Quadrature-based moment methods for the simulation of turbulent polydisperse multiphase systems

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Quadrature-based moment methods
for the simulation of turbulent polydisperse multiphase systems

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This presentation concerns the simulation of multiphase polydisperse systems with Eulerian methods. Disperse means that there is a continuous phase and one (or more) additional phases constituted by separate elements: particles, droplets or bubbles. The elements of the disperse phase are characterized by some properties, often referred to as internal coordinates, such as particle size, volume, momentum, mass, enthalpy. The system is polydisperse because each particle is characterized by a different value of these properties, resulting in distributions. Particle size distribution (PSD), bubble size distribution (BSD), crystal size distribution (CSD), particle velocity distribution (PVD).
Historically two approaches have been used:

- Focusing on the PSD and neglecting spatial inhomogeneities;
  population balance equation:

  \[
  \frac{\partial n(L)}{\partial t} + \frac{\partial}{\partial L} \left( G(L) n(L) \right) = S(L)
  \]

- Focusing on spatial inhomogeneities and neglecting the PSD;
  two-fluid model (all particles have size \( L = d_p \)):

  \[
  \frac{\partial \alpha_p}{\partial t} + \frac{\partial}{\partial x} \cdot (\alpha_p U_p) = 0
  \]

  \[
  \frac{\partial \alpha_p U_p}{\partial t} + \frac{\partial}{\partial x} \cdot (\alpha_p U_p U_p) = -\frac{\alpha_p}{\rho_p} \frac{\partial p_f}{\partial x} + \alpha_p g
  \]

  \[
  + \frac{3}{4} \rho_f C_D(d_p) |U_f - U_p| (U_f - U_p)
  \]
Introduction

- The PSD is related to $\alpha_p$ through the **moment transform**:

$$\alpha_p = k_V \int_0^\infty n(L)L^3 \, dL = k_V m_3$$

- In general, the moment transform of order $k$:

$$m_k = \int_0^\infty n(L)L^k \, dL$$

  can be used to define global properties from the PSD

- **Microscale model**: defined at the particle level
- **Mesoscale model**: defined at the population level
- **Macroscale model**: defined on the global properties of the population

- The PBE is a mesoscale model (defined through an **ensemble-average** from the microscale model), whereas the two-fluid model is a macroscale model (usually derived through **volume-average** directly from the microscale model)
Introduction

Quadrature-based moment methods

D. L. Marchisio

GPBE
Velocity disperse phase
Solution methods
Examples
Moment corruption
Spatial discretization in DQMOM
Conclusions

Microscale model
Fully resolved direct numerical simulation

Mesoscale model
Population balance equation
Boltzmann equation
Kinetic equation
(Lagrangian methods)

Macroscale model
Moment equations
Mixture model
Two-fluid model
(Eulerian methods)
Outline

1. GPBE
2. Velocity disperse phase
3. Solution methods
4. Examples
5. Moment corruption
6. Spatial discretization in DQMOM
7. Conclusions
GENERALIZED POPULATION BALANCE EQUATION
Generalized Population Balance Equation

- Independent variables are classified as external (space) and internal (velocity, size, composition) coordinates.
- In general other internal coordinates can be used and here particle velocity, size and composition are just an example.
- First we define a Number Density Function (NDF) so that:

\[ n^*(t, \mathbf{x}, \mathbf{v}_p, L, \phi_p) \, d\mathbf{x} \, d\mathbf{v}_p \, dL \, d\phi_p \]

represents the expected number of particles, droplets or bubbles, in the infinitesimal volume \( d\mathbf{x} = dx_1 \, dx_2 \, dx_3 \) around the physical point \( \mathbf{x} = (x_1, x_2, x_3) \) with velocity in between \( \mathbf{v}_p \) and \( \mathbf{v}_p + d\mathbf{v}_p \), size \( L \) and \( L + dL \), and composition (number of moles of a specific component) \( \phi_p \) and \( \phi_p + d\phi_p \).
The evolution of the NDF is dictated by the Generalized Population Balance Equation (GPBE):

$$\frac{\partial n^*}{\partial t} + \frac{\partial}{\partial x} \cdot (v_p n^*) + \frac{\partial}{\partial L} (\langle G \rangle n^*)$$

$$+ \frac{\partial}{\partial v_p} \cdot (\langle A_p \rangle n^*) + \frac{\partial}{\partial \phi_p} \cdot (\langle \dot{\phi}_p \rangle n^*) = h(t, x, L, v_p, \phi_p)$$

