldName "density";
};
```

- $U$  = name of the finite volume fluid velocity field
- *density* = name of the finite volume gravity field

### Examples:

```
forceModels
(
    SchillerNaumannDrag
);
SchillerNaumannDragProps
{
    velFieldName "U";
    densityFieldName "rho";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The SchillerNaumannDrag model is a model that calculates the particle based drag force following the correlation of Schiller and Naumann.

### Restrictions:

none.

### Related commands:

[forceModel](#)

## forceModel\_ShirgaonkarIB command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    ShirgaonkarIB
);
ShirgaonkarIBProps
{
    velFieldName "U";
    densityFieldName "density";
    pressureFieldName "pressure";
};
```

- *U* = name of the finite volume fluid velocity field
- *density* = name of the finite volume density field
- *pressure* = name of the finite volume pressure field

### Examples:

```
forceModels
(
    ShirgaonkarIB
);
ShirgaonkarIBProps
{
    velFieldName "U";
    densityFieldName "rho";
    pressureFieldName "p";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The ShirgaonkarIB model calculates the drag force (viscous and pressure force) acting on each particle in a resolved manner (see Shirgaonkar et al. (2009): "A new mathematical formulation and fast algorithm for fully resolved simulation of self-propulsion", Journal of Comp. Physics). This model is only suited for resolved CFD-DEM simulations where the particle is represented by immersed boundary method.

### References:

SHIRGAONKAR, A.A., MACIVER, M.A. and PATANKAR, N.A., (2009), "A new mathematical formulation and fast algorithm for fully resolved simulation of self-propulsion", J. Comput. Phys., 228, 2366-2390.

### Restrictions:

Only for immersed boundary solvers.

**Related commands:**

[forceModel](#)

## forceModel\_virtualMassForce command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    virtualMassForce
);
virtualMassForceProps
{
    velFieldName "U";
    densityFieldName "density";
};
```

- $U$  = name of the finite volume fluid velocity field
- *density* = name of the finite volume fluid density field

### Examples:

```
forceModels
(
    virtualMassForce
);
virtualMassForceProps
{
    velFieldName "U";
    densityFieldName "rho";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The virtualMassForce model calculates the virtual mass force for each particle.

### Restrictions:

Model not validated!

### Related commands:

[forceModel](#)



## forceModel\_viscForce command

### Syntax:

Defined in couplingProperties dictionary.

```
forceModels
(
    viscForce;
);
viscForceProps
{
    velocityFieldName "U";
    densityFieldName "density";
    interpolation;
};
```

- *U* = name of the finite volume fluid velocity field
- *density* = name of the finite volume gravity field
- *interpolation* = flag to use interpolate interpolated stress values (normally off)

### Examples:

```
forceModels
(
    viscForce;
);
viscForceProps
{
    velocityFieldName "U";
    densityFieldName "density";
}
```

### Description:

The force model performs the calculation of forces (e.g. fluid-particle interaction forces) acting on each DEM particle. The viscForce model calculates the particle based viscous force,  $-(\text{grad}(\tau)) * V_{\text{particle}}$  (see Zhou et al. (2010): "Discrete particle simulation of particle-fluid flow: model formulations and their applicability", JFM).

### Restrictions:

none.

### Related commands:

[forceModel](#)

## githubAccess\_public

---

### Description:

This routine describes how to setup a github account and pull repositories of the CFDEMproject. After setting some environment variables LIGGGHTS and CFDEMcoupling can be compiled

### Procedure:

Basically the following steps have to be performed:

- *git clone* the desired repository
- update your repositories by *git pull*
- set environment variables
- compile LIGGGHTS and CFDEMcoupling
- run your own cases

### *git clone* the desired repository:

If not already done, open a terminal and create a directory for LIGGGHTS in \$HOME:

```
cd  
  
mkdir LIGGGHTS  
  
cd LIGGGHTS
```

To clone the public LIGGGHTS repository, open a terminal and execute:

```
git clone git://github.com/CFDEMproject/LIGGGHTS-PUBLIC.git LIGGGHTS-PUBLIC
```

If not already done, open a terminal and create a directory for CFDEMcoupling in \$HOME:

```
cd  
  
mkdir CFDEM  
  
cd CFDEM
```

Make sure that OpenFOAM(R)-2.1.x is already set up correctly!

