## Documentation

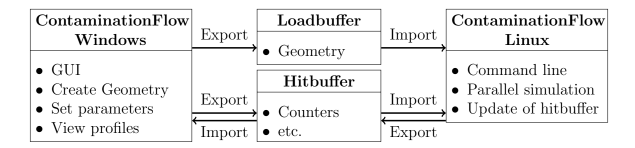
# ContaminationFlow on Linux and Windows

Hoai My Van, Rudolf Schönmann

# **Contents**

| 1. | Gen  | erai Structure                        | 1               |
|----|------|---------------------------------------|-----------------|
| 2. | Con  | taminationFlow Linux                  | 2               |
|    | 2.1. | Call of Application from Command line | 3               |
|    | 2.2. | Communication                         | 5               |
|    |      | Usage of boost Library                | 5               |
|    | 2.4. | New Quantities                        | 5               |
|    | 2.5. | Iterative Algorithm                   | 6               |
|    |      | 2.5.1. Initialization of simulation   | 6               |
|    |      | 2.5.2. Simulation on subprocesses     | 7               |
|    | 2.6. | Update main buffer                    | 9               |
|    | 2.7. | Summary                               | 10              |
| 3. | Con  | taminationFlow Windows                | 12              |
|    | 3.1. | Graphical User Interface              | 12              |
|    |      | -                                     | 12              |
|    | 3.3. | New Quantities                        | 12              |
|    |      | ·                                     | 13              |
| Α. | Forn | nulas for new Quantities              | 14              |
| В. | Data | atypes                                | 16              |
|    |      | ••                                    | 16              |
|    |      |                                       | 16              |
| c  | Ονο  | rview of new Classes and Functions    | 17              |
| С. |      |                                       | 17              |
|    |      |                                       | 19              |
|    | 0.2. |                                       | 19              |
|    |      | 11                                    | $\frac{10}{20}$ |
|    |      | 11                                    | $\frac{20}{20}$ |
|    |      | 11                                    | $\frac{20}{20}$ |
|    |      |                                       | $\frac{20}{21}$ |
|    |      | C.2.6. UpdateSubProcess.cpp           |                 |
|    |      |                                       | 22<br>22        |
|    |      | 0.2.1. ο ραωθοινιαπτι τουσορισρη      | 44              |

# 1. General Structure



General structure for ContaminationFlow simulation

- ullet Code adapted from Molflow
- $\bullet$  Contamination Flow Windows primary used to create Geometry and to view simulation results through the  ${\rm GUI}$
- ContaminationFlow Linux primary used for simulation and calculation of counters, profiles, etc.
- Loadbuffer contains information of geometry
- Hitbuffer contains information such as hit counters, profiles, etc.
- Import and export of buffer files for communication between ContaminationFlow Windows and ContaminationFlow Linux
- Export of simulationHistory for ContaminationFlow Linux

# 2. ContaminationFlow Linux

- Parallel simulation on several sub processes
- Processing and control of data in main process
- Update and accumulation of hit counters and other information such as profiles
- SimulationHistory, final hitbuffer and used parameters exported to results folder

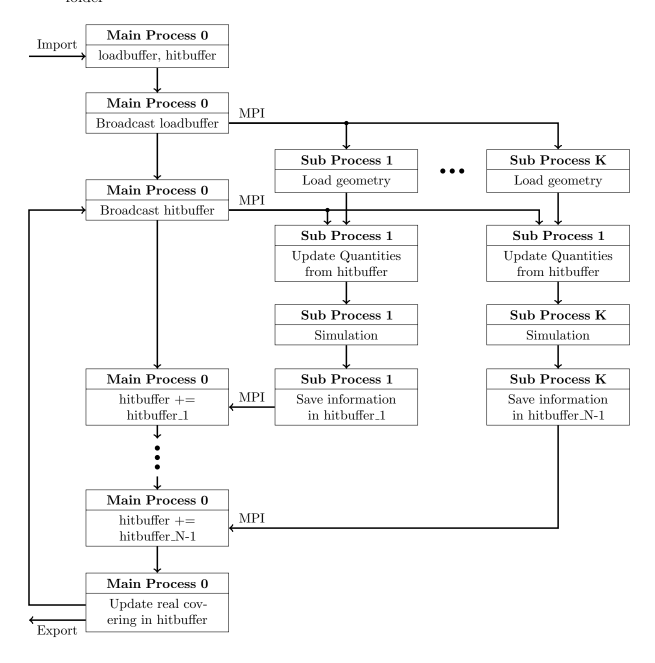


Figure 2.1.: Processing of data in main and sub processes

## 2.1. Call of Application from Command line

#### New class ProblemDef

- Defines parameters used for simulation
- Possible adaptation of default paramaters through input file
- Creates result folder for simulation if desired
  - Final resultbuffer
  - Final covering, input file and console output as text files