- $\langle G \rangle$ is the continuous growth rate, $\langle A_p \rangle$ is force per unit mass (i.e., continuous acceleration) acting on the particles, $\langle \dot{\phi}_p \rangle$ is the continuous rate of change of composition and $h$ is the discontinuous term accounting for the instantaneous changes in size, composition and momentum due to discrete events (such as collision, aggregation, breakage)

- $\langle G \rangle$, $\langle A_p \rangle$, $\langle \phi_p \rangle$, and $h$ are conditional phase-space velocities or mesoscale models for the population of particles
Example: drag force for bubbly flow

- Acceleration acting on one single spherical bubble:

\[
A_p = -\frac{1}{\rho_p} \frac{\partial p_f}{\partial x} + \frac{3}{4} \frac{C_D}{L} \frac{A_D}{V_p} \frac{\rho_f}{\rho_p} |U_f - U_p|(U_f - U_p)
\]

where for a contaminated system:

\[
C_D = \max \left[ \frac{24}{\text{Re}_p} (1 + 0.15\text{Re}_p^{0.687}), \frac{8}{3} \left( \frac{E_0}{E_0 + 4} \right) \right]
\]

- To calculate \( \langle A_p \rangle \) one needs to account for bubble interaction and microscale turbulence (\( \text{Re}_{\text{eff}} = L \rho_f |U_f - U_p|/\mu_{\text{eff}} \)):

\[
C_D = (1 - \alpha_p) C_1 \max \left[ \frac{24}{\text{Re}_{\text{eff}}} (1 + 0.15\text{Re}_{\text{eff}}^{0.687}), \frac{8}{3} \frac{E_0}{E_0 + 4} \right]
\]
VELOCITY OF THE DISPERSE PHASE
The problem of the velocity of the dispersed phase

- Particle velocity is an important internal coordinate since it is related to particle position (external coordinate) through Newton’s law!
- It can be included in the set of internal coordinates, resulting in mixed particle velocity moments\(^1\)
- It can be integrated out from the GPBE:

\[
n(t, x, L, \phi_p) = \int\int\int_{-\infty}^{+\infty} n^*(t, x, v_p, L, \phi_p) \, dv_p
\]

resulting in the following equation:

\[
\frac{\partial n}{\partial t} + \frac{\partial}{\partial x} \left( \langle U_p|L, \phi_p \rangle n \right) + \frac{\partial}{\partial L} \left( \langle G \rangle n \right) + \frac{\partial}{\partial \phi_p} \left( \langle \dot{\phi}_b \rangle n \right) = h
\]

- The conditional particle velocity has to be calculated from equations derived from the GPBE with the moment transform!

The problem of the velocity of the dispersed phase

There are three key parameters: particle loading, $\rho_p/\rho_f$, particle Stokes number, $St$, and polydispersity, PDI

1. **Diffusion equation**: sub-micron particles / Brownian motion

2. **Pseudo-homogeneous or dusty-gas model**: very small $St$ and limited PDI

3. **Algebraic Eulerian model with a single conditional velocity based on the mean particle size**: small $St$ and limited PDI

4. **Algebraic Eulerian model with conditional velocities**: small $St$ and non-negligible PDI

5. **Eulerian two-fluid model with particle-phase velocity based on the mean particle size**: moderate/large $St$ and limited PDI (in both dilute and dense systems)

6. **Eulerian multi-fluid model**: moderate/large $St$ and large PDI (in both dilute and dense systems)
SOLUTION METHODS
The mesoscale model is highly dimensional (1 time + 3 spatial coordinates + 1 size + 3 velocities + ...)

