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1 Introduction to hyFoam

hyFoam is a newly coded open-source single-temperature computational fluid dynamics (CFD) solver implemented within the OpenFOAM framework. It is based on the code developments made for hy2Foam. The code solves the Navier–Stokes–Fourier equations and it has the capability to model a chemically-reacting environment in the continuum regime. Various transport models have been implemented to determine species and mixture thermal properties at high-temperatures. There are also several new species diffusion models and boundary conditions available.

2 Installation of *hyFoam*

The installation of *hyFoam* is straightforward. Please make sure that the path to your OpenFOAM user directory is correct in the bash script called *install.sh* before execution. This script will install the new *strath* libraries located into the *src* folder, will install the *hyFoam* solver located into the *applications* folder (as well as a few other apps) and will copy some tutorials into your *run* directory.

3 hyFoam working environment

3.1 The main folder

The main working folder is no different from any other OpenFOAM solvers. The bash script *Allwelean* removes any temporary files within the main directory.

```
referenceCase
VincePC: referenceCase >ls
0 Allwclean constant gnuplot pv.foam system
VincePC: referenceCase >
```

Figure 1: Main folder contents

NB: The command line to run hyFoam is hy2Foam.

3.2 The θ folder

The $\boldsymbol{0}$ folder contains the traditional \boldsymbol{p} and \boldsymbol{U} files. The other mandatory files are the temperature, printed as \boldsymbol{Tt} , and the mass-fraction for each species. The prefix Y_{-} is omitted which means that the $\mathbf{N2}$ file is the mass-fraction of N_2 .

```
referenceCase
                                            Terminal
VincePC: referenceCase >ls 0
                                     Τt
                            02
                                            Tv N
                                                    Tv_N2
                            02+
             N2+
                   NO+
                        0+
                                     Tv e-
                                            Tv_N+
                                                   Tv_N2+
                                                            Tv_NO+
                                                                    Tv_0+
                                                                            Tv_02+
VincePC: referenceCase >ls 0/include/
boundaries initialConditions
VincePC: referenceCase >
```

Figure 2: The 0 folder contents

It is good practice to initialise the fields via an include statement to a dictionary and to set some of the boundary conditions by calling another dictionary. These dictionaries are regrouped into the 0/include folder. For instance, the initialConditions dictionary can be defined as follow.

```
1000;
inletPressure 1.01325e5:
inletVelocity (800 0 0);
initialVelocityField (0 0 0);
Y_N2
         0.767:
Y_02
         0.233;
Y NO
         0:
Y_C02
          0;
Y_H20
          0;
Y CO
          0;
Y_N
          0;
Y_0
    while the boundaries dictionary is for the case of an adiabatic heat bath
fixedWalls
    type
                    empty;
```

3.3 The *constant* folder

The **constant** folder regroups all the dictionaries relative to physicochemical phenomena occurring at supersonic/hypersonic speed and that have been implemented into *hy2Foam*. The mesh remains located into the **polyMesh** sub-folder.

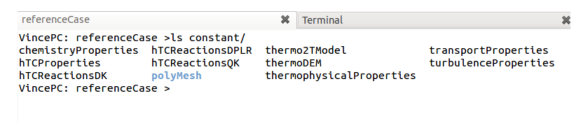


Figure 3: The *constant* folder contents

Recently, a new folder named *chemDicts* (not shown in Figure 3) has been created in *constant* to contain all chemistry dictionaries. It is advised to make a copy inside the *constant* directory before starting working on it.

A short description of the different dictionaries is given below. It is recommended to follow the order listed here while editing the entries of the dictionaries.

- 1) thermophysicalProperties: the standard OpenFOAM dictionary that prescribes which thermophysical models are going to be used and the location of some sub-dictionaries.
- 2) turbulenceProperties: the standard OpenFOAM dictionary that deals with turbulence modelling. In hy2Foam, the options are laminar and RAS2Model.