#### Application with custom parameters using input file

Call of ContaminationFlow Linux application in the command line:

```
$ module load mpi
$ mpirun -n N MolflowLinux inputfile save
```

with the following command line parameters:

- N: desired number of worker processes; simulation on K=N-1 worker processes
- MolflowLinux: path to application, e.g. ~/MolflowLinux/Debug/MolflowLinux
- inputfile: path to file that defines simulation parameters
- save: determines whether result directory is created (1: true, 0:false); default: 1 and the input file defining the following parameters:
  - loadbufferPath: path to loadbuffer file, contains geometry, e.g. ~/loadbuffer
  - hitbufferPath: path to hitbuffer file, contains counters, etc., e.g. ~/hitbuffer
  - simulationTime: simulation time per iteration step; default: 10.0
  - unit: simulation time unit; default: s
  - maxTime: maximum simulation time; default: 10.0
  - maxUnit: maximum simulation time unit; default: y
  - iterationNumber: number of iterations; default: 43200
  - particleDia: diameter of particles; default: 2.76E-12
  - $E_{de}$ : binding energy of a particle on pure substrate; default: 1E-21
  - $H_{vap}$ : vaporization enthalpy of a particle in case of multilayer contamination; default: 0.8E-19
  - $W_{tr}$ : transition width between monolayer and multilayer properties; default: 1
  - *sticking*: constant sticking coefficient for all facets, set to zero, not used at the moment; default: 0
  - targetPaticles: minimum number of desorbed particles per iter.; default: 1000
  - targetError: average statistical uncertainty (error) to be achieved for each iteration, calculated as the average (weighted with the facets area) of the normalized standard deviation of events per facet; default: 0.001
  - hitRatioLimit: Ratio at which hits are ignored, default: 1E-5
  - Tmin or  $t_{-}min$ : minimum time for step size; default: 1E-4
  - maxStepSize or t\_max: maximum time for step size; default: max
  - maxSimPerIt: maximum simulation steps per iteration; default: max
  - coveringMinThresh: minimum covering (through multiplication); default: 10000

- *histsize* : Size of history lists; default: max
- vipFacets: very important facets: facets with have their own target error. input in inputfile as alternating sequence of facet numbers and respective target errors seperated via blanks; default: []

#### **Terminology**

- Simulation time: desired computation time until check if target is reached for iteration
- Simulated time: physical time in the simulated system, e.g. flight time or residence time of a particle
- Maximum simulation time: desired total simulated time
- Step size: desired simulated time per particle for iteration

## 2.2. Communication

#### Import and export of buffer files

• New Databuff struct that replaces Dataport struct from MolFlow Windows

```
typedef unsigned char BYTE;
typedef struct {
  signed int size;
  BYTE *buff;
} Databuff;
```

- New functions importBuff(·) and exportBuff(·) for import of buffer files and export of Databuff struct
- New functions checkReadable(·) and checkWriteable(·) to check if file is readable or writeable

#### Communication between worker processes via MPI

- Main process 0 sends Databuff struct containing loadbuffer and Databuff struct containing hitbuffer and required simulationHistory values to sub processes using MPI\_Bcast(·)
- Sub processes send updated Databuff struct containing hitbuffer and required simulationHistory values to main process 0 using MPI\_Send(·) and MPI\_Recv(·)

## 2.3. Usage of boost Library

#### Multiprecision

- Increase precision for variables if required
- Avoid overflow for integer and underflow for floating point numbers

## 2.4. New Quantities

#### New counter covering

- Number of carbon equivalent particles on facet
- Increases with adsorption, decreases with desorption
- Extracted from new hitbuffer counter from Simulationcalc.cpp file in getCovering(·)

#### Coverage

- Number of monolayers of adsorbed particles
- Calculated from covering, particle diamieter (previously gas mass) and facet area
- Coverage computed from Simulationcalc.cpp file in calcCoverage(·)

#### Sticking factor

- Ratio adsorbed particles to impinging particles
- Set to 0, can be adapted for all facets through input file

#### Binding energy

- Calculated from  $E_{de}$ ,  $H_{vap}$  and  $W_{tr}$
- Energy computed from Simulationcalc.cpp file in calcEnergy(·)

