CFD-DEM study of mass transfer mechanisms in riser flow

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Abstract

In this paper we report a CFD-DEM simulation study on the interplay between mass transfer and a heterogeneous catalyzed chemical reaction in co-current gas-particle flows as encountered in risers. Slip velocity, axial gas dispersion, gas bypassing and particle mixing phenomena have been evaluated under riser flow conditions to study the complex system behavior in detail. The most important factors are found to be directly related to particle cluster formation. Low air to solids flux ratios lead to more heterogeneous systems, where the cluster formation is more pronounced and mass transfer more influenced.

Falling clusters can be partially circumvented by the gas phase, which therefore does not fully interact with the cluster particles, leading to poor gas-solid contact efficiencies. Cluster gas-solid contact efficiencies are quantified at several gas superficial velocities, reaction rates and dilution factors in order to gain more insight regarding the influence of clustering phenomena on the performance of riser reactors.
Introduction

Mass and heat transfer phenomena under riser flow conditions have been widely investigated during the last decades. The usage of mass and heat transfer coefficients is essential in phenomenological and computational models to estimate the performance of chemical processes in bubbling and fast fluidized bed reactors. However, there is still a lack of understanding regarding the applicability of the different correlations that could be employed to estimate mass and heat transfer coefficients in fluidized systems. Many mass transfer correlations for fluidized systems have been proposed to compute the dimensionless mass transfer coefficient of a particle in a fluidized system.\textsuperscript{1,2} Furthermore, Breault reported that Sherwood numbers in riser fluidized systems can differ in several orders of magnitude,\textsuperscript{3} so it is evident that there is not a unique and unequivocal equation to compute a mass transfer coefficient for fluidized systems. The reason for this is thought to be the presence of heterogeneities in the particle distributions that severely influence the flow patterns and consequently the mass transfer to particles.

Flow heterogeneities are especially prevalent in riser flows, which are characterized by a core annulus flow involving a rather dilute region in the core of the riser and a dense particulate phase close to the walls. The dense solids phase can be either formed by a falling solids film (annulus) or can be a more cluster-like flow, which intermittent behavior can be represented by, e.g., an intermittency index.\textsuperscript{4} When particle clusters are formed: low slip velocities, gas back-mixing and poor gas-solid contacting are key hydrodynamic phenomena that prevail. Besides heterogeneities, another cause of disagreement between mass transfer coefficients reported in literature is caused by different model interpretations and/or definitions of mass transfer coefficients. Thus, it should be noted that the obtained values of mass transfer coefficients can be completely meaningful within the context of their respective models, but these data should in principle not be used in other scenarios where (completely) different hypotheses and/or modeling assumptions have been taken.

In this paper, we generate insight about the mass transport mechanisms related to cluster
formation and the causes of the aforementioned disagreement. For this purpose we apply CFD-DEM simulations. CFD-DEM simulations not only can estimate the global process efficiency of a riser reactor, but also provide detailed information about cluster-related phenomena. The reason is that in CFD-DEM all particles are explicitly modeled. Therefore phenomena such as clustering of particles are emergent. This means that closures for mass transfer are needed at the particle level only. Extra mass transfer resistances related to the presence of particle clusters are a result of the simulations. Thus, by means of CFD-DEM simulations, a detailed analysis is performed regarding the influence of several mass transfer mechanisms on the performance of a pseudo-2D riser reactor.

Frössling-type correlations have been reported to predict accurately the mass transfer rate of a single particle immersed in a fluid flow.\textsuperscript{1,5,6} Convection has a strong impact on the thickness of the momentum and mass boundary layers, and for higher Reynolds number determines the mass transfer resistance at the particle level.\textsuperscript{7} Due to the thin boundary layer the particle Sherwood number of an isolated particle at larger Reynolds number will be bigger than 2. When particles are surrounded by other particles, such as in packed bed reactors, the mass transfer is influenced significantly compared to isolated particle mass transfer.\textsuperscript{8–10} Gunn’s correlation, which is widely used in literature,\textsuperscript{11} was developed from experiments in a packed bed reactor and a liquid fluidized bed reactor. The Gunn correlation has also been shown to represent well the fluid to particle mass/heat transfer in resolved simulations of dense\textsuperscript{12} and more dilute stationary particle arrangements\textsuperscript{13,14} as well as solid-liquid systems.\textsuperscript{15} Note that the dependence on solids-volume fraction is different if a particle that exchanges mass is surrounded by inert particles that do not exchange mass.\textsuperscript{1,16–19} Such ‘diluted’ systems have been employed to not reach saturated gas naphthalene concentrations\textsuperscript{1} or to not reach too high conversion rates. So, the presence of other particles significantly influences the mass transfer to individual particles. It has to be noted that e.g. Gunn’s correlation was obtained from experimental measurements in a packed bed and a liquid fluidized bed systems. However, in riser flows, particles may form heterogeneous structures
and the gas solid distributions may not be as homogeneous as in a fixed bed reactor or a bubbling liquid fluidized bed. It should be emphasized that Gunn’s correlation is valid if we assume that gas and solids distributions are relatively homogeneous. Thus, in our simulations, Gunn’s correlation is employed at a sufficiently small scale where the flow can be regarded as homogeneous, e.g. the CFD-DEM cell size.