To clone the public CFDEMcoupling repository, open a terminal and execute:

```
git clone git://github.com/CFDEMproject/CFDEMcoupling-PUBLIC.git CFDEMcoupling-PUBLIC-$WM_PROJECT_VERSION
```

Troubles? See Troubleshooting section below.

### Update your repositories by *git pull*:

To get the latest version, open a terminal, go to the location of your local installation and type: *Warning: git stash will remove your changes in \$HOME/CFDEM/CFDEMcoupling-PUBLIC-\$WM\_PROJECT\_VERSION !*

```
cd $HOME/CFDEM/CFDEMcoupling-PUBLIC-$WM_PROJECT_VERSION
git stash
git pull
```

### Set Environment Variables:

Now you need to set some environment variables in ~/.bashrc (if you use c-shell, manipulate ~/.cshrc accordingly). Open ~/.bashrc

```
gedit ~/.bashrc &
```

add the lines (you find them also in .../cfdemParticle/etc/bashrc and cshrc respectively):

```
#=====#
#- source cfdem env vars
export CFDEM_VERSION=PUBLIC
export CFDEM_PROJECT_DIR=$HOME/CFDEM/CFDEMcoupling-$CFDEM_VERSION-$WM_PROJECT_VERSION
export CFDEM_SRC_DIR=$CFDEM_PROJECT_DIR/src/laagrangian/cfdemParticle
export CFDEM_SOLVER_DIR=$CFDEM_PROJECT_DIR/applications/solvers
export CFDEM_DOC_DIR=$CFDEM_PROJECT_DIR/doc
export CFDEM_UT_DIR=$CFDEM_PROJECT_DIR/applications/utilities
export CFDEM_TUT_DIR=$CFDEM_PROJECT_DIR/tutorials
export CFDEM_PROJECT_USER_DIR=$HOME/CFDEM/$LOGNAME-$CFDEM_VERSION-$WM_PROJECT_VERSION
export CFDEM_bashrc=$CFDEM_SRC_DIR/etc/bashrc
export CFDEM_LIGGGHTS_SRC_DIR=$HOME/LIGGGHTS/LIGGGHTS-PUBLIC/src
export CFDEM_LIGGGHTS_MAKEFILE_NAME=fedora_fpic
export CFDEM_LPP_DIR=$HOME/LIGGGHTS/mylpp/src
export CFDEM_PIZZA_DIR=$HOME/LIGGGHTS/PIZZA/gran_pizza_17Aug10/src
. $CFDEM_bashrc
#=====#
```

Save the ~/.bashrc, open a new terminal and test the settings. The commands:

```
$CFDEM_PROJECT_DIR
$CFDEM_SRC_DIR
$CFDEM_LIGGGHTS_SRC_DIR
```

should give "...: is a directory" otherwise something went wrong and the environment variables in ~/.bashrc are not set correctly.

To specify the paths of pizza, please check the settings in \$CFDEM\_SRC\_DIR/etc/bashrc.

If \$CFDEM\_SRC\_DIR is set correctly, you can type

```
cfdemSysTest
```

to get some information if the paths are set correctly.

### Compile LIGGGHTS and CFDEMcoupling:

If above settings were done correctly, you can compile LIGGGHTS by typing:

```
git clone git://github.com/CFDEMproject/CFDEMcoupling-PUBLIC.git CFDEMcoupling-PUBLIC-$WM_PROJ
```

```
cfdemCompLIG
```

and you can then compile CFDEMcoupling by typing:

```
cfdemCompCFDEM
```

You can run the tutorial cases by executing `../etc/testTutorial.sh` through the alias `cfdemTestTUT`. Alternatively you can run each tutorial using the `Allrun.sh` scripts in the tutorial directories.

In case questions concerning the installation arise, please feel free to contact our forum at [www.cfdem.com](http://www.cfdem.com).

### Run Your Own Cases:

If you want to run your own cases, please do so in `$CFDEM_PROJECT_USER_DIR/run` which is automatically being generated. E.g. copy one of the tutorial cases there, adapt it to your needs. Changes in `$CFDEM_TUT_DIR` will be lost after every `git stash`!