#### **Desorption** [1/s]

- number of particles desorbing per second
- Calculated from binding energy, covering and temperature
- Desorption computed from Simulationcalc.cpp file in calcDesorption(·)

### **Desorption Rate [Pa m<sup>3</sup>/s]**

- Calculated from desorption, gas mass and facet area before each iteration
- Used to determine starting point for new particle
- Desorption rate computed from Simulationcalc.cpp file in calcDesorptionRate(·)

#### **Outgassing Rate**

- Calculated from facet outgassing and temerature defined in sHandle
- Outgassing rate computed from Worker.cpp file in CalcTotalOutgassingWorker()

#### K<sub>real/virtual</sub>

- Number of real particles represented by test particles
- Calculated from desorption & outgassing rate and number of desorbed molecules
- K<sub>real/virtual</sub> computed from Simulationcalc.cpp file in GetMoleculesPerTP(·)

#### Statistical Error

- Calculated from hits and desorbed particles (of facet and total)
- Used to determine significance of simulation results of iteration

## 2.5. Iterative Algorithm

#### 2.5.1. Initialization of simulation

#### New class to store Simulation History

• SimulationHistory class

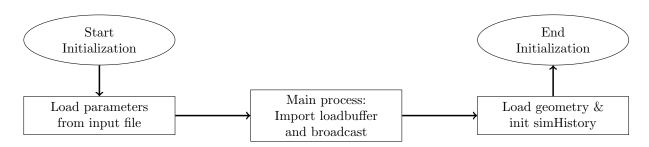


Figure 2.2.: Overview: Initialize simulation

```
class SimulationHistory {
public:
   SimulationHistory();
   SimulationHistory(Databuff *hitbuffer);

HistoryList<llong> coveringList;
HistoryList<llong> desorbedList;
HistoryList<double> hitList;
HistoryList<double> errorList;

double lastTime;
int currentStep;
};
```

```
template <typename T> class HistoryList {
public:
    HistoryList();
    std::vector<std::pair<double,std::vector<T>>> pointInTimeList;
    std::vector<T> currentList;
};
```

- In SimulationLinux.h and SimulationLinux.cpp file
- SimulationHistory updated after each iteration in UpdateCovering(·) from UpdateMainProcess.
- Currently recorded quantities: covering and error for each facet for each iteration, total hits and desorbed particles for each facet
- lastTime: simulated time (accumulated time steps) instead of computation time

## 2.5.2. Simulation on subprocesses

#### **Calculate Step Size**

- Use simHistory—currentStep to calculate logarithmic step size
- Calculation in UpdateMainProcess.cpp file in getStepSize()

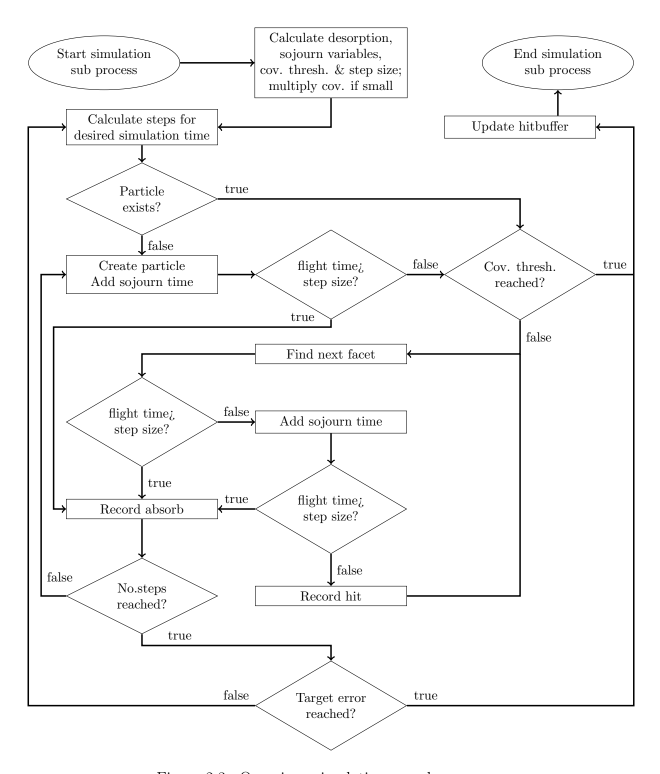


Figure 2.3.: Overview: simulation on sub processes

#### **Calculate Covering Threshold**

- Set lower threshold for covering for each facet to prevent covering getting negative
- Stop simulation once threshold is reached
- Threshold set in setCoveringThreshold(·) from Iteration.cpp file