- 3a) hTCProperties: the dictionary that allows the user to activate or deactivate the high-temperature chemistry (hTC) features and to output some desired species-related fields such as number densities or vibrational degrees of freedom.
- 3b) chemistryProperties: the dictionary that defines the different chemistry inputs
- 3c) hTCReactionsDPLR/QK/DK/ES/...: the list of the species composing the mixture is defined in those dictionaries. It is strongly recommended not to change the order of the species in the list (molecules ions atoms free electrons). Please uncomment the species and chemical reactions you may need.
- 4) *thermoDEM*: the dictionary that provides the species thermodynamic data for the different species loaded in dictionary 3c).
- 5) transportProperties: the dictionary that defines the models used for species diffusion and the mixing rule. Some parameters such as the mean free path or the local gradientlength Knudsen number can be printed from this dictionary.

In the subsequent subsections, the main features of the dictionaries listed above are reviewed.

3.3.1 The thermophysical Properties dictionary

```
FoamFile
   version
              2.0;
   format
              ascii:
              dictionary:
   class
   location
              "constant":
              thermophysicalProperties;
   obiect
    thermoType
                heRho2Thermo:
   type
               reacting2Mixture;
   mixture
                 BlottnerEucken;
   transport
                 decoupledEnergyModes;
   thermo
                 sensible2InternalEnergy;
   equationOfState perfect2Gas;
   specie
                 advancedSpecie;
hyLight yes;
downgradeToSingleTv no;
downgradeToSingleTemperature yes;
chemistryReader foam2ChemistryReader;
foamChemistryFile "$FOAM_CASE/constant/hTCReactionsJ92";
foamChemistryThermoFile "$FOAM_CASE/constant/thermoDEM";
temperatureBounds
{
   Tlow
          200:
         22000;
   Thigh
```

The thermoType defined in the thermophysicalProperties dictionary uses the new strath

thermophysical models. The user has the choice between the four transport models listed in Table 1.

Model name	shear viscosity	thermal diffusivity
constant	constant	constant Eucken
BlottnerEucken	Blottner	Eucken
powerLawEucken	power law	Eucken
SutherlandEucken	Sutherland	Eucken

Table 1: Available transport models in hyFoam

hyFoam is a single-temperature solver. The switch downgradeToSingleTemperature should always be turned to 'yes', as shown above. hyLight is an option that permits to run a light version of hyFoam, thus saving time. The temperature bounds are here to make the code more robust during the transient phase. Please make sure that the temperature field is not reaching these bounds once the desired solution is obtained.

The foamChemistryFile entry tells which set of chemical rate constants is going to be used by linking to the appropriate sub-dictionary. There are various sets of chemical rate constants available and their respective meanings are listed in Table 2.

Set	Meaning
DPLR	Park's rates (used in the DPLR solver), will soon be replaced
QK	Quantum Kinetics rates
DK	Dunn and Kang rates
ES	Evans and Shexnayder (1980)
J92	Jachimoski's rates (1992)
J92_fwd	Jachimoski's rates (1992), forward only
Mars94	Park's rates (1994)
Earth93	Park's rates (1993), available soon (to replace DPLR)

Table 2: Available chemical rate constant sets in hyFoam

3.3.2 The turbulenceProperties dictionary

As stated earlier, the turbulence models (simulationType) available in hyFoam are laminar and RAS2Model.

If RAS2Model is chosen, then the definition of a new dictionary, RASProperties, is required

```
//******************************//
FoamFile
{
   version   2.0;
   format   ascii;
```

The different choices for the RAS2Model entry are laminar, kOmegaSST, kEpsilon and SpalartAllmaras.

3.3.3 The hTCProperties dictionary

The high-temperature chemistry library is the high-level strath library. The hTC2Model entry should be consistent with the turbulent model chosen in turbulenceProperties. Please turn off the active switch if the chemistry module does not need to be loaded. The molar-fractions are printed with the prefix X_{-} , the number densities with the prefix nD_{-} , the partial densities with the prefix pD_{-} , and finally the partial pressures with the prefix pP_{-} .

```
FoamFile
   version
            2.0;
            ascii;
   format
   class
            dictionary;
   location
            "constant";
            hTCProperties;
   object
hTC2Model laminar2<rho2ChemistryHTC>;
active
       true:
laminar2Coeffs
{}
chemistryOutputs
   molarFraction
                  off;
   numberDensity
                  off:
   partialDensity
                  off;
   partialPressure
                  off;
```

3.3.4 The *chemistryProperties* dictionary

hyFoam uses finite rate chemistry. This translates into the use of the new class Euler2Implicit. Please turn on/off the chemistry switch to enable/disable chemical reactions. The initialChemicalTimeStep option should be available soon.