### Additional Installations:

Optionally you can install `lpp` which will help you convert the DEM (dump) data to VTK format. For standard CFD-DEM runs this will not be necessary. To get the DEM postprocessing tool "lpp" you need python-numpy package installed:

```
sudo apt-get install python-numpy
```

You can pull the latest version of `lpp` with:

```
cd $HOME/LIGGGHTS
```

```
git clone git://cfdem.git.sourceforge.net/gitroot/cfdem/lpp mylpp
```

### Troubleshooting:

- troubles with git clone?

**a)** The git protocol will not work if your computer is behind a firewall which blocks the relevant TCP port, you can use alternatively (write command in one line):

```
git clone https://user@github.com/CFDEMproject/CFDEMcoupling-PUBLIC.git  
CFDEMcoupling-PUBLIC-$WM_PROJECT_VERSION
```

**b)** If you face the error: "error: SSL certificate problem, verify that the CA cert is OK. Details: error:14090086:SSL routines:SSL3\_GET\_SERVER\_CERTIFICATE:certificate verify failed while accessing https://github.com/...",

please use: `env GIT_SSL_NO_VERIFY=true git clone https://github...`

(see <http://stackoverflow.com/questions/3777075/https-github-access>)

```
git clone git://github.com/CFDEMproject/CFDEMcoupling-PUBLIC.git CFDEMcoupling-PUBLIC-$WM_PROJECT_VERSION
```

## **IOModel\_basicIO command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
IOModel "basicIO";
```

### **Examples:**

```
IOModel "basicIO";
```

### **Description:**

The basic IO-model writes particle positions velocities and radii to files. The output directory (\$casePath/CFD/particles) is created automatically. Data is written every write time of the CFD simulation.

**Restrictions:** None.

### **Related commands:**

[IOModel](#)

## IOModel command

### Syntax:

Defined in couplingProperties dictionary.

```
IOModel "model";
```

- model = name of IO-model to be applied

### Examples:

```
IOModel "off";
```

Note: This examples list might not be complete - please look for other models (IOModel\_XY) in this documentation.

### Description:

The IO-model is the base class to write data (e.g. particle properties) to files.

### Restrictions:

none.

### Related commands:

Note: This examples list may be incomplete - please look for other models (IOModel\_XY) in this documentation.

**Default:** none.

## **IOModel\_noIO command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
IOModel "off";
```

### **Examples:**

```
IOModel "off";
```

### **Description:**

The noIO-model is a dummy IO model.

**Restrictions:** None.

### **Related commands:**

[IOModel](#)

## **IOModel\_trackIO command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
IOModel "trackIO";
```

### **Examples:**

```
IOModel "trackIO";
```

### **Description:**

The basic IO-model writes particle positions, velocities, index and radii to files. The output directories (\$casePath/CFD/\$timestep) are created automatically. Data is written every write time of the CFD simulation.

**Restrictions:** None.

**Related commands:**

[IOModel](#)



## liggghtsCommandModel\_execute command

### Syntax:

Defined in liggghtsCommmands dictionary.

```
liggghtsCommandModels
(
    execute
);
executeProps0
{
    command
    (
        run
        $couplingInterval
    );
    runFirst switch1;
    runLast switch2;
    runEveryCouplingStep switch3;
    runEveryWriteStep switch4;
}
```

- *command* = LIGGGHTS command to be executed. Each word in a new line, numbers and symbols need special treatment (e.g. \$couplingInterval will be replaced by correct coupling interval in the simulation)
- *switch1* = switch (choose on/off) if the command is executed only at first time step
- *switch2* = switch (choose on/off) if the command is executed only at last time step
- *switch3* = switch (choose on/off) if the command is executed at every coupling step
- *switch4* = switch (choose on/off) if the command is executed at every writing step

### Examples:

```
liggghtsCommandModels
(
    execute
    execute
);
executeProps0
{
    command
    (
        run
        $couplingInterval
    );
    runFirst off;
    runLast off;
    runEveryCouplingStep on;
}
executeProps1
{
    command
    (
        write_restart
        noBlanks
        dotdot
    )
}
```

```

        slash
        DEM
        slash
        liggghts.restart_
        timeStamp
    );
    runFirst off;
    runLast off;
    runEveryCouplingStep off;
    runEveryWriteStep on;
}

```

### **Description:**

The execute liggghtsCommand Model can be used to execute a LIGGGHTS command during a CFD run. In above example execute\_0 for instance executes "run \$couplingInterval" every coupling step. \$couplingInterval is automatically replaced by the correct number of DEM steps. Additionally execute\_1 executes "write\_restart ../DEM/liggghts.restart\_\$timeStamp" every writing step, where \$timeStamp is automatically set.

**These rather complex execute commands can be replaced by the "readLiggghts" and "writeLiggghts" commands!**

**Restrictions:** None.

**Related commands:**

[liggghtsCommandModel](#)

## liggghtsCommandModel command

### Syntax:

Defined in liggghtsCommmands dictionary.

