# Implementation Guide (for OpenFOAM-8 version):

Real-fluid thermophysical Models: An OpenFOAM-based library

for reacting flow simulations at high pressure

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To cite this article: D. N. Nguyen, K. S. Jung, J. W. Shim, C. S. Yoo, Real-fluid thermophysical Models: An OpenFOAM-based library for reacting flow simulations at high pressure, Computer Physics Communications 273 (2022) 108264.

Note: This document presents the implementation for only a set of widely used real-fluid models i.e., the Soave-Redlich-Kwong [1, 2] equation of state (EoS), Chung's model [3] for dynamic viscosity and thermal conductivity, mixture averaged model for mass diffusivity using Takahashi's correction [4] for binary diffusion coefficients at high pressure. For other models such as Peng-Robinson EoS [5] or the Standard Kinetic Theory model [6] for transport properties, readers are referred to 2-GuideForExtension.pdf file.

# Contents

1	Ар	rocedure for real-fluid models implementation	3
2	$\operatorname{Cre}$	ate runtime selectable packages of real-fluid models	7
	2.1	Making source files	7
	2.2	Making runtime selectable thermophysical model package	10
		2.2.1 Making interface inside thermophysical Models library	11
		2.2.2 Making interface outside of the thermophysical Models library	27
		2.2.3 Test new created runtime thermo-packages	36
3	Det	ailed implementation of real-fluid models	39
4	Det	ailed modification of existing classes	39
	4.1	Modification of <i>Transport</i> classes	39
	4.2	Modification of <i>Mixture</i> classes	40
	4.3	Modification of BasicThermo classes	44
	4.4	Modification of $Type$ classes	45
5	Usi	ng the new library	58

#### 1. A procedure for real-fluid models implementation

Figure 1 illustrates the class diagram of thermophysicalModels library in OpenFOAM (OF) 8.0 with sufficient classes associated with reacting flow simulations. The class diagram provides important features such as inheritance and the interface between classes which are necessary for code development of the thermophysicalModels library. In this diagram, the boxes with marked numbers denote the sample classes which can be one of the classes in side the box due to runtime selection mechanism. ThermoType Block and MixtureType Block contain all classes representing thermophysical (THERMPHYS) models and mixture models. ChemistryType Block consists of chemistry models while BasicType Block involves base classes for types of system (e.g., rho-based or psi-based).

Since the interface between classes in thermophysical Models is relatively complicated, we propose a simple procedure to implement real-fluid models as follows to minimize the debugging efforts during the implementation. The following proposed procedure is based on OF-8, it can also be referred to implement real-fluid models in other version of OF such as OF-7, OF-9. The procedure has three main steps:

- Step 1: Create runtime selectable packages including real-fluid models;
- Step 2: Implement real-fluid THERMPHYS models;
- Step 3: Modify related classes.

In the step 1, new classes representing to real-fluid models are created by copying from existing classes in the thermophysicalModels library without implementing any equations of real-fluid models into the code. In addition, some existing macros files need to be modified and some new macros files are created to form runtime selectable packages for real-fluid models. The detail instruction about the step 1 is presented in Sec. 2. In the step 2, the source code of new class are modified to deliver equations of new models. The detail instruction about the step 2 is shown in Sec. 3. In step 3, source code of several classes are modified to be compatible with new added classes due to the use of templates, polymorphism, and inheritance in OF. The detail instruction about the step 3 is presented in Sec. 4.

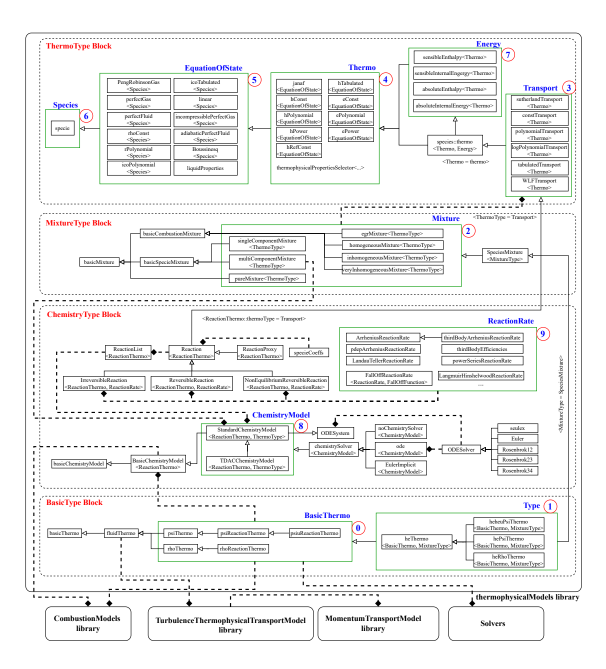


Figure 1: The class diagram of the *thermophysicalModels* library in OF-8. The boxes with marked numbers denote sample classes. The arrow-line denotes the inheritance relationship in which the direction of arrow is from a subclass to its base class. A dashed line denotes a class-class or class-solver interface in which one class is used in another or in a solver.

Figure. 2 shows the class diagram of the new thermophysical Models library of OF-8 including real fluid models. In this library, several new classes are created (in yellow boxes) such as rfSpecie, soaveRedlichKwong, rfJanafThermo, and chungTakaTransport representing for species, soave-Redlich-Kwong (SRK) equation of state (EoS), real-fluid JANAF-based, and Chung's models, respectively. Note that mixture averaged mass diffusivity model with Takahashi correlation for binary diffusion coefficients is included in chungTakaTransport class. The SRKchungTakaMixture is a new classes representing for mixture model which is implemented using a new algorithm proposed in our paper. The SRKchungTakaStandardChemistryModel is a new chemistry model class created by copying the original StandardChemistryModel class in OpenFOAM containing the interface with real fluid models. Furthermore, several existing classes in original thermophysicalModels library have also been modified (in gray boxes) to be compatible with new added classes.

New classes have been created:

```
// In thermophysicalModels/specie directory:
- specie/rfSpecie/rfSpecie
- specie/equationOfState/soaveRedlichKwong/soaveRedlichKwong
- specie/themo/rfJanaf/rfJanafThermo
- specie/transport/chungTaka/chungTakaTransport
// In thermophysicalModels/reactionThermo directory:
- reactionThermo/mixtures/SRKchungTakaMixture
// In thermophysicalModels/chemistryModel directory:
- chemistryModel/chemistryModel/SRKchungTakaStandardChemistryModel
```

Classes have been modified:

```
// In thermophysicalModels/reactionThermo/mixtures directory:
- reactionThermo/mixtures/multiComponentMixture/multiComponentMixture
- reactionThermo/mixtures/singleComponentMixture/singleComponentMixture
- reactionThermo/mixtures/egrMixture/egrMixture
- reactionThermo/mixtures/homogeneousMixture/homogeneousMixture
- reactionThermo/mixtures/inhomogeneousMixture/inhomogeneousMixture
- reactionThermo/mixtures/veryInhomogeneousMixture/
```

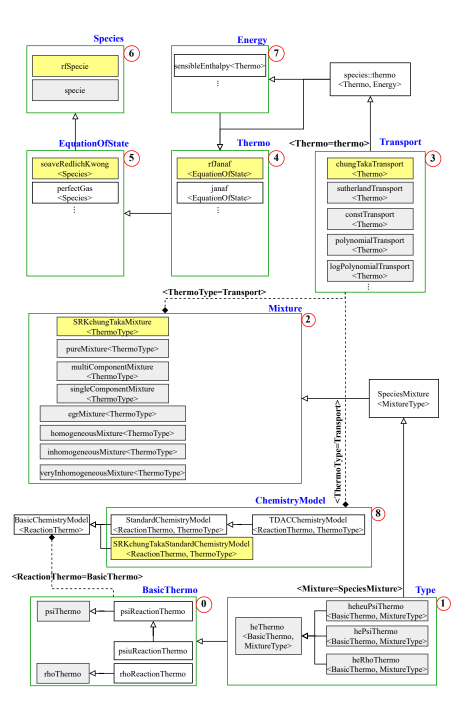


Figure 2: The class diagram of the real-fluid thermophysical Models library in OF-8. Yellow boxes are classes need to be created representing to real-fluid models. Gray boxes are existing classes in the original thermophysical Model library need to be modified. The arrow-line denotes the inheritance 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

```
veryInhomogeneousMixture
- reactionThermo/psiuReactionThermo/heheuPsiThermo
// In thermophysicalModels/basic directory:
- basic/mixtures/pureMixture
- basic/heThermo/heThermo
- basic/rhoThermo/rhoThermo
- basic/rhoThermo/heRhoThermo
- basic/psiThermo/psiThermo
- basic/psiThermo/hePsiThermo
```