#### Multiply small covering

- Multiply covering so that smallest covering  $\geq$  ProblemDef::coveringThreshMin
- Multiply covering threshold with same factor
- Calculation in checkSmallCovering(·) from SimulationLinux.cpp file

#### Calculate desorption

- Desorption rate calculated from current covering values
- Calculation in UpdateDesorptionRate(·) from UpdateSubProcess.cpp file

#### Calculate residence time and binding energy

- residence time of a particle on a surface calculated from binding energy, coverage and thermal oscillation frequency
- binding energy calculated using binding energy on pure substrate, vaporization enthalpy and coverage
- thermal oscillation frequency calculated using temperature, Boltzmann constant and Planck's constant
- Calculation in UpdateSojourn(·) from UpdateSubProcess.cpp file

#### Target error reached?

- Calculate statistical error in UpdateError() from UpdateSubProcess.cpp file
- Check if vip facets reached their own target error
- Check if normal facets total error reached target error
- Total error calculated from summing facet error weighted with facet area
- Set error of facets that reached ProblemDef::hitRatioLimit to inf
- Set error of facets with no hits and desorption to inf
- Facets with error=inf are not considered
  - if vip facet: target automatically reached
  - if normal facet: facet error and area not used for calculation
- Check in checkErrorSub(·) from UpdateSubProcess.cpp file

## 2.6. Update main buffer

#### Multiply small covering before summation of subprocesses

- Only if covering was multiplied in subprocesses
- Multiply covering in hitbuffer of main process before simulation using same factor

#### Divide small covering after summation of subprocesses

- Division by the small covering factor will be done when converting virtual test particles into real, physical particles
- Then, hitbuffer and hitbuffer\_sum of main process will be overwritten with the real, physical values anyway. So there is no need to divide by the small covering factor.

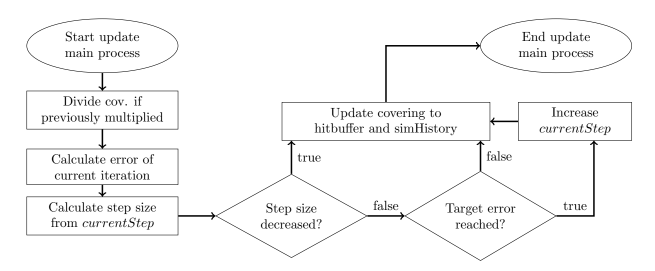


Figure 2.4.: Overview: update of covering in hitbuffer

#### **Error Calculation**

- Calculate statistical error of normal facets
- Facet error calculated from hits and desorbed particles, weighted with opacity
- Total error calculated from summing facet error weighted with facet area
- Set error of facets that reached ProblemDef::hitRatioLimit to inf
- Set error of facets with no hits and desorption to inf
- Facets with error=inf are not used for calculation of total error
  - if vip facet: target automatically reached
  - if normal facet: facet error and area not used for calculation
- Save error in simHistory→errorList
- Management in UpdateErrorMain(·) from UpdateMainProcess.cpp file

#### Management of Step Size

- Adapt step size if desorption × time step was larger than covering
- Increase simHistory -> currentStep if no adaptation & target errors reached
- Management in manageStepSize(·) from UpdateMainProcess.cpp file

#### **Update Covering**

- Use step size and  $K_{real/virt}$  to calculate new covering
- Calculation in UpdateCovering(⋅) from UpdateMainProcess.cpp file

## 2.7. Summary

#### **General Pipeline**

- Initialize MPI, ProblemDef p and SimulationHistory simHistory
- Load geometry into Simulation sHandle using LoadSimulation()
- Iteration until desired maximum simulation time is reached:
  - Reset hitbuffer counters using initbufftotero(·)