One interesting way to solve it is to resort to Lagrangian / Direct Simulation Monte Carlo (DSMC)

Equations for moments of the NDF (Eulerian / macroscale model): Quadrature Method of Moments (QMOM)

Inversion algorithms for univariate problems (only size): Product-Difference or Wheeler algorithms

Inversion algorithms for multivariate problems (size and composition): Brute force methods, tensor-product methods, Conditional QMOM (CQMOM)

The quadrature approximation can also be tracked directly by preserving an optimal moment set: Direct Quadrature Method of Moments (DQMOM)
Comparison DSMC with CQMOM/DQMOM

DQMOM $N = 3$

![Graph showing comparison between DSMC and CQMOM/DQMOM with data points highlighting bubble size $L$ vs. oxygen moles $\phi_b$.]

- Quadrature-based moment methods
- D. L. Marchisio
- GPBE
- Velocity disperse phase
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- Spatial discretization in DQMOM
- Conclusions
Comparison DSMC with CQMOM/DQMOM

CQMOM $N_1 = 3 \quad N_2 = 1$

Bubble size $L$

Oxygen Moles $\phi_b$

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Quadrature-based moment methods

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GPBE

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Comparison DSMC with CQMOM/DQMOM

CQMOM $N_1 = 3 \quad N_2 = 2$

Bubble size $L$

Oxygen Moles $\phi_b$

Examples

Solution methods

Moments corruption

Spatial discretization in DQMOM

Conclusions
Coalescence, breakage and mass transfer. Solid black line: DSMC method. Red line CQMOM with $N_1 = 3$ and $N_2 = 1$ and DQMOM with $N = 3$. 
EXAMPLES: SIMULATION PARTICLE LADEN FLOWS, FLUIDIZATION, GAS-LIQUID STIRRED TANKS AND BUBBLE COLUMNS
• In turbulent particle-laden flows a phenomenon called \textbf{turbophoresis} pushes particles towards the walls

• This is typically described with DNS simulations with Lagrangian particle tracking to represent polydispersity

• The GPBE (in the absence of aggregation, breakage or growth) for particle size only (\(\rightarrow\) after filtering) becomes:

\[
\frac{\partial n}{\partial t} + \frac{\partial}{\partial x} \cdot (U_P(L)n) = 0 \quad \rightarrow \quad \frac{\partial \overline{n}}{\partial t} + \frac{\partial}{\partial x} \cdot (U_P(L)\overline{n}) = 0
\]

• The evaluation of the advective term can be done with an \textbf{approximate deconvolution method (ADM)}, \(U_P(L)\) is calculated with the \textbf{algebraic slip model (ASM)} whereas polydispersity is accounted for with \textbf{DQMOM}
DQMOM-LES for particle-laden flow

DQMOM implementation in TransAT (ASCOMP) for particle-laden flow in channel with per. bound. cond.

Comparison between Lagrangian DNS (black) and LES/ASM/ADM with $k = 0$ (red), $k = 1$ (purple), $k = 5$ (green), finer grid $k = 0$ (blue), finer grid GR $k = 5$ (brown)

St = 0.2  St = 1.0

Contour plots of the mean particle size $d_{32}$ and PSD in the center of the channel and near the wall (DQMOM approximations, red line, and reconstructed PSD, blue line).
In fluidized suspensions particles are segregated (or mixed) according to their size: **flotsam** and **jetsam**

This can be described by using QMOM ($N = 2$) for size coupled with a multi-fluid model (in Fluent) with one continuous phase (gas) plus two disperse phases (nodes 1 and 2)\(^2\)

The resulting transport equation for the moment of order $k$ (again without aggregation nor breakage) is:

$$\frac{\partial M_k}{\partial t} + \frac{\partial}{\partial x} \cdot (U^k_p M_k) = 0$$

The velocity for the moment of order $k$ is defined (with two nodes) as follows:

$$U^k_p = \frac{w_1 U_p(L_1) L^k_1 + w_2 U_p(L_2) L^k_2}{M_k}$$

QMOM/multi-fluid for fluidization

$M_0$ (top) and $M_1$ (bottom) from 5, 10 and 15 cm/s (left to right)
Top and bottom layers at 5 cm/s (left) and 15 cm/s (right)
CQMOM for a stirred tank

Standard rushton turbine / gassing rate 0.018 - 0.093vvm / stirring rate 155 - 250 rpm / reactor volume ≈ 180 l resulting in hold-up values up to 1.5 % / Fluent

