User Guide for OpenFOAM-10

DTLreactingFoam: An efficient CFD tool for laminar reacting flow simulations using detailed chemistry and transport with time-correlated thermophysical properties

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Contents

1	General descriptions			3
	1.1	thermo	physicalModels library	3
	1.2	DTLrea	actingFoam solver	5
	1.3	preproc	essing utilities	5
		1.3.1	DTMchemkinToFoam	5
		1.3.2	FTMchemkinToFoam	5
2 Running simulations with DTLreactingFoam			mulations with DTLreactingFoam	5
	2.1	Simulations for general reacting flows using DTM		5
		2.1.1	Prepare input files	5
		2.1.2	Setup and run simulation	9
2.2 Simulations for general reacting flows using		Simulat	ions for general reacting flows using FTM	12
		2.2.1	Prepare input files	12
		2.2.2	Setup and run simulation	16

1. General descriptions

The framework is fully native OpenFOAM(OF) and comprises newly developed libraries, solvers, and utilities as follows:

- libraries:
 - thermophysicalModels
- solvers:
 - DTLreactingFoam
- utilities:
 - DTMchemkinToFoam
 - FTMchemkinToFoam

1.1. thermophysical Models library

The class diagram of a target library plays a crucial role for guiding code structure and modularity in OF code development (i.e., OOP). It outlines essential features such as inheritance hierarchies and class interfaces. Figure 1 depicts the class diagram of the updated thermophysicalModels library in the OF-10, highlighting key classes associated with reacting flow modeling. All classes can be organized in four conceptual blocks; the ThermoType block and MixtureType block representing various thermophysical models and mixture models; the ChemistryType Block, encompassing chemistry models; and the BasicType Block, which includes base classes for system types (e.g., rho-based or psi-based). While there are structural differences between the thermophysicalModels libraries in OF-10 and OF-6 (as presented in [1]), the core implementation philosophy of thermophysical property calculation remains consistent.

In the updated thermophysicalModels library, as shown in Figure 1, green boxes marked with numbers denote sample classes that typify groups of models, following the approach presented by [1]. Yellow boxes represent newly developed classes (i.e., DTM, FTM)

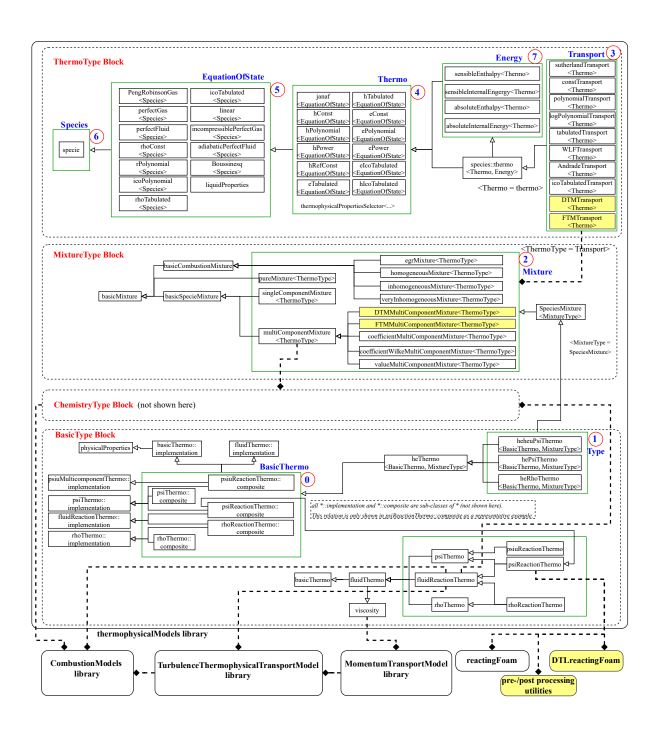


Figure 1: The class diagram of the updated thermophysicalModels library in the DTLreactingFoam-10 framework. The boxes with marked numbers denote sample classes. The arrow-line denotes the inheritance relationship in which the direction of arrow is from a subclass to its base class. A dashed line denotes a class-class or class-solver interface in which one class is used in another or in a solver.

added to the original thermophysicalModels library. The DTM and FTM are implemented following the methodology proposed by Nguyen et al. [1] to address limitations in OF's capability to support complex mixing rules in custom thermophysical models. The coTHERM method is subsequently integrated into these models via the hePsiThermo class.