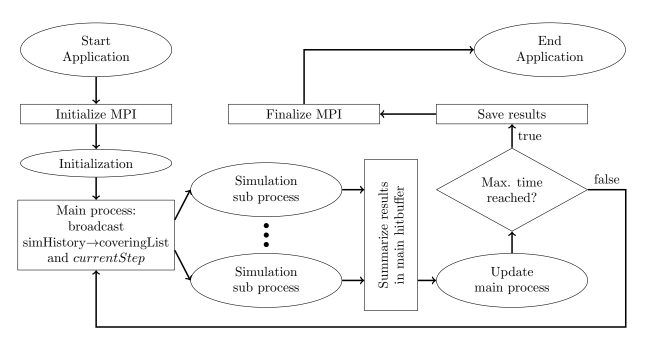


Figure 2.5.: Overview: ContaminationFlow application

- Broadcast simHistory→coveringList using MPI\_Bcast(·)
- Simulation in sub processes
  - Simulate until targetParticles and targetError or covthresh reached
  - Update hitbuffers of sub processes from sHandle using UpdateSubHits(·)
     from UpdateSubProcess.cpp
- Update Main process:
  - Send hitbuffer to main process using MPI\_Send(·) and MPI\_Recv(·)
  - Update of hitbuffer in UpdateMainHits(·) from UpdateMainProcess.cpp
- Update error of iteration using UpdateErrorMain(·) from UpdateMainProcess.cpp
- Calculate real covering in main process using  $K_{real/virtual}$  and simulated step size in UpdateCovering(·) from UpdateMainProcess.cpp, save in simHistory
- Update real covering in hitbuffer of main process in UpdateCoveringphys(·) from UpdateMainProcess.cpp
- Export final results (hitbuffer and simulationHistory) to results folder
- Close MPI

# 3. ContaminationFlow Windows

• Create Geometry and set parameters such as initial coverage and temperature

## 3.1. Graphical User Interface

Add screenshot of GUI

#### **New GUI elements**

- "Particles out" renamed to Contamination level
  - Text field for covering
  - Text field for coverage
- New facet properties
  - Effective surface factor
  - Facet depth and facet volume
  - Diffusion coefficient
  - Concentration and gas mass
- Window for CoveringHistory (reworked to SimulationHistory in ContaminationFlow Linux)
- PressureEvolution window expanded
  - Added list that contains information of graph
  - Option to show only selected facets or all
  - List exportable

### 3.2. Communication

#### Import and export of buffer files via GUI

• New Databuff struct

```
typedef unsigned char BYTE;
typedef struct
  signed int size;
  BYTE *buff;
Databuff;
```

- New functions importBuff(·) and exportBuff(·) for import and export of buffer files/Databuff struct
- New options in file menu: Export buffer and Import buffer

## 3.3. New Quantities

#### New counter covering

• Covering computed in SimulationMC.cpp file in updatecovering(·)

- Added covering counter to hitbuffer
- Added covering to GUI, can be defined through textfield

#### New facet property effetiveSurfaceFactor

• Defines increase of facet area due to texture

#### New facet property facetDepth

• Defines depth of facet

#### New facet property diffusionCoefficient

• Defines diffusion coefficient

#### New facet property concentration

• Defines concentration = mass of particles in volume

#### Removal of irrelevant quantities

- Sticking factor and pumping speed removed from GUI
- calcSticking() and calcFlow() in Molflow.cpp file not used anymore
- Flow not needed for iterative Algorithm

## 3.4. Iterative algorithm

#### New class to store covering for all facets at any time

- In HistoryWin.cpp and HistoryWin.h file
- std::vector<std::pair<double,std::vector<double>>> pointintime\_list to store points in time and respective covering for all facets
- New GUI option to add and remove entries for pointintime\_list
- New GUI option to export or import a complete list

# A. Formulas for new Quantities

#### **Constants**

$$carbondiameter = 2 \cdot 76E - 12$$

$$K_b = 1.38E - 23$$

$$h = 6.626E - 34$$
(A.1)

Number of carbon equivalent particles of one monolayer

$$N_{mono} = \frac{\text{Area of Facet [m^2]}}{carbondiameter^2} \tag{A.2}$$

Carbon equivalent relative mass factor

$$\Delta N_{surf} = \frac{\text{carbon equivalent gas mass}}{12.011} \tag{A.3}$$

Covering  $\theta^*$ 

$$\theta^* = N_{\text{particles on facet}}$$
 (A.4)

Coverage  $\theta$ 

$$\theta = \frac{\theta^*}{N_{mono}/\Delta N_{surf}} \tag{A.5}$$

**Step function**  $step(x, y_{start}, y_{end}, x_{turningpoint}, w)$ 

$$E = \frac{y_{start} - y_{end}}{2} \cdot \tanh\left(\left(x_{turningpoint} - x\right) \cdot \frac{5.4}{w}\right) + \frac{y_{start} + y_{end}}{2}$$
 (A.6)