CQMOM for a stirred tank

Results and comparison with experimental data

Reactor configuration 1
N = 250 RPM; 0.093 vvm

Mean bubble size, mm

BSD

d, mm

R2
R4
R8
R12
R9

Experimental data
Simulation results

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QMOM/CQMOM for a bubble column

Implementation of QMOM/CQMOM in openFOAM (compressibleTwoPhaseEulerFoam v 2.1)

Simulation of the bubble column investigated by Diaz et al., 2008. Chem. Eng. J., 139, 363-379
QMOM/CQMOM for a bubble column

Color mean bubble size
Implementation of QMOM/CQMOM in openFOAM
QMOM/CQMOM for a bubble column

Mean bubble size (m)
2.4 mm/s  11.9 mm/s  21.3 mm/s
QMOM/CQMOM for a bubble column

Sauter diameter $d_{32}$ (mm)

Superficial velocity (mm/s)

- ○ Experimental data
- ▲ Simulation
- ■ Inlet
QMOM/CQMOM for a bubble column

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- Simulation
- Experiment

Oxygen saturation (%)

Time (s)
ISSUES RELATED TO MOMENT CORRUPTION: QMOM/CQMOM
As we previously mentioned, given a distribution $n(L)$ the moments $M_k$ represent physical measurable quantities. For example, $M_0$ represents the total number of particles per unit volume. The first-order moment is related to the mean particle size: $M_1 / M_0$. The second order moment is related to the variance: $M_2 - M_1^2 / M_0$. The variance must be positive, therefore $M_2 \geq M_1^2 / M_0$. Since the moments are linked, they must respect some constraints, which are intuitive for lower-order moments and more exotic for higher-order moments.
Necessary validity condition: convexity of function \( \ln(m_k) \) with respect to \( k \):

\[
m_k m_{k-2} - m_{k-1}^2 \geq 0
\]
Spatial discretization in QMOM

- A moment set can be corrupted by time and space discretization schemes:

\[
\frac{\partial M_k}{\partial t} + U_p \frac{\partial M_k}{\partial x} = \overline{S}_k \rightarrow \frac{dM^P_k}{dt} = \overline{S}^P_k - \frac{U_p}{\Delta x} (M^e_k - M^w_k)
\]

- With first-order upwind $M^e_k = M^P_k$ and $M^w_k = M^W_k$

- Spatial discretization schemes based on first-order upwind always result in VALID moments. Higher-order schemes (second-order upwind, QUICK, MUSCL) always result in INVALID moments.
CORRECTION ALGORITHM BY MCGRAW

1. Build difference table and check if $d_2$ is negative
2. Identify moment order $k$ that causes the biggest change in $d_3$
3. Change moment (by multiplying it for a constant) in order to MINIMIZE $d_3$
4. Go back to point 1

CORRECTION ALGORITHM BY WRIGHT

1. Build difference table and check if $d_2$ is negative
2. Replace moments with those of a log-normal distribution with
   \[ \mu = \frac{j}{ij-i^2} \ln \left( \frac{M_i}{M_0} \right) + \frac{i}{ij-j^2} \ln \left( \frac{M_j}{M_0} \right) \text{ and} \]
   \[ \sigma^2 = \frac{1}{1-i/j} \left[ \frac{2}{j^2} \ln \left( \frac{M_j}{M_0} \right) - \frac{2}{ij} \ln \left( \frac{M_i}{M_0} \right) \right] \]
Spatial discretization in QMOM

Moment corruption detection and correction

- Negative values!
- Moment corruption
detection and correction

- Instability problems
- Residuals
- No instability problems
ISSUES RELATED TO SPATIAL DISCRETIZATION WITH DQMOM: DQMOM-FC
The original GPBE is:

\[ \frac{\partial n}{\partial t} + \frac{\partial}{\partial x} \cdot (U_p n) = S \]

With **DQMOM** the NDF is described in terms of weights, \( w_\alpha \), and nodes, \( L_\alpha - \phi_\alpha \), so that some moments \( M_{k,l} \) are correctly predicted:

\[ \frac{\partial w_\alpha}{\partial t} + \frac{\partial}{\partial x} \cdot (U_{p,\alpha} w_\alpha) = a_\alpha \]

however this does not conserve moments (in FV codes)