# 2. Create runtime selectable packages of real-fluid models

## 2.1. Making source files

Prepare a directory on your system, e.g., your Directory.

```
mkdir ~/OpenFOAM/yourDirectory
cd ~/OpenFOAM/yourDirectory/
```

Create /realFluidThermophysicalModels-8/src/ directory.

```
mkdir -p realFluidThermophysicalModels-8/src
```

Set an environment variable prescribing the path of the *src* directory.

```
echo "exportuLIB_rfFoam8_SRC=~/OpenFOAM/yourDirectory/
realFluidThermophysicalModels-8/src/" >> ~/.bashrc
source ~/.bashrc
```

Go to thermophysical Models directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
```

Copy ODE, thermphysical Models, Momentum Transport Models, Thermophysical Transport Models, radiation Models, combustion Models, fvOptions directories from original OpenFOAM into real Fluid Thermophysical Models - 8/src/ in your user directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
cp -rf -p $WM_PROJECT_DIR/src/ODE .
cp -rf -p $WM_PROJECT_DIR/src/thermophysicalModels .
```

```
cp -rf -p $WM_PROJECT_DIR/src/MomentumTransportModels .
cp -rf -p $WM_PROJECT_DIR/src/ThermophysicalTransportModels .
cp -rf -p $WM_PROJECT_DIR/src/radiationModels .
cp -rf -p $WM_PROJECT_DIR/src/combustionModels .
cp -rf -p $WM_PROJECT_DIR/src/fvOptions .
```

Then, remove unnecessary parts inside the *thermophysicalModels* and the *Momentum-TransportModels* such that there the following directories left after removal.

```
// In side thermophysicalModels directory
basic
chemistryModel
reactionModel
solidSpecie
solidThermo
specie
thermophysicalProperties
```

```
// In side MomentumTransportModels directory
compressible
momentumTransportModels
```

Then, go to /yourDirectory/.../src/thermophysicalModels/specie directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
thermophysicalModels/specie
```

Create rfSpecie class by copying from specie class.

```
cp -rf specie rfSpecie
cd rfSpecie
mv specie.H rfSpecie.H
mv specie.C rfSpecie.C
mv specieI.H rfSpecieI.H
```

Open these files and replace *specie* by *rfSpecie*. To make sure you do not miss any thing you should use a command to automatically find and replace a string, for instance use the following command if you are using vim text editor.

```
vi specie.H
:%s/specie/rfSpecie/g
```

Note that do not change *specie* as the name of a dictionary in a constructor of *rfSpecie* class as follows:

Do the same to create soaveRedlichKwong, rfJanafThermo, and chungTakaTransport classes from perfectGas, janafThermo, and sutherlandTransport classes inside equationOfS-tate, thermo, and transport directories, respectively. For instance:

```
cd equationOfState
cp -rf perfectGas soaveRedlichKwong
cd soaveRedlichKwong
mv perfectGas.H soaveRedlichKwong.H
mv perfectGas.C soaveRedlichKwong.C
mv perfectGasI.H soaveRedlichKwongI.H
vi soaveRedlichKwong.H
:%s/perfectGas/soaveRedlichKwong/g
...
```

Note that change *specie* into *rfSpecie* inside the *rfJanafThermo* and *chungTransport* classes since *rfSpecie* class is their base class. Do not confuse between *specie* (class's name) and *Specie* (name of type in templates), and the names at runtime of real-fluid models should be specified correctly in the \*.H files as follows:

```
/* In rfSpecie.H file: */
//- Runtime type information
ClassName("rfSpecie");
```

```
/* In soaveRedlichKwong.H file: */
//- Return the instantiated type name
static word typeName()
{
    return "soaveRedlichKwong<" + word(Specie::typeName_()) + '>';
}

/* In rfJanafThermo.H file: */
static word typeName()
{
    return "rfJanaf<" + EquationOfState::typeName() + '>';
}

/* In chungTransport.H file: */
static word typeName()
{
    return "chungTaka<" + Thermo::typeName() + '>';
}
```

# 2.2. Making runtime selectable thermophysical model package

In the OF-8, the THERMPHYS models are used in a reacting flow simulation by using runtime selection mechanism such that the *thermoType* dictionary is specified by user at runtime in the *constant/thermophysicalProperties* file as:

```
thermoType
{
                                            //(1) type of system
   type
                    hePsiThermo;
                    multiComponentMixture; //(2) mixture model
   mixture
                                            //(3) transport model
   transport
                    sutherland;
                    janaf;
                                            //(4) thermodynamic model
   thermo
                    sensibleEnthalpy;
                                            //(7) energy type model
   energy
                                            //(5) equation of state model
   equationOfState perfectGas;
                                            //(6) species model
   specie
                    specie;
```

A set of THERMPHYS models like that can be referred as a runtime thermo-package. Our goal is creating four thermo-packages including real-fluid THERMPHYS models in which the type of system can be either hePsiThermo (psi-based) or heRhoThermo (rho-based) and the energy type can be either sensibleEnthalpy or sensibleInternalEnergy as following:

```
thermoType
{
    type
                    hePsiThermo;
                                          //or "heRhoThermo"
                     SRKchungTakaMixture;
    mixture
    transport
                     chung Taka;
    thermo
                     rfJanaf;
                     sensibleEnthalpy; //or "sensibleInternalEnergy"
    energy
    equationOfState soaveRedlichKwong;
    specie
                     rfSpecie;
```

The structure of thermophysical Models library in the OF is relatively complicated since it is built based on a runtime selection mechanism using dynamic polymorphism, class templates, and macros. This instruction covers only models related to reacting flow simulations.

#### 2.2.1. Making interface inside thermophysical Models library

Copy the following directories from original OpenFOAM into thermophysical Models in your directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    thermophysicalModels/
cp -rf -p $WM_PROJECT_DIR/src/thermophysicalModels/basic .
cp -rf -p $WM_PROJECT_DIR/src/thermophysicalModels/reactionThermo .
cp -rf -p $WM_PROJECT_DIR/src/thermophysicalModels/chemistryModel .
cp -rf -p $WM_PROJECT_DIR/src/thermophysicalModels/
    thermophysicalProperties .
```

Then go to thermophyisicalModels/specie/include directory and make a new macros file for set of SRK-chung-Taka real fluid models named forSRKchungTakaGases.H by copying

from for Gases. H file.

```
cd ~/OpenFOAM/yourDirectory/src/thermophysicalModels/specie/include cp -rf forGases.H forSRKchungTakaGases.H
```

Then modify these following files in order.

```
// ======== The list files ==================================
// In thermophysicalModels/specie directory
1. thermophysicalModels/specie/Make/files
2. thermophysicalModels/specie/include/forSRKchungTakaGases.H //new file
   created
//--> compile specie to make specie.so
// In thermophysicalModels/thermophysicalProperties directory

    thermophysicalModels/basic/Make/files (+ options)

//--> compile basic to make thermophysicalProperties.so
// In thermophysical Models / basic directory

    thermophysical Models/basic/Make/files (+ options)

//--> compile basic to make fluidThermophysicalModels.so
// In thermophysical Models/reaction Thermo directory

    thermophysicalModels/reactionThermo/Make/files (+ options)

2. thermophysical Models/reaction Thermo/psiReaction Thermo/
   \verb"psiReactionThermos.C"
3. thermophysical Models/reaction Thermo/rhoReaction Thermo/
   rhoReactionThermos.C
4. thermophysical Models/reaction Thermo/chemistry Readers/chemistry Reader/
   makeChemistryReaders.C
//--> compile reactionThermo to make reactionThermophysicalModels.so
// In thermophysical Models / chemistry Model directory
1. thermophysical Models / chemistry Model / reactions / makeReactions . C
2. thermophysicalModels/chemistryModel/reactions/
   {\tt makeLangmuirHinshelwoodReactions.C}
3. thermophysicalModels/chemistryModel/chemistrySolver/
```

a. Modification in /yourDirectory/src/thermophysicalModels/specie directory:

In *Make/files* file, add *rfSpecie.C* into the list of source files and change the position to save the binary file, *specie.so*, at user directory.