Binding Energy E

$$E = step(\theta, E_{de}, H_{vap}, 1, W_{tr})$$

$$= \frac{E_{de} - H_{vap}}{2} \cdot \tanh\left((1 - \theta) \cdot \frac{5.4}{W_{tr}}\right) + \frac{E_{de} + H_{vap}}{2}$$
(A.7)

Residence

$$Frequency = \frac{K_b T}{h}$$

$$Energy = E$$
(A.8)

Residence Time

$$A = \exp\left(-\frac{Energy}{(k_bT)}\right)$$

$$residence \ time = \frac{-\ln(rnd)}{A \cdot Frequency}$$
(A.9)

#### **Desorption rate** des

$$\tau = \frac{h}{K_b T}$$

$$d = step(\theta, 1, 0, 1, W_{tr})$$

$$des = \begin{cases} \frac{1}{\tau} \theta^d \exp\left(-\frac{E}{K_b T}\right) \cdot \frac{N_{mono}}{\Delta N_{surf}} \cdot K_b T, & \text{if } \theta^* > 0\\ 0, & \text{otherwise} \end{cases}$$
(A.10)

#### Outgassing rate out

$$out = \frac{\text{Facet outgassing}}{K_b T} \tag{A.11}$$

K<sub>real/virtual</sub>

$$K_{\text{real/virtual}} = \frac{\sum_{\text{facets}} \left( out + \frac{des}{K_b T} \right)}{\text{number of total desorbed molecules}}$$
(A.12)

#### Step Size $T_{step}$

$$T_{min} = \text{Tmin}$$

$$T_i = T_{min} \cdot \exp\left(i \cdot \ln(\text{max. simulation time}/T_{min})/\text{max. } \# \text{ of steps}\right)$$

$$T_{step} = T_{currentStep+1} - T_{currentStep}$$
(A.13)

#### **Error**

$$error = \begin{cases} inf & \text{, if (hits+desorbed) on facet } = 0 \\ \left(\frac{1}{\text{(hits+desorbed) on facet}} \cdot \frac{1 - \text{(hits+desorbed) on facet}}{\text{total (hits+desorbed)}}\right)^{0.5} & \text{, else} \end{cases}$$
(A.14)

# **B.** Datatypes

# **B.1. Class Members**

| Name                            | Datatype                         | Alias    |
|---------------------------------|----------------------------------|----------|
| SimulationHistory::coveringList | boost::multiprecision::uint_128t | covBoost |
| FacetHitBuffer::covering        | llong                            | covLlong |
| FacetProperties::desorption     | boost::multiprecision::float128  | desBoost |
| Simulation::coveringThreshold   | llong                            |          |

# **B.2. Functions**

| Function              | Output Datatype                          | Relevant Input   |
|-----------------------|--|------------------|
| getCovering()         | boost::multiprecision::float128          | covBoost         |
| getCovering()         | llong                                    | covLlong         |
| calcCoverage()        | boost::multiprecision::float128 or llong | getCovering()    |
| calcDesorption()      | boost::multiprecision::float128          | calcCoverage()   |
| calcDesorptionRate()  | boost::multiprecision::float128          | calcDesorption() |
| calctotalDesorption() | boost::multiprecision::float128          | desBoost         |
| GetMoleculesPerTP()   | boost::multiprecision::float128          | desBoost         |

# C. Overview of new Classes and Functions

## C.1. New Classes

| SimulationHistory |   |  |
|-------------------|---|--|
| coveringList      | of class HistoryList, stores covering history   |  |
| errorList         | of class HistoryList, stores error history  |  |
| hitList           | of class HistoryList, stores hits for each facet  |  |
| desorbedList      | of class HistoryList, stores desorbed particles for each facet  |  |
| startNewParticle  | Determines wether to create a new particle for next iteration   |  |
| numFacet          | number of Facets  |  |
| numSubProcess     | number of sub processes used for simulation   |  |
| nbDesorbed_old    | number of total desorbed molecules of previous iteration $\Rightarrow$ To calculate difference between consecutive iterations |  |
| flightTime        | Simulated flight time for iteration   |  |
| nParticles        | Simulated particles for iteration   |  |
| lastTime          | Total simulated time = last time in Lists   |  |
| currentStep       | step of logarithmic time step calculation in getStepSize()  |  |
| stepSize          | current step size   |  |
| updateHistory()   | Reset and update from hitbuffer   |  |
| appendList()      | Updates coveringList from hitbuffer   |  |
| print()           | Print to terminal   |  |
| write()           | Write to file   |  |