Therefore convection is treated as a source term:

\[ \frac{\partial n}{\partial t} = - \frac{\partial}{\partial x} \cdot (U_p n) + S \]

Resulting in (**DQMOM-Fully-Conservative**):

\[ \frac{\partial w_\alpha}{\partial t} = a_\alpha^* + a_\alpha \]
Spatial discretization in DQMOM

Oxygen concentration in gas phase (mol m$^{-3}$)
No mass transfer

Results obtained with the commercial code Fluent
CONCLUSIONS
• The simulation of polydisperse multiphase systems can be efficiently done at the macroscale

• Quadrature-based moment methods (QBMM) can be used for reconstructing the NDF and overcoming the closure problem

• They combine accuracy and computational efficiency

• Currently implemented in Fluent, TransAT and openFoam

• DQMOM has many issues (the ones related to spatial discretization in finite volume codes are fixed by using DQMOM-FC)

• QMOM/CQMOM are way more efficient but attention should be paid to moment realizability/corruption

• Very important is also the problem of quadrature realizability

• Future work: development of mesoscale models for real systems
CQMOM for a stirred tank

Results and comparison with experimental data

- **N = 157 RPM; 0.052 vvm**
  - R2, R4, R8, R9, R12

- **N = 250 RPM; 0.052 vvm**
  - R2, R4, R8, R9, R12

- **N = 250 RPM; 0.072 vvm**
  - R2, R4, R8, R9, R12

Legend:
- Experimental
- Simulation (45°)
- Simulation (135°)
CQMOM for a stirred tank

Experimental and simulated mean bubble size (mm) in five different points of the stirred tank.

<table>
<thead>
<tr>
<th>SR, RPM</th>
<th>GR, vvm</th>
<th>R2</th>
<th>R4</th>
<th>R8</th>
<th>R9</th>
<th>R12</th>
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<tbody>
<tr>
<td>155</td>
<td>0.018</td>
<td>Exp.</td>
<td>2.37</td>
<td>2.48</td>
<td>2.29</td>
<td>1.65</td>
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<td></td>
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<td>Sim.</td>
<td>3.10</td>
<td>2.56</td>
<td>2.57</td>
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<td>Sim.</td>
<td>2.45</td>
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<td>250</td>
<td>0.093</td>
<td>Exp.</td>
<td>2.96</td>
<td>3.25</td>
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<td>Sim.</td>
<td>2.56</td>
<td>3.27</td>
<td>2.59</td>
<td>3.05</td>
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</tbody>
</table>
CQMOM for a stirred tank

Global gas hold-up for 390 rpm

- Experimental data
- Simulation
- DRAG CORRECTION

Graph: 
- Y-axis: global gas hold-up
- X-axis: gas flow rate, vvm
- Data points for different regimes:
  - Transitional regime
  - DRAG correction

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CQMOM for a stirred tank

Specific surface area of bubbles (m$^{-1}$) at 250 rpm and 0.052 vvm (left) and at 155 rpm and 0.018 vvm (right)

1.20e+02
1.08e+02
9.60e+01
8.40e+01
7.20e+01
6.00e+01
4.80e+01
3.60e+01
2.40e+01
1.20e+01
0.00e+00
CQMOM for a stirred tank

Mass transfer coefficient $k_l$ (m s$^{-1}$) at 250 rpm and 0.052 vvm (left) and at 155 rpm and 0.018 vvm (right)
CQMOM for a stirred tank

Oxygen concentration (mol m$^{-3}$) in the gas (left) and in the liquid (right) at 250 rpm and 0.052 vvm
DSMC for a stirred tank

Effect number of comp. on gas hold-up and Sauter diameter

![Graph showing the effect of the number of components on gas hold-up and Sauter diameter.](image-url)
Let us now consider a realistic stirred tank

- Rushton turbine (six blades) - reactor volume 15.3 l
- Gassing rate \( \approx 0.062 \) l/s; stirring rate 100-300 rpm
- Flow field simulation ANSYS/Fluent13 - DSMC in-house code