In *include/forRealGases.H* file, replace the existing code as following:

```
#ifndef forSRKchungTakaGases_H

#define forSRKchungTakaGases_H

#include "rfSpecie.H"

#include "soaveRedlichKwong.H"

#include "rfJanafThermo.H"

#include "sensibleEnthalpy.H"

#include "sensibleInternalEnergy.H"

#include "chungTakaTransport.H"

#include "thermo.H"
```

```
#include "forThermo.H"
#define forSRKchungTakaGasEqns(Mu, He, Cp, Macro, Args...)
   forThermo (Mu, He, Cp, soaveRedlichKwong, rfSpecie, Macro, Args)
#define forSRKchungTakaGasEnergiesAndThermos(Mu, Macro, Args...)
   forSRKchungTakaGasEqns(Mu, sensibleEnthalpy, rfJanafThermo, Macro,
      Args); \
   forSRKchungTakaGasEqns(Mu, sensibleInternalEnergy, rfJanafThermo,
      Macro, Args)
#define forSRKchungTakaGasTransports(Macro, Args...)
                                                                  \
   forSRKchungTakaGasEnergiesAndThermos(chungTakaTransport, Macro, Args)
#define forSRKchungTakaGases(Macro, Args...)
                                                                  \
   forSRKchungTakaGasTransports(Macro, Args)
#endif
```

Compile to make specie.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    thermophysicalModels/specie
wclean
wmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

b. Modification in /your Directory/.../src/thermophysical Models/thermophysical Properties directory:

In Make/files file, change the position to save the binary file, thermophysicalProperties.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libthermophysicalProperties
```

In *Make/options* file, change the path to include the header files of *specie.o* library as follows:

```
EXE_INC = \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude

LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
    -lspecie
```

Compile to make thermophysical Properties. so.

```
cd ~/OpenFOAM/yourDirectory/.../src/thermophysicalModels/
    thermophysicalProperties
wclean
wmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

c. Modification in /yourDirectory/.../src/thermophysicalModels/basic directory:

In Make/files file, change the position to save the binary file, fluidThermophysicalModels.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libfluidThermophysicalModels
```

In *Make/options* file, change the path to include the header files of *specie.o* and *thermo-physicalProperties.so* libraries as follows:

```
EXE_INC = \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/thermophysicalProperties/
    lnInclude \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude

LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
```

```
-lspecie \
-lthermophysicalProperties \
-lfiniteVolume \
-lmeshTools
```

Compile to make fluidThermophysicalModels.so.

```
cd ~/OpenFOAM/yourDirectory/.../src/thermophysicalModels/basic
wclean
wmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

d. Modification in your Directory/.../src/thermophysical Models/reaction Thermo directory:

Go to /yourDirectory/.../src/thermophysicalModels/reactionThermo/mixtures directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
thermophysicalModels/reactionThermo/mixtures
```

Create SRKchungTakaMixture class by copying from multiComponentMixture class.

```
cp -rf multiComponentMixture SRKchungTakaMixture
cd SRKchungTakaMixture
mv multiComponentMixture.H SRKchungTakaMixture.H
mv multiComponentMixture.C SRKchungTakaMixture.C
```

Open these files and replace multiComponentMixture by SRKchungTakaMixture. To make sure you do not miss any thing you should use a command to automatically find and replace a string, for instance use the following command if you are using vim text editor.

```
vi SRKchungTakaMixture.H
:%s/multiComponentMixture/SRKchungTakaMixture/g
```

In Make/files file, change the position to save the object file, reactionThermophysicalModels.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libreactionThermophysicalModels
```

In *Make/options* file, change the path to include the header files of *specie.so* and *flu-idThermophysicalModels.so* libraries as follows:

```
EXE_INC = \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_SRC)/finiteVolume/lnInclude

LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
    -lfluidThermophysicalModels \
    -lspecie \
    -lfiniteVolume
```

In psiReactionThermo/psiReactionThermos.C, include the header files of new classes and then create new forSRKchungTakaGases macros as follows:

```
#include "SRKchungTakaMixture.H"

...
#include "forSRKchungTakaGases.H"

...
namespace Foam
{
...
    // macros for set of SRK-chung-Taka real-fluid models
    forSRKchungTakaGases(makePsiReactionThermos, SRKchungTakaMixture); //
}
```

In rhoReactionThermo/rhoReactionThermos.C, include the header files of new classes and then create new forSRKchungTakaGases macros as follows:

```
#include "SRKchungTakaMixture.H"

...

#include "forSRKchungTakaGases.H"

...
```

```
namespace Foam
{
...
    // macros for set of SRK-chung-Taka real-fluid models
    forSRKchungTakaGases(makeRhoReactionThermos, SRKchungTakaMixture); //
}
```

Compile to make reaction Thermophysical Models. so.

```
cd ~/OpenFOAM/yourDirectory/.../src/thermophysicalModels/reactionThermo
wclean
wmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

e. Modification in /your Directory/.../src/thermophysical Models/chemistry Model directory:

Create SRKchungTakaStandardChemistryModel class by copying from StandardChemistryModel class.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    thermophysicalModels/chemistryModel/chemistryModel/
cp -rf StandardChemistryModel SRKchungTakaStandardChemistryModel
cd SRKchungTakaStandardChemistryModel
mv StandardChemistryModel.H SRKchungTakaStandardChemistryModel.H
mv StandardChemistryModel.C SRKchungTakaStandardChemistryModel.C
mv StandardChemistryModelI.H SRKchungTakaStandardChemistryModelI.H
```

Open these files and replace StandardChemistryModel and multiComponentMixture by SRKchungTakaStandardChemistryModel and SRKchungTakaMixture, respectively. To make sure you do not miss any thing you should use a command to automatically find and replace a string, for instance use the following command if you are using vim text editor.

```
vi SRKchungTakaStandardChemistryModel.H
:%s/StandardChemistryModel/SRKchungTakaStandardChemistryModel/g
:%s/multiComponentMixture/SRKchungTakaMixture/g
```

It is of importance to note that the name of this class in runtime is defined in the SRKchungTakaStandardChemistryModel.H file as follows:

```
//- Runtime type information
TypeName("SRKchungTakaStandard");
```

In chemistry Model/chemistry Model/basic Chemistry Model/basic Chemistry Model Templates. C file, change the method Name variable to be abe to recognize the name of new SRK chung-Taka Standard Chemistry Model class created so far as follows:

```
. . .
     const word& methodName
 76
 77
 78
       chemistryTypeDict.lookupOrDefault <word>
 79
 80
          "method",
          chemistryTypeDict.lookupOrDefault <bool >("TDAC", false)
 81
 82
 83
          : chemistryTypeDict.lookupOrDefault<bool>("SRKchungTakaStandard"
    , false)
          ? "SRKchungTakaStandard"
 84
          : "standard"
 85
 86
      )
 87);
```