1.2. DTLreactingFoam solver

The DTLreactingFoam solver is developed based on the realFluidReactingFoam solver [1], which was originally implemented on OF-6 (i.e., extending reactingFoam), for both transient and steady-state laminar reacting flow simulations incorporating DTM/FTM with the coTHERM method to reduce computational time while preserving accuracy.

1.3. preprocessing utilities

1.3.1. DTMchemkinToFoam

Based on the original OF chemkinToFoam utility, a new DTMchemkinToFoam pre-processing utility is created to accommodate the newly developed DTM. It automates the conversion of input files from CHEMKIN to OF format.

1.3.2. FTMchemkinToFoam

Similarly, dedicated FTMchemkinToFoam pre-processing utility is created to convert input files from CHEMKIN to OF format for the newly developed FTM.

2. Running simulations with DTLreactingFoam

2.1. Simulations for general reacting flows using DTM

2.1.1. Prepare input files

Before running simulation using the DTM, an input file, say thermo.DTM, is needed. It involves essential parameters for transport property calculations and must be in the following format:

```
02
{
...
```

```
transport
    {
        // input parameters for DTM
        linearity
                          1;
        epsilonOverKb
                          107.4;
        sigma
                          3.458;
        dipoleMoment
                          0;
                          1.6;
        alpha
        Zrot
                          3.8;
    }
}
```

The thermo.DTM input file can be generated using DTMchemkinToFoam utility with the following command:

```
DTMchemkinToFoam <input1> <input2> <input3> <input4> <output1> <output2> where:
```

- input1: chemical mechanism file in CHEMKIN format, e.g., chem.inp
- input2: data file for thermodynamics in CHEMKIN format, e.g., therm.dat
- input3: data file for DTM transport in CHEMKIN format, e.g., trans.dat
- input4: data file for Sutherland transport, e.g., transportSutherland
- output1: chemical mechanism file in OF format, e.g., reactions
- output2: data file for thermodynamic and transport properties using DTM, e.g., thermo.DTM

The three first input files in CHEMKIN format can be taken directly from the desired chemical mechanism available in the literature. The input4 file must be in OF format as following:

```
".*"
{
```

```
transport
{
         As 1.512e-06;
         Ts 120.;
}
```

At DTLreactingFoam-10/tutorials/Mech/DTMchemkinToFoam/example/ directory, an example is provided to demonstrate the use of the DTMchemkinToFoam utility. It is important to note that using DTMchemkinToFoam requires only four input files placed in the case directory. In other words, 0, constant, and system directories are not needed in this step. To generate the thermo.DTM file, first go to the case directory by:

```
cd ~/OpenFOAM/yourDirectory/DTLreactingFoam-10/tutorials/Mech/
DTMchemkinToFoam/example/
```

Then execute the following command in the terminal:

```
// Orders of files are important and need to be specified correctly
DTMchemkinToFoam chem.inp therm.dat trans.dat transportSutherland
  reactions thermo.DTM
```