DSMC for a stirred tank

Compartment model

<table>
<thead>
<tr>
<th>Tot. numb. comp.</th>
<th>Turb. diss. rate $m^2s^{-3}$</th>
<th>$N \approx 2$</th>
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DSMC for a stirred tank

Compartment model

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<td>$N \approx 5$</td>
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DSMC for a stirred tank

Compartment model

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D. L. Marchisio
DSMC for a stirred tank

Bubble size-composition in two regions
DSMC for a stirred tank

Effect of chemical reaction \( (H_a = \sqrt{kC_{O_2}D/k_l}) \)

\begin{align*}
\text{Example 1:} & \quad L, \text{ m} \\
C, \text{ mol/m}^3 & \quad L, \text{ m} \\
0.00 \times 10^0 & \quad 0.00 \times 10^0 \\
0.80 \times 10^0 & \quad 0.80 \times 10^0 \\
0.60 \times 10^0 & \quad 0.60 \times 10^0 \\
0.40 \times 10^0 & \quad 0.40 \times 10^0 \\
0.20 \times 10^0 & \quad 0.20 \times 10^0 \\
0.00 \times 10^0 & \quad 0.00 \times 10^0 \\
\end{align*}

\begin{align*}
\text{Example 2:} & \quad L, \text{ m} \\
C, \text{ mol/m}^3 & \quad L, \text{ m} \\
0.00 \times 10^0 & \quad 0.00 \times 10^0 \\
0.80 \times 10^0 & \quad 0.80 \times 10^0 \\
0.60 \times 10^0 & \quad 0.60 \times 10^0 \\
0.40 \times 10^0 & \quad 0.40 \times 10^0 \\
0.20 \times 10^0 & \quad 0.20 \times 10^0 \\
0.00 \times 10^0 & \quad 0.00 \times 10^0 \\
\end{align*}
Comparison DSMC with CQMOM/DQMOM

**Table:** Mean percentage error of QBMM for the examined case at steady-state.

<table>
<thead>
<tr>
<th></th>
<th>$M_{0,0}$</th>
<th>$M_{1,0}$</th>
<th>$M_{2,0}$</th>
<th>$M_{3,0}$</th>
<th>$M_{4,0}$</th>
<th>$M_{5,0}$</th>
<th>$M_{6,0}$</th>
<th>$M_{7,0}$</th>
</tr>
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<tbody>
<tr>
<td>$N=2$</td>
<td>10.1</td>
<td>7.9</td>
<td>4.6</td>
<td>0.5</td>
<td>6.5</td>
<td>22.3</td>
<td>38.3</td>
<td>50.1</td>
</tr>
<tr>
<td>$N=3$</td>
<td>7.9</td>
<td>5.2</td>
<td>2.3</td>
<td>0.5</td>
<td>3.3</td>
<td>6.2</td>
<td>8.2</td>
<td>6.9</td>
</tr>
<tr>
<td>$N=4$</td>
<td>7.7</td>
<td>5.1</td>
<td>2.4</td>
<td>0.4</td>
<td>3.3</td>
<td>6.4</td>
<td>9.5</td>
<td>12.7</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th></th>
<th>$M_{0,1}$</th>
<th>$M_{1,1}$</th>
<th>$M_{2,1}$</th>
<th>$M_{3,1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N=2$</td>
<td>2.1</td>
<td>6.5</td>
<td>21.4</td>
<td>44.6</td>
</tr>
<tr>
<td>$N=3$</td>
<td>1.9</td>
<td>1.8</td>
<td>4.7</td>
<td>6.2</td>
</tr>
<tr>
<td>$N=4$</td>
<td>3.2</td>
<td>1.7</td>
<td>3.4</td>
<td>6.4</td>
</tr>
</tbody>
</table>
Analogously the mass transfer coefficient is calculated with Danckwerts theory:\(^3\):

\[ k_L = 1.13 \mathcal{D}^{0.5} \left( \frac{\epsilon \rho_c}{\mu_c} \right)^{0.25} \]

Consequently the mass transfer rate reads as follows:

\[ \dot{\phi}_b = k_L k_a L^2 \left( C_c - H \frac{\phi_b}{k_v L^3} \right) \]

whereas the rate of change of bubble size:

\[ G = \frac{2k_L M_w}{\rho_b} \left( C_c - H \frac{\phi_b}{k_v L^3} \right) \]

---

Discontinuous term meso-scale model

Coalescence kernel

A kernel written in terms of bubble size is used:\n
$$a(\lambda, L) = 0.88 \epsilon^{1/3} (\lambda + L)^2 (\lambda^{2/3} + L^{2/3})^{1/2} \eta(\lambda, L)$$

where it is implicitly assumed that turbulent fluctuations are the driving force for bubble collisions.