In chemistry Model/reactions/make Reactions. C file, include the header files of for SRK chung-Taka Gases. H and then create for SRK chung Taka Gases macros to make reactions types integrating with real-fluid models as follows:

```
#include "forSRKchungTakaGases.H"

...
namespace Foam
{
...
forSRKchungTakaGases(defineReaction, nullArg);
```

```
. . .
// Irreversible/reversible/non-equilibrium-reversible reactions
for SRK chung Taka Gases (\verb|makeIRNR| eactions|, Arrhenius Reaction Rate);
forSRKchungTakaGases(makeIRNReactions, infiniteReactionRate);
for SRK chung Taka Gases (\verb|makeIRNR| eactions|, Landau Teller Reaction Rate);
for SRK chung Taka Gases (make IRN Reactions, third Body Arrhenius Reaction Rate)
   ;
. . .
// Irreversible/reversible reactions
forSRKchungTakaGases(makeIRReactions, JanevReactionRate);
forSRKchungTakaGases(makeIRReactions, powerSeriesReactionRate);
// Pressure dependent reactions
{\tt for SRKchung Taka Gases}
    makeIRRPressureDependentReactions,
    FallOffReactionRate,
    ArrheniusReactionRate,
    LindemannFallOffFunction
);
forSRKchungTakaGases
    makeIRRPressureDependentReactions,
    FallOffReactionRate,
    ArrheniusReactionRate,
    TroeFallOffFunction
);
forSRKchungTakaGases
(
    makeIRRPressureDependentReactions,
    FallOffReactionRate,
```

```
ArrheniusReactionRate,
    SRIFallOffFunction
);
forSRKchungTakaGases
    makeIRRPressureDependentReactions,
    ChemicallyActivatedReactionRate,
    ArrheniusReactionRate,
    LindemannFallOffFunction
);
{\tt for SRKchung Taka Gases}
(
    makeIRRPressureDependentReactions,
    ChemicallyActivatedReactionRate,
    ArrheniusReactionRate,
    TroeFallOffFunction
);
forSRKchungTakaGases
    {\tt makeIRRPressureDependentReactions}\ ,
    ChemicallyActivatedReactionRate,
    ArrheniusReactionRate,
    SRIFallOffFunction
);
. . .
```

In chemistry Model/reactions/make Langmuir Hinshelwood Reactions. C file, include the header files of for SRK chung Taka Gases. H and then create for SRK chung Taka Gases macros to make reactions types integrating with real-fluid models as follows:

```
...
#include "forSRKchungTakaGases.H"
...
namespace Foam
```

```
{
...
forSRKchungTakaGases(makeIRReactions, LangmuirHinshelwoodReactionRate);
}
```

In chemistryModel/chemistrySolver directory, create a new macros file named makeRe-alFluidChemistrySolver.H to build up chemistry solver models to incorporate the real fluid models as follows.

```
cd ~/OpenFOAM/yourDirectory/.../src/thermophysicalModels/chemistryModel/
    chemistrySolver
cp -rf makeChemistrySolver.H makeRealFluidChemistrySolver.H
```

In makeRealFluidChemistrySolver.H file, replace the old definition of macros by the new one as follows:

```
//*----
#ifndef makeRealFluidChemistrySolver_H
#define makeRealFluidChemistrySolver_H
#include "addToRunTimeSelectionTable.H"
// This can be used for all real-fluid models
#define defineRealFluidChemistrySolver(Model, ReactionThermo,
   ThermoPhysics)
   typedef Model < Reaction Thermo, ThermoPhysics >
       Model ## ReactionThermo ## ThermoPhysics;
   defineTemplateTypeNameAndDebugWithName
   (
       Model ## ReactionThermo ## ThermoPhysics,
           word(Model##ReactionThermo##ThermoPhysics::typeName_())\
            + "<" + ReactionThermo::typeName + ","
            + ThermoPhysics::typeName() + ">"
```

```
).c_str(),
   );
// This also can be used for all real-fluid models
#define makeRealFluidChemistrySolver(Solver, Model, ReactionThermo,\
    ThermoPhysics)
    typedef Solver < Model < Reaction Thermo , ThermoPhysics >>
        Solver##Model##ReactionThermo##ThermoPhysics;
    defineTemplateTypeNameAndDebugWithName
        Solver##Model##ReactionThermo##ThermoPhysics,
            word(Solver##Model##ReactionThermo##
            ThermoPhysics::typeName_())
          + "<" + word(Model < ReactionThermo,
            ThermoPhysics >:: typeName_())
          + "<" + word(ReactionThermo::typeName_()) + ","
          + ThermoPhysics::typeName() + ">>"
        ).c_str(),
   );
    BasicChemistryModel <ReactionThermo >::
        addthermoConstructorToTable
        <Solver##Model##ReactionThermo##ThermoPhysics>
        add##Solver##Model##ReactionThermo##ThermoPhysics##\
thermoConstructorToTable_;
// This is only for SRKchungTaka combined model
```

```
// You can make the same macro for new combined real-fluid model
#define defineSRKchungTakaChemistrySolvers(ReactionThermo,
                                                                      \
    ThermoPhysics)
    defineRealFluidChemistrySolver
        {\tt SRKchungTakaStandardChemistryModel}\ ,
        ReactionThermo,
        ThermoPhysics
    )
// This is only for SRKchungTaka combined model
// You can make the same macro for new combined real-fluid model
#define makeSRKchungTakaChemistrySolvers(Solver, ReactionThermo,
    ThermoPhysics)
    makeRealFluidChemistrySolver
                                                                      \
    (
        Solver,
        SRKchungTakaStandardChemistryModel,
        ReactionThermo,
        ThermoPhysics
    )
#endif
```

In chemistryModel/chemistrySolver/chemistrySolver/chemistrySolvers. C file, include new header files and then create forSRKchungTakaGases macros to make chemistry solver type integrating with real-fluid models as follows:

```
...
#include "SRKchungTakaStandardChemistryModel.H"

#include "forSRKchungTakaGases.H"

#include "makeRealFluidChemistrySolver.H"

...

namespace Foam
```

```
forSRKchungTakaGases(defineSRKchungTakaChemistrySolvers,
    psiReactionThermo);

forSRKchungTakaGases(defineSRKchungTakaChemistrySolvers,
    rhoReactionThermo);
}
```

In chemistryModel/chemistrySolver/EulerImplicit/EulerImplicitChemistrySolvers. C file, include new header files and then create forSRKchungTakaGases macros to make EulerImplicit chemistry solver integrating with real-fluid models as follows:

```
#include "SRKchungTakaStandardChemistryModel.H"

#include "forSRKchungTakaGases.H"

#include "makeRealFluidChemistrySolver.H"

...

namespace Foam

{
    ...
    forSRKchungTakaGases(makeSRKchungTakaChemistrySolvers, EulerImplicit, psiReactionThermo);
    forSRKchungTakaGases(makeSRKchungTakaChemistrySolvers, EulerImplicit, rhoReactionThermo);
}
```

In chemistryModel/chemistrySolver/ode/odeChemistrySolvers.C file, include new header files and then create forSRKchungTakaGases macros to make ode chemistry solver integrating with real-fluid models as follows:

```
#include "SRKchungTakaStandardChemistryModel.H"

#include "forSRKchungTakaGases.H"

#include "makeRealFluidChemistrySolver.H"

...

namespace Foam
```

In chemistryModel/chemistrySolver/noChemistrySolver/noChemistrySolvers.C file, include new header files and then create forSRKchungTakaGases macros to make noChemistry chemistry solver integrating with real-fluid models as follows:

Go to thermophysicalModels/chemistryModel/Make/ directory:

```
cd ~/OpenFOAM/yourDirectory/.../src/thermophysicalModels/chemistryModel/
    Make/
```

In *Make/files* file, change the position to save the binary file, *chemistryModel.so*, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libchemistryModel // save at user directory
```

In *Make/options* file, change the path to include the header files of *specie.so*, *fluidTher-mophysicalModels.so*, and *reactionThermophysicalModels* libraries as follows:

```
EXE_INC = \setminus
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/reactionThermo/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_SRC)/thermophysicalModels/functions/Polynomial \
    -I$(LIB_rfFoam8_SRC)/ODE/lnInclude \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude
LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
    -lfluidThermophysicalModels \
    -lreactionThermophysicalModels \
    -lspecie \
    -10DE \
    -lfiniteVolume \
    -lmeshTools
```

Compile to make chemistryModel.so.