After executing that command, two output files named reactions and thermo.DTM are generated as:

```
// This is inside the 'reactions' file:
reactions
{
     un-named-reaction-0
     {
           type
                                   reversibleArrhenius;
           reaction
                                   "CH4_{\sqcup} + _{\sqcup} 202_{\sqcup} = _{\sqcup} C02_{\sqcup} + _{\sqcup} 2H20";
           Α
                                   1.2e + 11;
           beta
                                   -1;
           Τa
                                   0;
     }
}
```

```
Tlow 200;
Thigh 3500;
```

```
// This is inside the 'thermo.DTM' file:
        5 ( CH4 H2O O2 CO2 N2 );
species
N2
{
   specie
   {
       molWeight 28.0134;
   }
    thermodynamics
    {
       Tlow
                       200;
                       5000;
       Thigh
       Tcommon
                      1000;
       highCpCoeffs ( 2.92664 0.0014879768 -5.68476e-07 1.0097038e-10
           -6.753351e-15 -922.7977 5.980528 );
        lowCpCoeffs (3.298677 0.0014082404 -3.963222e-06 5.641515e-09
            -2.444854e-12 -1020.8999 3.950372 );
   }
   transport
    {
                       1.512e-06;
        Αs
       Тs
                       120;
       linearity
                       1;
        epsilonOverKb
                     97.53;
        sigma
                       3.621;
        dipoleMoment
                       0;
       alpha
                       1.76;
       Zrot
                       4;
   }
    elements
    {
```

```
N 2;
}

CO2

...

...
```

These two files now are ready for use.

Examples with script Allrun to automate converting files for several detailed chemical mechanisms are also available in DTLreactingFoam-10/tutorials/Mech/DTMchemkinToFoam/directory.

2.1.2. Setup and run simulation

A test case of 2-D counterflow diffusion flame of $\mathrm{CH_4/air}$, which is available in tutorials directory of the original OF, is selected to demonstrate the setting and running simulation using the DTLreactingFoam solver with the DTM and coTHERM method in this manual (see the DTLreactingFoam-10/tutorials/counterFlowFlame2D_example directory). To create this test case, first, go to the tutorials in your directory. Then copy the original counterFlowFlame2D case from OpenFOAM into your directory as:

```
cd ~/OpenFOAM/yourDirectory/DTLreactingFoam-10/tutorials/
cp -rf ~/OpenFOAM/OpenFOAM-10/tutorials/combustion/reactingFoam/laminar/
    counterFlowFlame2D/ .
```

Then go to the counterFlowFlame2D directory:

```
cd ~/OpenFOAM/yourDirectory/DTLreactingFoam - 10/tutorials/
counterFlowFlame2D/
```

Then, copy two files (e.g., reactions and thermo.DTM) which are generated using the DTMchemkinToFoam utility, as described in Section 2.1.1, into the constant directory.

In this directory, the physicalProperties dictionary file must be modified to be compatible with the DTLreactingFoam solver. Particularly, in the physicalProperties dictionary file, add new essential keywords and modify the thermoType entry as follows:

```
usingDetailedTransportModel
                             true;
                                     //newly added
usingCoTHERM
                                    //newly added
                              true:
majorSpeciesForCoTHERM (CH4 02 CO2 H20); //newly added
epsilonT
                  0.2;
                         // [K], newly added
epsilonP
                  100; // [Pa], newly added
epsilonS
                  0.001; // [-], in mass fraction, newly added
thermoType
{
    type
                    hePsiThermo;
                    DTMMultiComponentMixture; //for DTM
    mixture
                    DTMTransport;
                                              //for DTM
    transport
    thermo
                    janaf;
    energy
                    sensibleEnthalpy;
    equationOfState perfectGas;
    specie
                    specie;
}
defaultSpecie N2; //this can be changed
//#include "thermo.compressibleGas" //original OF
#include "thermo.DTM" //for DTM
```

where meaning and options of newly added keywords are explained as below:

- usingDetailedTransportModel: It must be explicitly set to be true when using DTM. Otherwise the thermophysical properties will be not updated in each time step during simulation. The default value of this keyword is false, appropriate for using original STM since our code does not eliminate any functionality of original OF.
- usingCoTHERM: It must be explicitly set to be true if you want to apply the coTHERM method with DTM for your simulation to reduce the computational time. The default

value of this keyword is false, appropriate for using the DTM alone.