3.3.5 The hTCReactionsDPLR/QK/DK/ES dictionaries

QK reactions should be defined as irreversible while Park (DPLR) reactions can be considered either as reversible or irreversible. Printed below is an example of the implementation of one chemical reaction in the hTCReactionsDPLR dictionary

```
FoamFile
  version
           2.0:
          ascii;
  format
           dictionary;
  class
  object
           hTCReactionsDPLR;
             species
   //N2
  02
  //NO
  //N2+
  //02+
  //NO+
  Π
  //N+
  //0+
   //e-
/* REACTIONS NO 1 TO 15 AND 18 TO 32 -> DISSOCIATION
  REACTIONS NO 16 AND 17
                         -> EXCHANGE
  REACTION NO 33
                          -> ELECTRON IMPACT DISSOCIATION
                          -> ELECTRON IMPACT IONIZATION
  REACTIONS NO 34 AND 35
  REACTIONS NO 36 TO 38
                          -> ASSOCIATIVE IONIZATION
  REACTIONS NO 39 TO 49
                          -> CHARGE EXCHANGE
reactions
  // Reaction no 1
  \verb"oxygenAtomicNitrogenReaction"
           reversibleArrheniusReaction;
     reaction "02 + N = 20 + N";
     Α
           1.0e19;
            -1.5;
     beta
            59500:
     Ta
            (1e20 1e21 1e22 1e23 1e24 1e25); // local mixture number density in 1/m<sup>3</sup>
     ni
```

Please uncomment only the necessary species, here O_2 , N, and O. Please uncomment the appropriate chemical reactions, for instance Reaction no 1. The *type* of the reaction specifies whether the reaction is irreversible or reversible. A, beta, and Ta are the standard Arrhenius coefficients. The equilibrium constant is a function of the local number density of the mixture. The lists of coefficients A0 to A4 are linearly interpolated if the local mixture number density is within the range $[1 \times 10^{20}, 1 \times 10^{25}]$ m^{-3} . If not, the interpolation is bounded.

3.3.6 The *thermoDEM* dictionary

This dictionary provides the species thermodynamic data for the different species loaded in hTCReactionsDPLR. The dictionary structure is given here for the N_2 molecule.

```
FoamFile
                2.0;
    version
    format
                ascii:
                dictionary;
                "constant";
    location
    object
                thermoDEM:
N2
{
    specie
        nMoles
                        28.0134;
        molWeight
        particleType
                        2:
        charge
                        0;
                        4.17e-10;
        diameter
                        3.36e7:
        dissocEnergy
                        0.74;
        omega
        noVibTemp
                        1:
        noElecLevels
                        15:
    thermodynamics
        Tlow
                             200;
                             40000:
        Thigh
        decoupledCvCoeffs
                             (1.5 1 1 0 0 0 0 );
        vibrationalList
                             (1 3371);
        electronicList
                                3 7.223157e4
                                6 8.577863e4
                                6
                                  8.605027e4
                                3 9.535119e4
                                  9.805636e4
                                  9.968268e4
                                2 1.048976e5
                                5 1.116490e5
                                1
                                   1.225836e5
                                6
                                   1.248857e5
                                6 1.282476e5
                                10 1.338061e5
                                6 1.404296e5
```

```
1.504959e5
   }
   transport
   {
       constant
                     0:
          mu
       SutherlandEucken
                     1.41e-6;
                     111.0;
          Ts
       BlottnerEucken
       {
                     2.68e-2:
          В
                     3.18e-1;
          С
                    -1.13e1;
       }
       CEA
       {
          temp
               (200 1000 5000 15000);
          visco (
                  (0.62526577 -31.779652 -1640.7983 1.7454992)
                 (0.87395209 561.52222 -173948.09 -0.39335958)
                 (0.88503551 909.02171 -731290.61 -0.53503838)
               );
          kappa (
                 (0.85439436\ 105.73224\ -12347.848\ 0.47793128)
                 (0.88407146 133.57293 -11429.64
                 (2.4176185 8047.7749 3105580.2 -14.517761)
               );
       }
   }
}
```