| HistoryList          |  |  |
|----------------------|--|--|
| pointInTimeList      | list containing history respective facet values          |  |
| currentList          | list containing facet values at current step             |  |
| currIt               | current iteration number                                 |  |
| appendCurrent()      | Appends currentList to pointInTimeList                   |  |
| appendList()         | Append input list to pointInTimeList                     |  |
| convertTime()        | Converts time for better clarity                         |  |
| printCurrent()       | Print currentList as table to terminal, optional message |  |
| print()              | Print pointInTimeList as table to terminal, optional msg |  |
| write(), read()      | Write to file, read from file                            |  |
| set/getCurrent()     | Set/get value of desired facet in currentList            |  |
| setLast(), getLast() | Set/get value of desired facet from pointInTimeList      |  |

| ProblemDef  |   |
|---|---|
| resultpath  | Path of result folder                                       |
| outFile   | Path of file that contains terminal output                  |
| loadbufferPath  | Path of loadbuffer file                                     |
| hitbufferPath   | Path of hitbuffer file                                      |
| $simulationTime$ , unit $\Rightarrow simulationTimeMS$                                | Computation time of each iteration in milliseconds          |
| $\begin{array}{c} \text{maxTime, maxUnit} \\ \Rightarrow \text{maxTimeS} \end{array}$ | Maximal total simulated time in seconds                     |
| iterationNumber   | Number of iterations  |
| particleDia   | Diameter of particles                                       |
| E_de, H_vap, W_tr   | Parameters to calculate binding energy, see equation A.7    |
| sticking  | Sticking factor for all facets                              |
| targetParticles/-Error  | Target values for each iteration                            |
| hitRatioLimit   | threshold of hitratio at which hits are ignored             |
| coveringMinThresh   | Minimum covering, multiplication to this if covering low    |
| Tmin, maxStepSize   | Minimum/ Maximum step size                                  |
| maxSimPerIt   | Maximun simulation steps per iteration                      |
| histSize  | Size of history lists (most recent values in memory)        |
| vipFacets   | alterning: vip facet and target error, e.g. 1 0.001 3 0.002 |
| readInputfile()   | Initialization from input file                              |
| printInputfile()  | Print to terminal   |

# C.2. New Functions

# C.2.1. molflowlinux\_main.cpp

| Preprocessing        |  |
|----------------------|--|
| parametercheck()     | Checks validity of input parameters from input file Defines values for ProblemDef object p |
| importBuff()         | Import load- and hitbuffer to main process   |
| MPI_Bcast()          | Send loadbuffer to sub processes   |
| LoadSimulation()     | Load geometry from loadbuffer  |
| initCoveringThresh() | Initialize covering threshold  |
| simHistory           | Initialize SimulationHistory object  |

| Simulation Loop   |  |  |
|---|--|--|
| initbufftozero()  | Reset all hitbuffer counters except covering                             |  |
| MPI_Bcast()   | Send simHistory—coveringList and simHistory—currentStep to sub processes |  |
| setCoveringThreshold()  | Sets covering threshold for each facet                                   |  |
| UpdateSojourn()   | Sets sojourn variables for each facet                                    |  |
| UpdateDesorptionRate()  | Sets desorption for each facet, ends simulation if 0                     |  |
| simulateSub()   | Simulation on sub processes  |  |
| MPI_Send(), MPI_Recv()  | Send sub hitbuffer to main process                                       |  |
| UpdateMCMainHits()  | Add simulation results from sub hitbuffer to main hitbuffer              |  |
| UndoSmallCovering()   | Divide covering if it was multiplied before                              |  |
| UpdateErrorMain()   | Calculate and save error of iteration to simHistory                      |  |
| UpdateCovering()  | Calculate and save new covering to simHistory                            |  |
| UpdateCoveringphys()  | Saves current covering to hitbuffer                                      |  |
| $	ext{simHistory} {	o} 	ext{coveringList} $ $	ext{simHistory} {	o} 	ext{errorList}$ | Adapt size to p→histSize if necessary                                    |  |
| End simulation if maximum simulation time is reached                                |  |  |

| Postprocessing     |                           |
|--------------------|---------------------------|
| exportBuff()       | Export final hitbuffer    |
| simHistory→write() | Export simulation history |