- majorSpeciesForCoTHERM: This is a list of selected species in the chemical mechanism considered as representative major species for the mixture when using the coTHERM method. Note that species selection is case dependent. Typically, it includes dominant species in both unburnt and burnt regions, for example of CH₄/air flame: CH₄, O₂, N₂, CO₂, H₂O, OH, HO₂, CH₂O [2].
- epsilonT: This is threshold value (in [K]) to control residual of temperature, corresponding to ε_T in the coTHERM method. For a general reacting flow, we recommend this value is set to be 0.2 K for computational efficiency without sacrificing accuracy. If this value is not specified, the program will automatically use 0.2 K as a default value for ε_T .
- epsilonP: This is threshold value (in [Pa]) to control residual of pressure, corresponding to ε_P in the coTHERM method. For a general reacting flow, we recommend this value is set to be 100 Pa for computational efficiency without sacrificing accuracy. If this value is not specified, the program will automatically use 100 Pa as a default value for ε_P .
- epsilonS: This is threshold value (dimensionless) to control residual of species mass fraction, corresponding to ε_S in the coTHERM method. For a general reacting flow, we recommend this value is set to be 0.001 for computational efficiency without sacrificing accuracy. If this value is not specified, the program will automatically use 0.001 as a default value for ε_S .

All other settings are identical to those used in the reactingFoam solver.

It is important to note that the default keyword of the divSchemes entry in the fvSchemes dictionary file, placed in the system directory, needs to be changed from none to a specific type, for example:

divSchemes

```
{
    // default none; // original OF
    default Gauss linear; // change from 'none' to 'Gauss linear'
    ...
}
...
```

Everything now is ready for use. For running simulation, first run blockMesh to generate geometry, then type the solver name in the terminal (in the same way with reactingFoam) as:

```
blockMesh
DTLreactingFoam
```

2.2. Simulations for general reacting flows using FTM

Simulations for general reacting flows using FTM are the same as using DTM except for input files and keyword setup for models combined with the FTM.

2.2.1. Prepare input files

Before running simulation using the FTM, an input file, say thermo.FTM, is needed. It involves essential parameters for transport property calculations and must be in the following format:

```
(-27.6164 4.47195 -0.36524 0.0158195)
(-25.3927 3.75885 -0.275697 0.0120551)
...
);
```

The thermo.FTM input file can be generated using the FTMchemkinToFoam utility.

Unlike the DTMchemkinToFoam utility, the FTMchemkinToFoam requires a case directory that already works with the DTLreactingFoam solver using the DTM. In other words, the case directory involves 0, constant, and system directories, as the same as the standard OF case. Especially, in the constant directory, the thermo.DTM file must be provided as an input for fitting procedure.

The counterFlowFlame2D case in the Section 2.1.2 can be used to demonstrate the procedure of generating the thermo.FTM file. To do so, first go to that case as:

```
cd ~/OpenFOAM/yourDirectory/DTLreactingFoam-10/tutorials/
counterFlowFlame2D/
```

Then open the constant/physicalProperties file and modify as following:

```
// All setting of coTHERM are same as applied for DTM
usingDetailedTransportModel true; //must be true
usingCoTHERM
                            false; //must be false
usingPreProcessingFTM
                            true;
                                   //must be true, default is false
thermoType
{
   mixture
                   DTMMultiComponentMixture; //for DTM
                                             //for DTM
   //transport
                    DTMTransport;
   transport
                   preprocessingFTMTransport; //for preprocessing only
    . . .
defaultSpecie N2;
#include "thermo.DTM" //The thermo file for DTM must be included
```

where usingPreProcessingFTM keyword must be added, and must be explicitly set to be true. The default value of this keyword is false.

