MD - Bulk Evaluation

Program to evaluate MD output files of bulk simulations Notes:

- usable for LAMMPS simulations in metal units
- required files provided by LAMMPS scipt "in.output"

Manual

Required Folder structure:

- "STATE_1"
 - "Sim_001"
 - "Sim_002"
 - o ...
- "STATE_1"
 - "Sim_001"
 - "Sim_002"
 - o ...
- ...

Input File

Name: 'INPUT.txt' (has to be in same folder as 'Master' file)

Format:

• first line: '#' + 'keyword'

• second line: value assigned to keyword

Keywords:

Name	Туре	Description
folder	string	path to main folder containing all simulation data of one thermodynamic state (can occur multiple times)
ensemble	string ["NVT"	"NVE"
timesteps_EQU	interger (≥ 0)	number of timesteps to ignore at the start of each simulation
DO_evalulation	integer [0	1]
DO_state	integer [0	1]
N_boot	integer (≈ No. simulations)	number of bootstrapping repetitions

Example:

#folder

F:/MD_Bulk/Others/Methane/2020-04-16_trappe-ua/SIM_T_273.15K_rho_0.1gml #folder

```
F:/MD_Bulk/Others/Methane/2020-04-16_trappe-ua/SIM_T_273.15K_rho_0.3gml
#folder
F:/MD_Bulk/Others/Methane/2020-04-16_trappe-ua/SIM_T_273.15K_rho_0.5gml
#ensemble
NVT
#timesteps_EQU
0
#DO_evaluation
1
#DO_state
1
#N_boot
```

Output

Units:

• Length: [L] = m

• Mass: [m] = g/mol

• Time: [t] = s

• Temperature: [T] = K

• Pressure: [p] = MPa

• Density: $[\rho] = g/ml$

• Energy: [E] = eV

Viscosity: [η] = Pa*s

• Diffusion Coefficient: [D] = m^2/s

• Thermal Conductivity: $[\lambda] = W/(m^*K)$