Inside the specie sub-dictionary, new entries have been added. The particle Type gives the type of the particle: electron (0), atom (1), molecule (2), atom (3). The charge entry indicates the charge of particle: atom and molecule (0), ion (1), electron (-1). The diameter of the particle and its temperature coefficient of viscosity, omega, are also specified. The dissociation energy of the particle, named dissocEnergy, is given in particular for use in hy2Foam. The last two entries indicate the number of vibrational modes (noVibTemp) and electronic levels (noElecLevels) of the particle. Further detail about vibrational and electronic energy modes are then given into the thermodynamics sub-directory thanks to the vibrationalList and electronicList lists. These two lists contain a sequence of scalars grouped by pair: the degeneracy and the associated characteristic temperature for a given mode or level.

The following decoupledCvCoeffs list has seven elements. Their meaning is detailed in Table 3.

```
decoupledCvCoeffs ( 1.5 1 1 0 0 0 0 );
```

Element number	Meaning
1	translational coefficient
2	rotational coefficient
3	vibrational coefficient
4	electronic coefficient
5	electron coefficient
6	chemical enthalpy at 298.15 K
7	chemical entropy at 298.15 K (unused)

Table 3: The decoupledCvCoeffs list

Hence, the scalar number 4 should be set to 0 to deactivate the electronic energy mode in a simulation. The same can be repeated for the scalar number 3 to have vibrationless nitrogen.

The transport sub-dictionary specifies some constants depending on the transport model that has been chosen in the thermophysicalProperties dictionary. Using the constant transport model in combination with the following set-up in thermoDEM

```
constant
{
    mu    0;
}
```

allows the user to run inviscid calculations. As and Ts are the standard coefficients to determine the viscosity from the Sutherland's law while coefficients A, B, and C coefficients appear in Blottner's formula. CEA stands for the CEA2 (Chemical Equilibrium with Applications) library from NASA Glenn Research Center.

3.3.7 The transportProperties dictionary

This dictionary is composed of two sections. The first one concerns the quantities used to gauge the degree of rarefaction of a gas. The characteristic length of the case scenario should be entered here. There are two mean free path models (mfpModel) available: variableHardSphere and hardSphere. The rarefied parameters can be printed from this dictionary. KnGLL is the local gradient-length Knudsen number defined as the maximum of the three local gradient-length Knudsen numbers based on the density, velocity, and temperature fields. Turning the writeKnGLL components switch on and the three KnGLL numbers will be printed.

The second section of the transportProperties dictionary completes the information already given in thermophysicalProperties and thermoDEM about the different transport models.

```
FoamFile
                2.0;
    version
    format.
                ascii:
                dictionary;
    location
                "constant";
    object
                transportProperties;
rarefiedParameters
    computeFieldAndBoundaries
    computeMfpBoundaries
                                    true;
    mfpModel
                                    variableHardSphere;
    writeMfpSpecies
                                    off:
```

```
writeMfpMixture
                              off;
   characteristicLength
                              1e-5:
   writeKn_overall
                              off;
   writeKnGLL
                              off:
   writeKnGLL_components
                              off;
transportModels
   viscosityAndThermalConductivityModels
                                      "see thermophysicalProperties";
   mixingRule
                              molar;
   writeViscositySpecies
   writeViscosityMixture
                              off:
   {\tt write Thermal Conduc Species}
                              off;
   writeThermalConducMixture
                              off;
   writeHeatFluxes
                              off;
   writeWallHeatFlux
                              on;
   multiSpeciesTransport
                              LewisNumber;
   binaryDiffusivityModel
                              noBinaryDiffusivityModel;
```