The coalescence efficiency $\eta(\lambda, L)$ can be written as follows:\n
$$\eta(\lambda, L) = \exp \left( -6 \cdot 10^9 \frac{\mu_c \rho_c \epsilon}{\sigma^2} \left( \frac{\lambda L}{\lambda + L} \right)^4 \right)$$


Discontinuous term meso-scale model

Breakage kernel

\[ b(L) = 6.0 \epsilon^{1/3} \text{erfc} \left( \sqrt{0.04 \frac{\sigma}{\rho_c \epsilon^{2/3} L^{5/3}}} + 0.01 \frac{\mu_c}{\sqrt{\rho_c \rho_b} \epsilon^{1/3} L^{4/3}} \right) \]

Daughter distribution function

Generic $\beta$-PDF for bubble size (and bubble composition) where $C = 2$ for binary breakage:

\[
P(L|\lambda) = \frac{1}{12} (C + 1)^2 (C + 2)^2 (C + 3)^2 (C + 4)^2 \times \frac{L^2}{\lambda^3} \left( \frac{L^3}{\lambda^3} \right)^2 \left( 1 - \frac{L^3}{\lambda^3} \right)^C
\]

Comparison DSMC with CQMOM/DQMOM

- With DSMC the evolution of the population of particles is represented through its stochastic equivalent (with \( N \approx 100 \) notional particles)
- With QMOM/CQMOM/DQMOM the evolution is tracked deterministically with \( N \approx 3 - 6 \) macro-classes!
- These macro-classes are centered at the quadrature nodes
- Therefore all the unclosed terms appearing in the equations for the moments are closed with great accuracy
- The two distributions share the same moments of the NDF:

\[
M_{k,l}(t, x) = \int \int n(t, x, L, \phi_p) L^k \phi_p^l \, dL \, d\phi_p
\]

- Sharing the same moments does not imply that the quadrature is realizable (for example for specific applications some moments have to be included!)
Differences between QMOM/CQMOM/DQMOM

- When transporting moments (QMOM/CQMOM) the governing equations are solved for CONSERVATIVE variables, ensuring the conservation of important properties (in finite volume codes)

- Time and space discretization can CORRUPT a moment set causing instabilities when the inversion algorithm is applied

- Moment corruption is significant when adopting higher-order spatial discretization (and interpolation) schemes, whereas is limited when using first-order upwind schemes

- Several algorithms for detecting moment corruption and for correcting an invalid moment set have been developed

- Also ad-hoc accurate spatial discretization schemes that preserve the consistency of a moment set have been developed

---

Differences between QMOM/CQMOM/DQMOM

- When transporting the quadrature approximation (DQMOM) the equations are solved for PRIMITIVE variables making non-trivial the conservation of moments.

- When numerical diffusion (always present in finite volume codes) dominates, even the conservation of low-order moments is not guaranteed.

- The presence of diffusion generates spurious terms that must be corrected; the correction is readily calculated when the diffusion term is exactly known but in the case of numerical diffusion it can be very problematic.

- A new method has been developed: DQMOM fully-conservative (DQMOM-FC).

- Another well known problem of DQMOM is that the algorithm becomes unstable anytime two nodes of the quadrature approximation are characterized by the same values.