# C.2.2. SimulationLinux.cpp

| simulateSub()                             |   |  |
|---|---|--|
| targetParticles, targetError              | Calculate target values from overall target and number sub processes                  |  |
| <pre>simHistory-&gt;updateHistory()</pre> | Reset and Update SimulationHistory object from hitbuffer                              |  |
| smallCovering, smallCoveringFactor        | If smallCovering: Covering is multiplied by smallCoveringFactor to improve statistics |  |
| SimulationRun()                           | Simulate for desired simulation time  |  |
| UpdateError()                             | Calculate current error of sub process  |  |
| CheckErrorSub()                           | Checks if normal facets reached targetError and if vip facets reached own target      |  |
| UpdateMCSubHits()                         | Save simulation results to hitbuffer  |  |

| Small covering       |   |
|----------------------|---|
| CheckSmallCovering() | If smallCovering, find smallCoveringFactor to reach p→coveringMinThresh |
| UndoSmallCovering()  | If smallCovering, divide by previously determined smallCoveringFactor   |

# C.2.3. Iteration.cpp

| Set Covering Threshold to avoid negative covering |   |
|---|---|
| init Covering Thresh()                            | Initializes size of covering threshold vector |
| setCoveringThreshold()                            | Sets covering threshold for each facet        |

# C.2.4. Buffer.cpp

| Buffer functions  |  |
|-------------------|--|
| Databuff struct() | signed int size BYTE *buff                           |
| checkReadable()   | Checks if file can be opened for reading             |
| checkWriteable()  | Checks if file can be openend or created for writing |
| importBuff()      | Imports buffer file to Databuff struct               |
| exportBuff()      | Exports Databuff struct to buffer file               |

# C.2.5. Calculations in SimulationCalc.cpp etc.

| SimulationCalc.cpp                     |   |
|--|---|
| getCovering()                          | Get covering from hitbuffer                           |
| getHits()                              | Get number of hits from hitbuffer                     |
| getnbDesorbed()                        | Get number of total desorbed molecules from hitbuffer |
| calcNmono()                            | see equation A.2                                      |
| calcdNsurf()                           | see equation A.3                                      |
| calcCoverage()                         | see equation A.5                                      |
| calcStep()                             | see equation A.6                                      |
| calcEnergy()                           | see equation A.7                                      |
| calcStickingnew()                      | sets sticking coefficient to p—sticking               |
| calcDesorption(), calcDesorptionRate() | see equation A.10                                     |
| GetMoleculesPerTP()                    | see equation A.12                                     |
| calctotalDesorption                    | calculates desorption for startFromSource()           |
| calcPressure(), calcParticleDensity()  | TODO has to be verified                               |

| worker.cpp                    |  |
|-------------------------------|--|
| CalcTotalOutgassingWorker()   | see equation A.11, calculates outgassing |
| Calc rotaroutgassing worker() | for startFromSource()                    |

| SimulationLinux.cpp |  |
|---------------------|--|
| convertunit()       | Converts simutime*unit to milliseconds |

# C.2.6. UpdateSubProcess.cpp

| Update sHandle paramters from hitbuffer |                                      |
|---|--------------------------------------|
| UpdateSticking()                        | Updates sticking                     |
| UpdateDesorptionRate()                  | Updates desorption rate              |
| UpdateSojourn()                         | Updates sojourn frequency and energy |

| Error calculations |   |
|--------------------|---|
| UpdateErrorSub()   | Updates error for current iteration, see equation A.14  |
| UpdateError()      | Sums up error of normal facets & weights by facet area  |
| CheckErrorSub()    | Checks if normal & vip facets reached respective target |

| Update hitbuffer  |  |
|-------------------|--|
| initbufftozero()  | Sets hitbuffer except covering to zero               |
| UpdateMCSubHits() | Saves simulation results from sHandle into hitbuffer |

# C.2.7. UpdateMainProcess.cpp

| Update main hitbuffer from sub hitbuffer |   |
|--|---|
| UpdateMCMainHits()                       | Add simulation results from sub hitbuffer to main hitbuffer |

| Update real covering in hitbuffer |   |
|-----------------------------------|---|
| getStepSize()                     | Calculates step size for current step, see equation A.13                                |
| ${\rm manageStepSize}()$          | Adapts step size if desRate $\cdot$ step size $>$ than covering                         |
| UpdateCovering()                  | Uses step size and Krealvirt to calculate new covering Saved to simHistory—coveringList |
| UpdateCoveringphys()              | Saves current real covering to hitbuffer  |
| UpdateErrorMain()                 | Calculates total error, see equation A.14 Saved to simHistory→errorList                 |
| CalcPerIteration()                | Calculates total error and covering over all facets per iteration                       |