Additionally, the FittingDict dictionary file is needed for fitting procedure. It is placed in the constant directory and has the following format:

```
----*- C++ -*-
 _____
 \\ / F ield
                        | OpenFOAM: The Open Source CFD Toolbox
           O peration | Website: https://openfoam.org
          And
                        Version:
           M anipulation
FoamFile
{
   format
            ascii;
   class
              dictionary;
   location
             "constant";
   object
              FittingDict;
}
                     * * * * * * * * * * * * * * * * * * //
numberOfPoint 101; // [points] don't change this value
                3000; // [K] don't change this value
maxT
minT
                300; // [K] this may be changed
                101325; // [Pa] this is case dependent
pressure
```

where:

- numberOfPoint: The number of temperature sampling points used in the polynomial fitting process. The number of 101 points is found to be optimal value for this process. Higher values does not improve fitting accuracy but increase unnecessarily computational cost.
- maxT: The upper bound of the fitting temperature range $[T_{\min}, T_{\max}]$ over which trans-

port properties are sampled and fitted. This value should be chosen to cover the expected flame temperature range.

- minT: The lower bound of the fitting temperature range $[T_{\min}, T_{\max}]$ for sampling. This value should capture the range of cold reactants or ambient conditions.
- pressure: The reference pressure at which all transport properties are evaluated and fitted. This is actually the operating pressure condition of your problem.

Now everything is ready. Then execute the following command in the terminal to run the FTMchemkinToFoam:

```
cd ~/OpenFOAM/yourDirectory/DTLreactingFoam-10/tutorials/
    counterFlowFlame2D/
FTMchemkinToFoam
```

After execution, the thermo.FTM file is generated, and placed in the constant directory. It is ready for use in simulations with the FTM.

It is important to note that, after running the FTMchemkinToFoam, a new file named TransportCoeffDict is generated in the constant directory. It has the following format:

```
// in side 'TransportCoeffDict' file:
species
(
    CH4
    H20
     02
     C02
     N2
);

CH4
{
    fittingTranCoeff
    {
        muCoeffs (-22.2457 3.54364 -0.383872 0.0169568);
}
```

```
kappaCoeffs
                          (0.812318 - 4.54742 \ 0.975145 - 0.0536628);
         DijCoeffs
                        (
                           (-9.21034 -6.44736e-26 9.25847e-27 -4.40439e-28)
                           (-30.6998 \ 5.62052 \ -0.483426 \ 0.0197159)
                           (-25.3927 \ 3.75885 \ -0.275697 \ 0.0120551)
                           (-28.4471 \ 4.81747 \ -0.406845 \ 0.0174929)
                           (-25.0853 \ 3.63653 \ -0.260141 \ 0.0113939)
                        );
    }
};
H20
{
    fittingTranCoeff
}
```

This file is generated for only purpose of checking the fitting transport coefficients of the FTM in case it needs double check. Otherwise, this file can be ignored.

2.2.2. Setup and run simulation

To run simulation with the FTM, new essential keywords for coTHERM are required in the physicalProperties dictionary file, as explained in Section 2.1.2. The dictionary file need to be modified to be compatible with the FTM as follows:

```
thermoType
{
    type
                     hePsiThermo;
                     FTMMultiComponentMixture; //for FTM
    mixture
                     FTMTransport; //for FTM
    transport
    thermo
                     janaf;
    energy
                     sensibleEnthalpy;
    equationOfState perfectGas;
    specie
                     specie;
}
defaultSpecie N2;
#include "thermo.FTM" //for FTM
```

All other settings are identical to those used in the DTLreactingFoam solver with DTM model.

Everything now is ready for use. For running simulation, type the solver name in the terminal (in the same way with reactingFoam) as:

```
DTLreactingFoam
```

References

- D. N. Nguyen, K. S. Jung, J. W. Shim, C. S. Yoo, Real-fluid thermophysical Models library: An OpenFOAM-based library for reacting flow simulations at high pressure, Comput. Phys. Commun. 273 (2022) 108264.
- [2] S. Yang, R. Ranjan, V. Yang, S. Menon, W. Sun, Parallel on-the-fly adaptive kinetics in direct numerical simulation of turbulent premixed flame, Proc. Combust. Inst. 36 (2017) 2025–2032.