```
liggghtsCommandModels
(
    model_x
    model_y
);
```

- model = name of the liggghtsCommandModel to be applied

### Examples:

```
liggghtsCommandModels
(
    runLiggghts
    writeLiggghts
);
```

Note: This examples list might not be complete - please look for other models (liggghtsCommandModel\_XY) in this documentation.

### Description:

The liggghtsCommandModel is the base class to execute DEM commands within a CFD run.

### Restrictions:

Works only with MPI coupling.

**Default:** none.

## liggghtsCommandModel\_readLiggghtsData command

### Syntax:

Defined in liggghtsCommmands dictionary.

```
liggghtsCommandModels
(
    readLiggghtsData
);
readLiggghtsDataProps0
{
    ???
}
```

### Examples:

```
liggghtsCommandModels
(
    readLiggghtsData
    readLiggghtsData
);
readLiggghtsDataProps0
{
    ???
}
```

### Description:

The readLiggghtsData liggghtsCommand Model can be used to ???

### Restrictions:

Note: Model is not up to date.

### Related commands:

[liggghtsCommandModel](#)

## liggghtsCommandModel\_runLiggghts command

### Syntax:

Defined in liggghtsCommmands dictionary.

```
liggghtsCommandModels
(
    runLiggghts
);
//- optional
runLiggghtsProps
{
    preNo true;
}
```

### Examples:

```
liggghtsCommandModels
(
    runLiggghts
);
```

### Description:

The liggghtsCommand models can be used to execute a LIGGGHTS command during a CFD run. The "runLiggghts" command executes the command "run \$nrDEMsteps", where \$nrDEMsteps is automatically set according to the coupling intervals, every coupling step. Optionally a dictionary called runLiggghtsProps can be specified where the "preNo" switch can be set, which uses the command "run \$nrDEMsteps pre no" for every time step except the first.

**Restrictions:** Warning: the "pre no" option can cause troubles (dump data of particles changin the domain might be erroneous)!

### Related commands:

[liggghtsCommandModel](#)

## liggghtsCommandModel\_writeLiggghts command

### Syntax:

Defined in liggghtsCommmands dictionary.

```
liggghtsCommandModels
(
    writeLiggghts
);
//- optional
writeLiggghtsProps
{
    writeLast switch1;
    writeName "name";
    overwrite switch2;
}
```

- *switch1* = switch (choose on/off) to select if only last step is stored or every write step. "off" is not recommended (DEM data might get lost)
- *name* = name of the restart file to be written in /\$caseDir/DEM/ default default "liggghts.restartCFDEM"
- *switch2* = switch (choose on/off) to select if only one restart file \$name or many files \$name\_\$timeStamp are written

### Examples:

```
liggghtsCommandModels
(
    runLiggghts
    writeLiggghts
);
```

### Description:

The liggghtsCommand models can be used to execute a LIGGGHTS command during a CFD write. The "writeLiggghts" command executes the command "write\_restart \$name", where \$name is the name of the restart file, every write step.

**Restrictions:** None.

**Related commands:**

[liggghtsCommandModel](#)

## locateModel\_engineSearch command

### Syntax:

Defined in couplingProperties dictionary.

```
locateModel engine;  
engineProps  
{  
    treeSearch switch1;  
}
```

- *switch1* = switch to use tree search algorithm

### Examples:

```
locateModel engine;  
engineProps  
{  
    treeSearch false;  
}
```

### Description:

The locateModel "engine" locates the CFD cell and cellID corresponding to a given position. The engineSearch locate Model can be used with different settings to use different algorithms:

- treeSearch false; will execute some geometric (linear) search using the last known cellID (recommended)
- treeSearch true; will use a recursive tree structure to find the cell.

**Restrictions:** none.

### Related commands:

[locateModel](#)

## locateModel\_engineSearchIB command

### Syntax:

Defined in couplingProperties dictionary.