```
cd ~/OpenFOAM/yourDirectory/.../src/thermophysicalModels/chemistryModelwcleanwmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

### 2.2.2. Making interface outside of the thermophysical Models library

Momentum Transport Models, Thermophysical Transport Models, and combustion Models libraries must be updated since the thermophysical Models library is adopted in these libraries. To update these libraries, we should copy them from original OpenFOAM into your directory, then modify the path to link to the new thermophysical Models library that has been made as described above.

Go to /yourDirectory/.../src directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src
cp -rf -p $WM_PROJECT_DIR/src/MomentumTransportModels .
cp -rf -p $WM_PROJECT_DIR/src/ThermophysicalTransportModels .
cp -rf -p $WM_PROJECT_DIR/src/combustionModels .
```

a. Modification in /yourDirectory/.../src/MomentumTransportModels directory.

 $\label{local-constraint} \mbox{Go to } / your Directory/.../src/Momentum Transport Models / momentum Transport Models directory.$ 

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
MomentumTransportModels/momentumTransportModels
```

In momentumTransportModels/Make/files file, change the position to save the object file, momentumTransportModels.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libmomentumTransportModels
```

Compile to make momentum Transport Models.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    MomentumTransportModels/momentumTransportModels
wclean
wmake libso
```

Go to /yourDirectory/.../src/MomentumTransportModels/compressible directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
MomentumTransportModels/compressible
```

In compressible/Make/files file, change the position to save the object file, fluidThermo-MomentumTransportModels.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libfluidThermoMomentumTransportModels
```

In compressible/Make/options file, change the path to include the header files of specie.so and fluidThermophysicalModels.so libraries as follows:

```
EXE_INC = \
   -I../momentumTransportModels/lnInclude \
```

```
-I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
-I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
-I$(LIB_rfFoam8_SRC)/thermophysicalModels/solidThermo/lnInclude \
-I$(LIB_rfFoam8_SRC)/thermophysicalModels/solidSpecie/lnInclude \
-I$(LIB_SRC)/finiteVolume/lnInclude \
-I$(LIB_SRC)/meshTools/lnInclude \

LIB_LIBS = \
-L$(FOAM_USER_LIBBIN) \
-lfluidThermophysicalModels \
-lsolidThermo \
-lmomentumTransportModels \
-lspecie \
-lfiniteVolume \
-lmeshTools
```

Compile to make fluidThermoMomentumTransportModels.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    MomentumTransportModels/compressible
wclean
wmake libso
```

b. Modification in /yourDirectory/.../src/ThermophysicalTransportModels directory. Go to /yourDirectory/.../src/ThermophysicalTransportModels directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
ThermophysicalTransportModels
```

In Make/files file, change the position to save the object file, thermophysicalTransport-Models.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libthermophysicalTransportModels
```

In *Make/options* file, change the path to include the header files of *specie.so*, *fluidTher-mophysicalModels.so*, and *momentumTransportModels.so* libraries as follows:

```
EXE_INC = \
```

```
-I$(LIB_rfFoam8_SRC)/MomentumTransportModels/momentumTransportModels/
     lnInclude \
  -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/compressible/lnInclude \
  -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
  -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
  -I$(LIB_rfFoam8_SRC)/thermophysicalModels/solidThermo/lnInclude \
  -I\$(LIB\_rfFoam8\_SRC)/thermophysicalModels/solidSpecie/lnInclude \ \setminus \\
  -I$(LIB_SRC)/finiteVolume/lnInclude \
  -I$(LIB_SRC)/meshTools/lnInclude \
LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
    -lfluidThermophysicalModels \
    -lsolidThermo \
    -1momentumTransportModels \
    -lspecie \
    -lfiniteVolume \
    -lmeshTools
```

Compile to make thermophysical Transport Models.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    ThermophysicalTransportModels
wclean
wmake libso
```

Then, go to /your Directory/.../src/Thermophysical Transport Models/psiReaction Thermo directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
ThermophysicalTransportModels/psiReactionThermo
```

In Make/files file, change the position to save the object file, psiReactionThermophysicalTransportModels.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libpsiReactionThermophysicalTransportModels
```

In *Make/options* file, change the path to include the header files of *specie.so*, *fluidTher-mophysicalModels.so*, *reactionThermophysicalModels*, and *momentumTransportModels.so* libraries as follows:

```
EXE_INC = \setminus
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/momentumTransportModels/
       lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/compressible/lnInclude \
    -I\$(LIB\_rfFoam8\_SRC)/ThermophysicalTransportModels/lnInclude \ \setminus \\
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/reactionThermo/lnInclude \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude \
LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
    -lfluidThermophysicalModels \
    -lreactionThermophysicalModels \
    -1momentumTransportModels \
    -lspecie \
    -lfiniteVolume \
    -lmeshTools
```

Compile to make psiReactionThermophysicalTransportModels.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    ThermophysicalTransportModels/psiReactionThermo/
wclean
wmake libso
```

Then, go to /your Directory/.../src/Thermophysical Transport Models/rhoReaction Thermodirectory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
ThermophysicalTransportModels/rhoReactionThermo
```

In Make/files file, change the position to save the object file, rhoReactionThermophysicalTransportModels.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/librhoReactionThermophysicalTransportModels
```

In *Make/options* file, change the path to include the header files of *specie.so*, *fluidTher-mophysicalModels.so*, *reactionThermophysicalModels*, and *momentumTransportModels.so* libraries as follows:

```
EXE_INC = \setminus
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/momentumTransportModels/
       lnInclude \
   -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/compressible/lnInclude \
    -I$(LIB_rfFoam8_SRC)/ThermophysicalTransportModels/lnInclude \
   -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
   -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
   -I$(LIB_rfFoam8_SRC)/thermophysicalModels/reactionThermo/lnInclude \
   -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude \
LIB_LIBS = \
   -L$(FOAM_USER_LIBBIN) \
   -lfluidThermophysicalModels \
   -lreactionThermophysicalModels \
   -lmomentumTransportModels \
   -lspecie ∖
   -lfiniteVolume \
    -lmeshTools
```

Compile to make rhoReactionThermophysicalTransportModels.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    ThermophysicalTransportModels/rhoReactionThermo/
wclean
wmake libso
```

c. Modification in /yourDirectory/.../src/radiationModels directory.