---

Coupling QMOM/CQMOM/DQMOM with particle advection

- QMOM/CQMOM/DQMOM can be used with the assumption that all the particles are moving with the same average velocity\(^6\)
- DQMOM can account for the fact that particles move with different velocities
- QMOM/CQMOM can also be used with particle velocity that depends on particle size, resulting in advection terms that are different for different moments\(^7\)
- CQMOM/DQMOM can be used to describe the NDF including particle velocity\(^8\)
- Moreover being most of these systems turbulent: DNS, LES or RANS can be used

\(^7\) Mazzei, Marchisio, Lettieri, 2012. AIChE J., 58, 3054-3069.
A moment set is said to be valid, if the Hankel-Hadamard determinants are all non-negative:

\[ \Delta_{k,l} = \begin{vmatrix} M_k & M_{k+1} & \cdots & M_{k+l} \\ M_{k+1} & M_{k+2} & \cdots & M_{k+l+1} \\ \vdots & \vdots & \ddots & \vdots \\ M_{k+l} & M_{k+l+1} & \cdots & M_{k+l+l} \end{vmatrix} \geq 0 \]

for \( k = 0, 1 \) and \( l \geq 0 \). A less stringent condition is that of convexity of the function \( \ln(M_k) \) with respect to \( k \):

\[
\frac{\ln(M_k) + \ln(M_{k-2})}{2} \geq \ln(M_{k-1}) \quad k = 1, 2, \ldots;
\]

or, equivalently,

\[
M_k M_{k-2} - M_{k-1}^2 \geq 0 \quad k = 1, 2, \ldots.
\]
Spatial discretization in QMOM

The convexity of the function $\ln(M_k)$ with respect to $k$ can be easily verified by building a difference table of $\ln(M_k)$.

Example: VALID SET; moment of a Gaussian distribution ($M_0 = 1, M_1 = 5, M_2 = 26, M_3 = 140, M_4 = 778, M_5 = 4450, M_6 = 26140, M_7 = 157400$)

<table>
<thead>
<tr>
<th>$k$</th>
<th>$d_0 = \ln(M_k)$</th>
<th>$d_1$</th>
<th>$d_2$</th>
<th>$d_3$</th>
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<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1.609</td>
<td>0.039</td>
<td>-0.0043</td>
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<td>0.034</td>
<td>-0.0033</td>
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<td>1.683</td>
<td>0.031</td>
<td>-0.0027</td>
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<td>-0.0022</td>
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Spatial discretization in QMOM

The convexity of the function $\ln(M_k)$ with respect to $k$ can be easily verified by building a difference table of $\ln(M_k)$. Example: INVALID SET (modify the second-order moment from 26 to 25, corresponding to a difference of only 4 %)

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<tr>
<th>$k$</th>
<th>$d_0 = \ln(m_k)$</th>
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<td>0.113</td>
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<td>1.609</td>
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Spatial discretization in QMOM

- Since it is very easy to CORRUPT the moments it is important to have CORRECTION algorithms
- If we transform the moment set so that \( d_2 \) is positive, we are almost sure that the moment set is valid
- But how positive?
- The moments of a log-normal distribution have the smallest \( d_3 \)

\[
n(L) = \frac{N_T}{\sigma \sqrt{2\pi}} \exp \left( \frac{-(\ln(L) - \mu)^2}{2\sigma^2} \right),
\]

\[
M_k = N_T \exp \left( k\mu + \frac{k^2\sigma^2}{2} \right),
\]

- The log-normal distribution is the smoothest distribution!
Spatial discretization in QMOM

- One solution would be to evaluate the moments at the faces $M^e_k$ and $M^w_k$ through the quadrature approximation.
- We know the value of the moments at the center of the cells $M^W_k, M^P_k, M^E_k$.

From these moments we can evaluate the corresponding weights $w^P_\alpha$ and abscissas $L^P_\alpha$.

If weights and abscissas at the center of the face are interpolated with $p^{th}$-order spatial reconstruction the resulting moments will be valid.

This allows to improve the numerical accuracy preserving the moments!
Spatial discretization in DQMOM

Steady state solution for bubbles undergoing aggregation and breakage → DQMOM/DQMOM-FC
CQMOM for a stirred tank

Results and comparison with experimental data

Reactor configuration 1 – N = 250 RPM; 0.093 vvm

Bubble volume density vs d, mm

- Experimental data
- Simulation results

Spatial discretization in DQMOM

Conclusions
CQMOM for a stirred tank

Time evolution of oxygen concentration in the liquid

250 rpm, 0.052 vvm

155 rpm, 0.018 vvm
CQMOM for a stirred tank

Example of the poor performance of fixed-constant bubble 3-D simulation for mass transfer!