```
locateModel engineIB;
engineIBProps
{
    engineProps
    {
        treeSearch switch1;
    }
    zSplit value1;
    xySplit value2;
}
```

- *switch1* = names of the finite volume scalar fields to be temporally averaged
- *value1* = number of z-normal layers for satellite points
- *value2* = number of satellite points in each layer

### Examples:

```
locateModel engineIB;
engineIBProps
{
    engineProps
    {
        treeSearch false;
    }
    zSplit 8;
    xySplit 16;
}
```

### Description:

The locateModel "engine" locates the CFD cell and cellID corresponding to a given position. This locate model is especially designed for parallel immersed boundary method. Each particle is represented by "satellite points" if it is distributed over several processors.

The engineSearchIB locate Model can be used with different settings to use different algorithms:

- treeSearch false; will execute some geometric (linear) search using the last known cellID (recommended)
- treeSearch true; will use a recursive tree structure to find the cell.

This model is a modification of the engine search model. Instead of using the centre-cell as starting point for the engine search, further satellite points located on the surface of the sphere are checked. This makes sure that (parts of) spheres can be located even when their centre is on another processor. This is especially important for parallel computations, when a sphere is about to move from one processor to another.

### Restrictions:



Only for immersed boundary solvers!

**Related commands:**

[locateModel](#)

## locateModel command

### Syntax:

Defined in couplingProperties dictionary.

```
locateModel model;
```

- model = name of the locateModel to be applied

### Examples:

```
locateModel engine;
```

Note: This examples list might not be complete - please look for other models (locateModel\_XY) in this documentation.

### Description:

The locateModel is the base class for models which search for the CFD cell and cellID corresponding to a position. In general it is used to find the cell a particle is located in.

**Restrictions:** none.

**Default:** none.

## **locateModel\_standardSearch command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
locateModel standard;
```

### **Examples:**

```
locateModel standard;
```

### **Description:**

The locateModel "standard" locates the CFD cell and cellID corresponding to a given position. A very straight-forward (robust!) locate algorithm is used.

**Restrictions:** none.

### **Related commands:**

[locateModel](#)

## locateModel\_turboEngineSearch command

### Syntax:

Defined in couplingProperties dictionary.

```
locateModel turboEngine;  
turboEngineProps  
{  
    treeSearch switch1;  
}
```

- *switch1* = switch to use tree search algorithm

### Examples:

```
locateModel turboEngine;  
turboEngineProps  
{  
    treeSearch false;  
}
```

### Description:

The locateModel "turboEngine" locates the CFD cell and cellID corresponding to a given position. The algorithm is improved compared to engine search to show better parallel performance.

The turboEngineSearch locate Model can be used with different settings to use different algorithms:

- faceDecomp false; treeSearch false; will execute some geometric (linear) search using the last known cellID (recommended)
- faceDecomp false; treeSearch true; will use a recursive tree structure to find the cell.

**Restrictions:** none.

**Related commands:**

[locateModel](#)

## meshMotionModel command

### Syntax:

Defined in couplingProperties dictionary.

```
meshMotionModel model;
```

- model = name of the meshMotionModel to be applied

### Examples:

```
meshMotionModel noMeshMotion;
```

Note: This examples list might not be complete - please look for other models (meshMotionModel\_XY) in this documentation.

### Description:

The meshMotionModel is the base class for models which manipulate the CFD mesh according to the DEM mesh motion.

**Restrictions:** none.

**Default:** none.

## **meshMotionModel\_noMeshMotion command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
meshMotionModel noMeshMotion;
```

### **Examples:**

```
meshMotionModel noMeshMotion;
```

### **Description:**

The noMeshMotion-model is a dummy meshMotion model.

**Restrictions:** None.

### **Related commands:**

[meshMotionModel](#)

## **momCoupleModel\_explicitCouple command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
momCoupleModels
(
    explicitCouple
);
explicitCoupleProps
{
    fLimit vector;
}
```

- *vector* = limiter vector for explicit force term (default (1e10,1e10,1e10) )

### **Examples:**

```
momCoupleModels
(
    explicitCouple
);
explicitCoupleProps
{
    fLimit (1e3 1e2 1e4);
}
```

### **Description:**

The explicitCouple-model is a momCoupleModel model providing an explicit momentum source term for the CFD solver.