To this date, there are three mixing rule implemented in hyFoam: a simple and generally unsuitable molar average using the keyword molar, and those of Wilke and ArmalySutton. The viscosity and thermal conductivity can be printed, either for each species or for the mixture. The transportModels sub-dictionary also prescribes the multi-species transport model used in hyFoam. The different possibilities are listed in Table 4.

multiSpeciesTransport entry	Meaning	
noSpeciesDiffusion	disables species diffusion	
Lewis Number	uses the Lewis number	
modified Lewis Number	see Gollan's PhD thesis	
Fick	generalisation of Fick's binary formula for a mixture	
binary Diffusivity Model entry	Meaning	
no Binary Diffusivity Model	no binary diffusivity model	
constant	constant binary diffusion coefficients (used with Fick only)	
Hirschfelder	Hirschfelder model (used with Fick only)	
Stephani	see K. A. Stephani's PhD thesis (used with Fick only)	
Gupta	available soon, see Gupta 1989 (used with Fick only)	

Table 4: Available multi-species transport models in hy2Foam

3.4 The *system* folder

A few modifications have been made to the system folder. They are located in fvSchemes and fvSolution. Local time stepping can be switched on by simply selecting the localEuler rDeltaT temporal scheme in place of the traditional Euler scheme. Three divergence scheme entries have been added.

```
referenceCase

VincePC: referenceCase >ls system/
controlDict decomposeParDict fvSchemes fvSolution
VincePC: referenceCase >
```

Figure 4: The **system** folder

4 New boundary conditions

4.1 Velocity field

$4.1.1 \quad rampInlet$

The time-dependent rampInlet boundary condition is aiming at reducing the very steep initial velocity gradients and thus falicitating the convergence of hypervelocity simulations. Its implementation is given hereafter

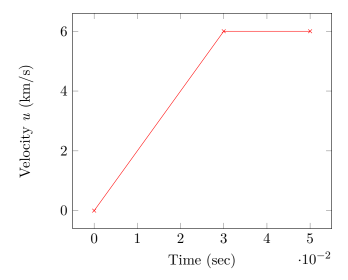
Using the rampInlet boundary type, the inlet velocity vector is defined as follow

$$U(t) = (u(t), v(t), w(t))^{T}$$

$$= \left(amplitude \times refValue - offset \odot refValue\right)$$

$$\times min(t/tRamp, 1) + offset \odot refValue$$
(1)

where \odot is the Hadamard product.



$4.1.2 \quad non Eq Max well Slip U$

The original maxwellSlipU velocity boundary condition implemented in rhoCentralFoam is modified and takes the following form in hy2Foam

```
wall
{
                            nonEqMaxwellSlipU;
        type
        Uwall
                            uniform (0 0 0);
                            uniform (0 0 0);
        refValue
        accommodationCoeff 1.0;
        valueFraction
                            uniform 1.0;
        thermalCreep
                            off;
        curvature
                            off;
        value
                            $Uwall;
}
```

4.2 Temperature field

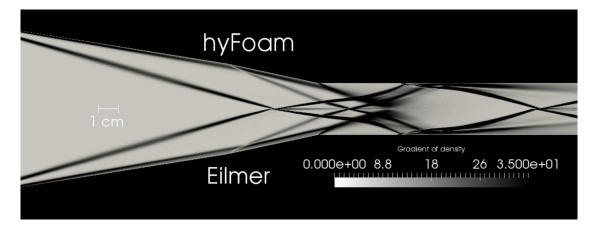
$4.2.1 \quad nonEqSmoluchowskiJumpT$

The original smoluchowskiJumpT temperature boundary condition implemented in rhoCentralFoam is modified and takes the following form in hyFoam

These formulations are using the overall free path boundary field.

5 Coming soon

A new section to describe the tutorials that have been uploaded.



Credit: J.-J. Hoste et al. 'Numerical Modeling and Simulation of Supersonic Flows in Propulsion Systems by Open-Source Solvers', AIAA Hypersonics, Xiamen, China, 6-9 March 2017)