Go to /yourDirectory/.../src/radiationModels directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/radiationModels
```

In *Make/files* file, change the position to save the object file, *radiationModels.so*, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libradiationModels
```

In Make/options file, change the path to include the header files of chemistryModel.so, momentumTransportModels.so, and fluidThermoMomentumTransportModels.so libraries as follows:

```
EXE_INC = \setminus
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/solidThermo/lnInclude \
    -I$(LIB_SRC)/thermophysicalModels/SLGThermo/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/thermophysicalProperties/
       lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/reactionThermo/lnInclude \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude
LIB_LIBS = \
    -L$(FOAM_USER_LIBBIN) \
    -lfluidThermophysicalModels \
    -lspecie ∖
    -lsolidThermo \
    -1SLGThermo \
    -lthermophysicalProperties \
    -lfiniteVolume \
    -lmeshTools
```

Compile to make radiationModels.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
  radiationModels
wclean
wmake libso
```

d. Modification in /yourDirectory/.../src/combustionModels directory.

Go to /yourDirectory/.../src/combustionModels directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
combustionModels
```

In *Make/files* file, change the position to save the object file, *combustionModels.so*, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libcombustionModels
```

In Make/options file, change the path to include the header files of chemistryModel.so, momentumTransportModels.so, fluidThermoMomentumTransportModels.so, and radiation-Models.so libraries as follows:

```
EXE_INC = \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/reactionThermo/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/chemistryModel/lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/momentumTransportModels/
    lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/compressible/lnInclude \
    -I$(LIB_rfFoam8_SRC)/radiationModels/lnInclude \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude \
```

```
-lchemistryModel \
-lradiationModels \
-lfiniteVolume \
-lmeshTools
```

Compile to make combustionModels.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
    combustionModels
wclean
wmake libso
```

e. Modification in /yourDirectory/.../src/fvOptions directory.

Go to /yourDirectory/.../src/fvOptions directory.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/fvOptions
```

In Make/files file, change the position to save the object file, fvOptions.so, at user directory.

```
...

LIB = $(FOAM_USER_LIBBIN)/libfvOptions
```

In *Make/options* file, change the path to include the header files of *momentumTransport-Models.so*, *fluidThermoMomentumTransportModels.so*, and *thermophysicalTransportModels.so* libraries as follows:

```
EXE_INC = \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude \
    -I$(LIB_SRC)/sampling/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/solidThermo/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/momentumTransportModels/
    lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/compressible/lnInclude \
    -I$(LIB_rfFoam8_SRC)/ThermophysicalTransportModels/lnInclude
```

```
LIB_LIBS = \
-L$(FOAM_USER_LIBBIN) \
-lfiniteVolume \
-lsampling \
-lmeshTools \
-lmomentumTransportModels \
-lfluidThermoMomentumTransportModels \
-lthermophysicalTransportModels
```

Compile to make fvOptions.so.

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/fvOptions
wclean
wmake libso
```

#### 2.2.3. Test new created runtime thermo-packages

reactingFoam solver is selected to check the availability of real-fluid runtime thermopackages, in which 2D laminar counterflow non-premixed flame is solved as a test case.

To use reactingFoam, we need to link it to the new libraries that we created so far. First, go to reactingFoam directory.

```
sol
cd combustion/reactingFoam
```

Modify *Make/options* file as follows:

```
EXE_INC = \
    -I$(LIB_SRC)/finiteVolume/lnInclude \
    -I$(LIB_SRC)/meshTools/lnInclude \
    -I$(LIB_SRC)/sampling/lnInclude \
    -I$(LIB_SRC)/sampling/lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/momentumTransportModels/
    lnInclude \
    -I$(LIB_rfFoam8_SRC)/MomentumTransportModels/compressible/lnInclude \
    -I$(LIB_rfFoam8_SRC)/ThermophysicalTransportModels/lnInclude \
    -I$(LIB_rfFoam8_SRC)/ThermophysicalTransportModels/psiReactionThermo/
    lnInclude \
```

```
-I$(LIB_rfFoam8_SRC)/thermophysicalModels/specie/lnInclude \
   -I$(LIB_rfFoam8_SRC)/thermophysicalModels/reactionThermo/lnInclude \
   -I$(LIB_rfFoam8_SRC)/thermophysicalModels/basic/lnInclude \
   -I$(LIB_rfFoam8_SRC)/thermophysicalModels/chemistryModel/lnInclude \
   -I$(LIB_rfFoam8_SRC)/ODE/lnInclude \
   -I$(LIB_rfFoam8_SRC)/combustionModels/lnInclude
EXE_LIBS = \
   -L$(FOAM_USER_LIBBIN) \
   -lfiniteVolume \
   -lfvOptions \
   -lmeshTools \
   -lsampling \
   -lmomentumTransportModels \
   -lfluidThermoMomentumTransportModels \
   -lthermophysicalTransportModels \
   -lpsiReactionThermophysicalTransportModels \
   -lreactionThermophysicalModels \
   -lspecie \
   -lfluidThermophysicalModels \
   -lchemistryModel \
   -10DE \
    -lcombustionModels
```

Then, recompile it.

```
sol
cd combustion/reactingFoam
wclean
wmake
```

Now, reactingFoam is ready to use. Go to 2D laminar counterflow flame test case in OpenFOAM.

```
tut
cd combustion/reactingFoam/laminar/counterFlowFlame2D
```

Generate mesh and check with ideal gas models first.

```
blockMesh reactingFoam
```

It should be executed without error.

Specify the thermoType dictionary in constant/thermophysicalProperties dictionary file to use new created runtime thermo-packages as follows:

```
//This is a new thermo packages we have created so far
thermoType
{
                    hePsiThermo;
    type
    mixture
                     SRKchungTakaMixture;
    transport
                     chung Taka;
                    rfJanaf;
    thermo
                     sensibleEnthalpy;
    energy
    equationOfState soaveRedlichKwong;
    specie
                     rfSpecie;
```

Specify the chemistry Type dictionary in constant/chemistry Properties dictionary file to use new created runtime thermo-packages as follows:

```
chemistryType
{
    solver EulerImplicit;
    method SRKchungTakaStandard; //new chemistry model for real fluid
}
```

Then, execute the reactingFoam again.

```
reactingFoam
```

It should be executed without error. So far, we have successfully created new runtime selectable thermo-packages for real-fluid THERMPHYS models. Note that the THERMPHYS models of these new thermo-packages here are ideal gas models since we have not implemented the actual real-fluid models yet.

### 3. Detailed implementation of real-fluid models

The details of real-fluid models are described in [1–4]. Since the implemented source code of real-fluid models are too long to be described in details in this document, readers are recommended referring directly to our source code for more convenience. The following classes should be replaced by our source files: rfSpecie, soaveRedlichKwong, rfJanafThermo, chung-TakaTransport, and SRKchungTakaMixture. It is of importance to note that SRKchung-TakaStandardChemistryModel is also a new created class but it does not need to be changed so far.

# 4. Detailed modification of existing classes

There are several groups of classes have to be modified to be compatible with new real-fluid model classes due to the use of templates in OpenFOAM.

#### 4.1. Modification of Transport classes

All classes inside *Transport* group involving *constTransport*, *logPolynomialTransport*, *polynomialTransport*, *sutherlandTransport*, etc., have to be modified as follows:

In \*.H file, create a new function as:

```
public:
...
// Species diffusivity
inline scalar Dimix(label speciei, const scalar p, const scalar T) const
{
    return 1.0;
}
...
```

Goto thermophysical Models/specie and compile this library again after modifying the code:

```
cd ~/OpenFOAM/yourDirectory/src/thermophysicalModels/specie
wmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

### 4.2. Modification of Mixture classes

In thermophysical Models/reaction Thermo/mixtures directory, modify single Component-Mixture, egr Mixture, homogeneous Mixture, inhomogeneous Mixture, and very Inhomogeneous-Mixture classes as follows:

### In \*.H file:

```
#include "PtrList.H"
private:
    // Number of species
    label numberOfSpecies_;
    // List of specie thermo
    PtrList < ThermoType > specieThermos_;
public:
    // Return the raw specie thermodynamic data
    const PtrList < ThermoType > & specieThermos()
    {
        return specieThermos_;
    }
    // Return the number of species in the mixture
    inline const label& numberOfSpecies() const
        return numberOfSpecies_;
    }
```

In \*.C file:

```
// In constructor:
. . .
{
. . .
    specieThermos_(1);
    specieThermos_.set
       (
          0,
          new ThermoType(thermoDict.subDict("mixture"))
      );
       // note that subDict's name here differs from class by class.
      // Particularly, it is:
      // "mixture" in singleComponentMixture class;
       // "fuel" in egrMixture class;
      // "reactants" in homogeneousMixture class;
       // "fuel" in inhomogeneousMixture class;
       // "fuel" in veryInhomogeneousMixture class;
   numberOfSpecies_ = specieThermos_.size();
```

In thermophysical Models/reaction Thermo/mixtures directory, modify the multiCompo-nent Mixture class as follows:

In multiComponentMixture.H file:

```
protected:
    label numberOfSpecies_;
...
public:
...
```

```
inline const label& numberOfSpecies() const
{
    return numberOfSpecies_;
}
```

In multiComponentMixture.C file:

```
// In constructor:
template < class ThermoType >
Foam::multiComponentMixture < ThermoType > ::multiComponentMixture
    const dictionary& thermoDict,
    const fvMesh& mesh,
    const word& phaseName
    basicSpecieMixture
    (
        thermoDict,
        thermoDict.lookup("species"),
        mesh,
        phaseName
    ),
    specieThermos_(readSpeciesData(thermoDict)),
    {\tt speciesComposition\_(readSpeciesComposition(thermoDict, species())),}
    mixture_("mixture", specieThermos_[0]),
    mixtureVol_("volMixture", specieThermos_[0]),
    numberOfSpecies_(species_.size()) //
{
    correctMassFractions();
```

In thermophysical Models/basic/mixtures directory, modify the pure Mixture class as follows:

In pureMixture.H file:

```
// private data
// - Number of species
    label numberOfSpecies_;

//- List of specie thermo
PtrList<ThermoType> specieThermos_;

...

public:
...

// - Return the number of species in the mixture
inline const label& numberOfSpecies() const
{
    return numberOfSpecies_;
}

//- Return the raw specie thermodynamic data
    const PtrList<ThermoType>& specieThermos() const
{
    return specieThermos_;
}
...
```

In pureMixture.C file:

Goto thermophysical Models/basic and compile this library again after modifying the code:

```
cd ~/OpenFOAM/yourDirectory/src/thermophysicalModels/basic
wmake libso
```

If errors occur, check carefully your steps again to make sure you do not miss any thing.

## 4.3. Modification of BasicThermo classes

In psiThermo and rhoThermo class, add these following public virtual functions into \*.H files. These functions will be overridden by functions in their child classes due to polymorphism.

```
//- Diffusion coefficient of specie ith in the mixture [m^2/s]
virtual tmp<volScalarField> Dimix(const label speciei) const = 0;

//- Diffusion coefficient of specie ith in the mixture for patch [m^2/s]
virtual tmp<scalarField> Dimix(const label speciei, const label patchi)
        const = 0;

//- New functions for realFluidRhoReactingFoam solver
```

```
//- Enthalpy/Internal energy of specie ith [J/kg]
virtual tmp<volScalarField> hei(label speciei) const = 0;

//- Enthalpy/Internal energy of specie ith for patch [J/kg]
virtual tmp<scalarField> hei(label speciei, const label patchi) const = 0;
...
```

## 4.4. Modification of Type classes

In heThermo class, add a these following code.

In heThermo.H file:

```
protected:
//- Store List of energy field of indivisual species
PtrList<volScalarField> heList_;

public:
...
// Access to thermodynamic state variables
//- New functions for realFluidReactingFoam solver
//- Return he[J/kg] of indivisual speciei
    virtual tmp<volScalarField> hei(label speciei) const;
//- Return he[J/kg] of indivisual speciei
    virtual tmp<scalarField> hei(label speciei)
```

In heThermo.C file:

```
// In both constructor functions
...
:
...
heList_(MixtureType::numberOfSpecies())
{
    forAll(heList_, i)
```

```
{
    heList_.set
    (
        i,
        new volScalarField
            IOobject
            (
                "hei",
                mesh.time().timeName(),
                mesh,
                IOobject::NO_READ,
                IOobject::NO_WRITE
            ),
            mesh,
            he_.dimensions()
       )
    );
}
forAll(WList_, i)
{
    WList_.set
        i,
        new volScalarField
        (
            IOobject
            (
                 mesh.time().timeName(),
                 mesh,
                IOobject::NO_READ,
                IOobject::NO_WRITE,
```

```
false
            ),
            mesh,
            dimMass/dimMoles
        )
    );
}
//- Calculate he_i, Wi internal fields for individual species
const PtrList<typename MixtureType::thermoType>& speciesData_ = this->
   specieThermos();
forAll(heList_, i)
    forAll(heList_[i].primitiveFieldRef(), celli)
    {
        heList_[i].primitiveFieldRef()[celli]
      = speciesData_[i].HE(pCells[celli], TCells[celli]);
        WList_[i].primitiveFieldRef()[celli]
      = speciesData_[i].W();
    }
}
//
//- Calculate he_i, Wi boundary fiels for individual species
forAll(WList_, i)
{
    forAll(WList_[i].boundaryFieldRef(), patchi)
       forAll(WList_[i].boundaryFieldRef()[patchi], facei)
       {
           WList_[i].boundaryFieldRef()[patchi][facei]
         = speciesData_[i].W();
```

```
heList_[i].boundaryFieldRef()[patchi][facei]
             = speciesData_[i].HE
                    this ->p_.boundaryField()[patchi][facei],
                    this ->T_.boundaryField()[patchi][facei]
               );
           }
        }
    }
    //
// New functions for realFluidReactingFoam solver
//- Return he_i of individual species ith
template < class BasicThermo, class MixtureType >
Foam::tmp<Foam::volScalarField>
Foam::heThermo <BasicThermo, MixtureType >::hei
    label speciei
) const
    return heList_[speciei];
template < class BasicThermo, class MixtureType >
Foam::tmp<Foam::scalarField>
Foam::heThermo <BasicThermo, MixtureType >::hei
(
   label speciei,
   const label patchi
) const
```

```
{
    return heList_[speciei].boundaryField()[patchi];
}
```

In hePsiThermo class, add a these following code. In hePsiThermo.H file:

```
private:
//- Store list of mass diffusion coefficients
PtrList<volScalarField> Dimix_;
...

public:
...
//- Diffusion coefficient of specie ith in the mixture [m^2/s]
virtual tmp<volScalarField> Dimix(const label speciei) const
{
    return Dimix_[speciei];
}

//- Diffusion coefficient of specie ith in the mixture for patch [m^2/s]
virtual tmp<scalarField> Dimix(const label speciei, const label patchi)
    const
{
    return Dimix_[speciei].boundaryField()[patchi];
}
...
```

In hePsiThermo.C file:

```
// In constructor function
:
...
Dimix_(MixtureType::numberOfSpecies())
{
    forAll(Dimix_, i)
    {
```

```
Dimix_.set
        (
            i,
            new volScalarField
                IOobject
                (
                     this->phasePropertyName("thermo:Dimix"),
                     mesh.time().timeName(),
                    mesh,
                    IOobject::NO_READ,
                    IOobject::NO_WRITE
                ),
                mesh,
                dimensionSet(0, 2, -1, 0, 0)
            )
        );
    }
}
//In calculate() function
const PtrList<typename MixtureType::thermoType>& speciesData_ = this->
   specieThermos();
forAll(TCells, celli)
{
    forAll(Dimix_, i)
    {
        Dimix_[i].primitiveFieldRef()[celli]
      = mixture_.Dimix(i, pCells[celli], TCells[celli]);
    }
```

```
forAll(this->T_.boundaryField(), patchi)
    . . .
    if (pT.fixesValue())
    {
        forAll(pT, facei)
             . . .
            forAll(Dimix_, i)
            {
                 Dimix_[i].boundaryFieldRef()[patchi][facei]
                 = mixture_.Dimix(i, pp[facei], pT[facei]);
            }
        }
    }
    else
    {
        forAll(pT, facei)
        {
            forAll(Dimix_, i)
            {
                 Dimix_[i].boundaryFieldRef()[patchi][facei]
                 = mixture_.Dimix(i, pp[facei], pT[facei]);
            }
        }
    }
```

In heRhoThermo class, add a these following code. In heRhoThermo.H file:

```
private:
//- Store list of mass diffusion coefficients
PtrList < volScalarField > Dimix_;
```

```
public:
...
//- Diffusion coefficient of specie ith in the mixture [m^2/s]
virtual tmp<volScalarField> Dimix(const label speciei) const
{
    return Dimix_[speciei];
}

//- Diffusion coefficient of specie ith in the mixture for patch [m^2/s]
virtual tmp<scalarField> Dimix(const label speciei, const label patchi)
    const
{
    return Dimix_[speciei].boundaryField()[patchi];
}
...
```

In heRhoThermo.C file:

```
mesh.time().timeName(),
                     mesh,
                     IOobject::NO_READ,
                     IOobject::NO_WRITE
                ),
                mesh,
                dimensionSet(0, 2, -1, 0, 0)
            )
        );
    }
. . .
//In calculate() function
const PtrList<typename MixtureType::thermoType>& speciesData_ = this->
   specieThermos();
//Diffusion coefficient has not been implemented yet for rho-based systems
forAll(TCells, celli)
{
   forAll(Dimix_, i)
        Dimix_[i].primitiveFieldRef()[celli] = 1.0;
    }
. . .
}
forAll(this->T_.boundaryField(), patchi)
{
    . . .
   if (pT.fixesValue())
```

```
forAll(pT, facei)
    {
        forAll(Dimix_, i)
             Dimix_[i].boundaryFieldRef()[patchi][facei] = 1.0;
        }
    }
}
else
{
    forAll(pT, facei)
    {
    . . .
        forAll(Dimix_, i)
        {
             Dimix_[i].boundaryFieldRef()[patchi][facei] = 1.0;
        }
    }
}
```

In thermophysical Models/reaction Thermo/psiuReaction Thermo directory, modify hehe-uPsiThermo class as follows: In heheuPsiThermo.H file:

```
private:
//- Store list of mass diffusion coefficients
PtrList < volScalarField > Dimix_;
...
public:
...
//- Diffusion coefficient of specie ith in the mixture [m^2/s]
virtual tmp < volScalarField > Dimix (const label speciei) const
{
    return Dimix_[speciei];
```

```
//- Diffusion coefficient of specie ith in the mixture for patch [m^2/s]
virtual tmp<scalarField> Dimix(const label speciei, const label patchi)
    const
{
    return Dimix_[speciei].boundaryField()[patchi];
}
```

In heheuPsiThermo.C file:

```
// In constructor
Dimix_(MixtureType::numberOfSpecies())
{
. . .
    this->heuBoundaryCorrection(this->heu_);
    //
    forAll(Dimix_, i)
    {
        Dimix_.set
        (
            i,
            new volScalarField
                IOobject
                 (
                     this->phasePropertyName("thermo:Dimix"),
                     mesh.time().timeName(),
                     mesh,
                     IOobject::NO_READ,
                    IOobject::NO_WRITE
                ),
```

```
mesh,
                dimensionSet(0, 2, -1, 0, 0)
            )
        );
   }
    //
    calculate();
//In calculate() function
//Diffusion coefficient has not been implemented yet for psiu-based
const PtrList<typename MixtureType::thermoType>& speciesData_ = this->
   specieThermos();
forAll(TCells, celli)
{
. . .
   forAll(Dimix_, i)
    {
        Dimix_[i].primitiveFieldRef()[celli] = 1.0;
    }
forAll(this->T_.boundaryField(), patchi)
{
   if (pT.fixesValue())
        forAll(pT, facei)
        {
            forAll(Dimix_, i)
```

```
{
        Dimix_[i].boundaryFieldRef()[patchi][facei] = 1.0;
}

}
else
{
    forAll(pT, facei)
    {
        ...
        forAll(Dimix_, i)
        {
            Dimix_[i].boundaryFieldRef()[patchi][facei] = 1.0;
        }
    }
}
```

Compile all libraries again after implementing real-fluid models and modifying some existing classes as mentioned above as follows (in order):

```
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
thermophysicalModels/basic
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
thermophysicalModels/reactionThermo
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
thermophysicalModels/chemistryModel
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
MomentumTransportModels/momentumTransportModels
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels -8/src/
MomentumTransportModels/compressible
```

```
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
   Thermophysical Transport {\tt Models}
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
   ThermophysicalTransportModels/psiReactionThermo
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
   ThermophysicalTransportModels/rhoReactionThermo
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
   radiationModels
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/
   combustionModels
wmake libso
cd ~/OpenFOAM/yourDirectory/realFluidThermophysicalModels-8/src/fvOptions
wmake libso
```

If there is no error, the real-fluid based thermophysical Models now is ready to use. It is of importance to note that any solver that utilize thermophysical Models, momentum Transport Models, fluid Momentum Transport Models, thermophysical Transport Models, and combustion Models should be recompiled after implementing real-fluid based thermophysical Models.

### 5. Using the new library

The new library can be used for any reacting flow solver in OpenFOAM 8.0 that adopt a set of implemented real-fluid models by using either psiReactionThermo or rhoReactionThermo classes. To use these classes, their header files (psiReactionThermo.H and rhoReactionThermo.H) should be included in a source file of the solver. When an object of these classes is created, we can call their functions to return the corresponding thermophysical properties. The following piece of code demonstrates how to utilize the new library using the psiReactionThermo class in realFluidReactingFoam developed from reactingFoam:

```
//Create an object named thermo of psiReactionThermo type.
autoPtr < psiReactionThermo > pThermo (psiReactionThermo :: New (mesh));
psiReactionThermo & thermo = pThermo();
...
// call functions to return THERMPHYS properties
thermo.rho(); // return the density field
thermo.Dimix(i); // return the mass diffusivity of specie ith
```

In the running case directory, the *thermoType* dictionary needs to be specified in the *constant/thermophysicalProperties* file as follows:

```
thermoType
{
   type
                   hePsiThermo;
                                      // heRhoThermo for rho-based system.
                    SRKchungTakaMixture;
   mixture
   transport
                    chungTaka;
                    rfJanaf;
   thermo
                    sensibleEnthalpy; //or sensibleInternalEnergy
   energy
   equationOfState soaveRedlichKwong;
   specie
                    rfSpecie;
```

In the running case directory, the *chemistryType* dictionary needs to be specified in the constant/chemistryProperties file as follows:

```
chemistryType
{
    solver EulerImplicit; //or "ode" or "none"
    method SRKchungTakaStandard;
}
```

Some input data for real-fluid calculations also need to be specified at dataForRealFluid entry in the constant/thermo.compressibleGas dictionary file for each species in the following format:

```
dataForRealFluid // of Oxygen species
```

```
Тс
       154.58;
                // the critical temperature, K
Рς
       5.043;
                // the critical pressure, MPa
                // the critical volume, cm^3/mol
Vс
       73.529;
                // the dimensionless acentric factor
omega
       0.025;
kappai 0.0;
                // the dimensionless association factor
miui
       0.0;
                   the dimensionless dipole moment
sigmvi 16.6;
                // the dimensionless diffusion volume
```

Readers are referred to [7] to find input data of other species, and referred to the source code of the *realFluidReactingFoam* and tutorials for test cases provided in the our repository for better understanding of using the new library in a solver.

The original reactingFoam solver can be used for testing real fluid implementation processes as mentioned in sec-2.2.3 but the result is not guaranteed since its governing equations have not been validated for real fluid models yet. We highly recommend readers using provided realFluidReactingFoam solver. It is a quasi direct numerical simulation (quasi-DNS) solver developed based on a work of Li et al. [8] in which its governing equations have been validated at low pressure. In the current work, we have validated for laminar reacting flows under both low pressure and supercritical conditions, up to 200 atm (see our paper for details of the validation). This solver will be updated and validated for reacting turbulence flows in the future.

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