### **Restrictions:**

Only for solvers that include explicit momentum exchange.

### **Related commands:**

[momCoupleModel](#)

## **momCoupleModel command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
momCoupleModels
(
    model
);
```

- model = name of the momCoupleModel to be applied

### **Examples:**

```
momCoupleModels
(
    implicitCouple
);
```

Note: This examples list might not be complete - please look for other models (momCoupleModel\_XY) in this documentation.

### **Description:**

The momCoupleModel is the base class for momentum exchange between DEM and CFD simulation.

**Restrictions:** none.

**Default:** none.



## **momCoupleModel\_implicitCouple command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
momCoupleModels
(
    implicitCouple
);
implicitCoupleProps
{
    velFieldName "U";
    granVelFieldName "Us";
    voidfractionFieldName "voidfraction";
}
```

- *U* = name of the finite volume fluid velocity field
- *Us* = name of the finite volume granular velocity field
- *voidfraction* = name of the finite volume voidfraction field

### **Examples:**

```
momCoupleModels
(
    implicitCouple
);
implicitCoupleProps
{
    velFieldName "U";
    granVelFieldName "Us";
    voidfractionFieldName "voidfraction";
}
```

### **Description:**

The implicitCouple-model is a momCoupleModel model providing an implicit momentum source term for the CFD solver.

### **Restrictions:**

Only for solvers that include implicit momentum exchange.

### **Related commands:**

[momCoupleModel](#)

## **momCoupleModel\_noCouple command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
momCoupleModels
(
    off
);
```

### **Examples:**

```
momCoupleModels
(
    off
);
```

### **Description:**

The noCouple-model is a dummy momCoupleModel model providing an no momentum source term for the CFD solver.

### **Restrictions:**

Only for solvers that include no momentum exchange, e.g. immersed boundary.

### **Related commands:**

[momCoupleModel](#)

## **regionModel\_allRegion command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
regionModel allRegion;
```

### **Examples:**

```
regionModel allRegion;
```

### **Description:**

The allRegion-model is a region model including the whole CFD region for the coupling.

**Restrictions:** None.

### **Related commands:**

[regionModel](#)

## regionModel command

### Syntax:

Defined in couplingProperties dictionary.

```
regionModel model;
```

- model = name of the regionModel to be applied

### Examples:

```
regionModel allRegion;
```

Note: This examples list might not be complete - please look for other models (regionModel\_XY) in this documentation.

### Description:

The regionModel is the base class for region models to select a certain region for coupled simulation.

**Restrictions:** none.

**Default:** none.

## voidfractionModel\_bigParticleVoidFraction command

### Syntax:

Defined in couplingProperties dictionary.

```
voidfractionModel bigParticle;  
bigParticleProps  
{  
    maxCellsPerParticle number1;  
    alphaMin number2;  
    scaleUpVol number3;  
}
```

- *number1* = max number of cells covered by a particle (search will fail when more than *number1* cells are covered by the particle)
- *number2* = minimum limit for voidfraction
- *number3* = diameter of the particle's representation is artificially increased according to *number3* \*  $V_{\text{particle}}$ , volume remains unaltered!

### Examples:

```
voidfractionModel bigParticle;  
bigParticleProps  
{  
    maxCellsPerParticle 1000;  
    alphaMin 0.10;  
    scaleUpVol 5.0;  
}
```

### Description:

The bigParticle voidFraction model is supposed to be used when a particle (or its representation) is bigger than a CFD cell. The voidfraction field is set in those cell whose centres are inside the particle which results in a stairstep representation of the bodies within the mesh (i.e. voidfraction is either 1 (fluid) or zero (solid)). For achieving accurate results, approx. 8 cells per particle diameter are necessary.

The region of influence of a particle can be increased artificially by "scaleUpVol", which blows up the particles, but keeps their volume (for voidfraction calculation) constant.

Code of this sub-model contributed by Alice Hager, JKU.

**Restrictions:** none.

**Related commands:**

[voidfractionModel](#)

## voidfractionModel\_centreVoidFraction command

### Syntax:

Defined in couplingProperties dictionary.

```
voidfractionModel centre;  
centreProps  
{  
    alphaMin value;  
}
```

- *value* = minimum limit for voidfraction

### Examples:

```
voidfractionModel centre;  
centreProps  
{  
    alphaMin 0.1;  
}
```

### Description:

The centre voidFraction model calculates the voidfraction in a CFD cell accounting for the volume of the particles whose centres are inside the cell.

**Restrictions:** none.

**Related commands:**

[voidfractionModel](#)

## voidfractionModel\_dividedVoidFraction command

### Syntax:

Defined in couplingProperties dictionary.

```
voidfractionModel divided;
dividedProps
{
    alphaMin number1;
    scaleUpVol number2;
    interpolation;
}
```

- *number1* = minimum limit for voidfraction
- *number2* = diameter of the particle's representation is artificially increased according to *number2* \*  $V_{\text{particle}}$ , volume remains unaltered!
- *interpolation* = flag to interpolate voidfraction to particle positions (normally off)

### Examples:

```
voidfractionModel divided;
dividedProps
{
    alphaMin 0.2;
    scaleUpVol 1.0;
}
```

### Description:

The divided voidFraction model is supposed to be used when a particle (or it's representation) is in the size range of a CFD cell. Satellite points are used to divide the particle's volume to the touched cells.

The region of influence of a particle can be increased artificially by "scaleUpVol", which blows up the particles, but keeps their volume (for voidfraction calculation) constant.

**Restrictions:** none.

### Related commands:

[voidfractionModel](#)

## voidfractionModel\_GaussVoidFraction command

### Syntax:

Defined in couplingProperties dictionary.

```
voidfractionModel Gauss;
GaussProps
{
    maxCellsPerParticle number1;
    alphaMin number2;
    scaleUpVol number3;
}
```

- *number1* = max number of cells covered by a particle (search will fail when more than *number1* cells are covered by the particle)
- *number2* = minimum limit for voidfraction
- *number3* = diameter of the particle's representation is artificially increased according to *number3* \*  $V_{\text{particle}}$ , volume remains unaltered!

### Examples:

```
voidfractionModel Gauss;
GaussProps
{
    maxCellsPerParticle 1000;
    alphaMin 0.10;
    scaleUpVol 5.0;
}
```

### Description:

The Gauss voidFraction model is supposed to be used when a particle (or its representation) is bigger than a CFD cell. The voidfraction field is set in those cell whose centres are inside the particle. The volume is here distributed according to a Gaussian distribution.

The region of influence of a particle can be increased artificially by "scaleUpVol", which blows up the particles, but keeps their volume (for voidfraction calculation) constant.

**Restrictions:** none.

**Related commands:**

[voidfractionModel](#) , [bigParticle](#)



## **voidfractionModel command**

### **Syntax:**

Defined in couplingProperties dictionary.

```
voidfractionModel model;
```

- model = name of the voidfractionModel to be applied

### **Examples:**

```
voidfractionModel centre;
```

Note: This examples list might not be complete - please look for other models (voidfractionModel\_XY) in this documentation.

### **Description:**

The voidfractionModel is the base class for models to represent the DEM particle's volume in the CFD domain via a voidfraction field.

**Restrictions:** none.

**Default:** none.

## voidfractionModel\_IBVoidFraction command

### Syntax:

Defined in couplingProperties dictionary.

```
voidfractionModel IB;
IBProps
{
    maxCellsPerParticle number1;
    alphaMin number2;
    scaleUpVol number3;
}
```

- *number1* = max number of cells covered by a particle (search will fail when more than *number1* cells are covered by the particle)
- *number2* = minimum limit for voidfraction
- *number3* = diameter of the particle's representation is artificially increased according to *number3* \*  $V_{\text{particle}}$ , volume remains unaltered!

### Examples:

```
voidfractionModel IB;
IBProps
{
    maxCellsPerParticle 1000;
    alphaMin 0.10;
    scaleUpVol 5.0;
}
```

### Description:

The IB voidFraction model is supposed to be used when a particle (or its representation) is bigger than a CFD cell. The voidfraction field is set in those cells whose centres are inside the particle. The model is specially designed for cfemSolverIB and creates a smooth transition of the voidfraction at the particle surface. Cells which are only partially covered by solid are marked by void fraction values between 0 and 1 respectively.

The region of influence of a particle can be increased artificially by "scaleUpVol", which blows up the particles, but keeps their volume (for voidfraction calculation) constant.

Code of this sub-model contributed by Alice Hager, JKU.

**Restrictions:** none.

**Related commands:**

[voidfractionModel](#)