For heterogeneous systems, the distinction between the local particle mass transfer coefficient and the global or overall (i.e. reactor length scale) mass transfer coefficient should be made. A particle-based mass transfer coefficient is interpreted as the mass transfer rate of a single particle that exchanges mass with the surrounding fluid phase, where the driving force is defined by local concentration differences. At the same time, an overall mass transfer coefficient can well represent the mass transfer rate at the reactor scale. Here the driving force for mass transfer is globally defined e.g. by means of a gas concentration cup-averaged over the reactor cross section. However, different hydrodynamic models can be found in literature and consequently definitions of global mass transfer coefficients could also differ. Thus, the measurements of global mass transfer coefficients in fluidized systems (e.g., assuming a 1D plug flow interpretation model) can be clearly different to Gunn’s correlation predictions (using a global driving force) because the flow is heterogeneous and local particle-scale driving forces are different from the global reactor-scale driving forces.\textsuperscript{20,21} Larger (than particle level) scale hydrodynamic resistances can play an important role in mass transfer phenomena.\textsuperscript{22–25}

Breault et al. reported that Sherwood numbers in riser fluidized systems can differ in several orders of magnitude.\textsuperscript{3} This spread in reported values is likely due to the presence of different levels of heterogeneity in different experiments. However, another important cause of these differences is that different models and definitions have been employed to report such Sherwood numbers. For instance, Subbarao and Gambhir\textsuperscript{26} utilized naphthalene deposition on sand particles at the bottom of a riser as a model experimental system to develop a correlation to compute an overall mass transfer coefficient. In their work, circulation pat-
tern effects are lumped into global parameters of gas superficial velocity and solids mass flux. Wang and Li\textsuperscript{27} used a TFM coupled to a Energy-Minimization Multi-Scale (EMMS) method to model the local heterogeneous flow structure. The model was validated, using a particle-based Sherwood correlation,\textsuperscript{28} with experimental measurements of naphthalene sublimation.\textsuperscript{27} Venderbosch et al.\textsuperscript{16} employed CO oxidation as model reaction to determine mass transfer-limited rates of a diluted (chemically active and inert particles) fluidized system, where platinum supported FCC particles (catalytically active) were mixed with unsupported FCC particles (inert). Sherwood numbers reported by the previously mentioned authors can easily differ in several orders of magnitude and all might be correct for their respective models. In Table S4 (in the Supporting Information), an overview of relevant publications related to mass transfer phenomena on riser flow is provided. It can be noticed that CFD computational strategies, which account for the flow heterogeneous, have become more popular in recent years. However, the computational expenses of these models are still (too) high to apply them at large industrial scale.

Thus, some attempts have been made to develop correlations that compute overall mass transfer coefficients that could be easily estimated under riser flow conditions with known operating parameters such as the gas superficial velocity and solids mass flux of a riser.\textsuperscript{23,26,29} Other authors suggested that the cluster-bulk mass transfer coefficient is proportional to the cluster surface renewal rate and that this is defined by their respective size and velocity.\textsuperscript{30–33} A lot of core-annular models have been developed to predict the solids and gas distributions under riser flow conditions\textsuperscript{34–37} and to estimate the performance of a riser reactor. However, these are mainly used to model only fully developed sections of a riser and do not model complex flow structures, which are highly dependent on the physical properties of the particles and the reactor geometry. It is known that riser hydrodynamics can be highly affected by the riser diameter, entrance and exit effects and therefore more detailed models are required. Thus, high precision CFD models are expected to describe riser flow hydrodynamics more accurately and provide a complete picture of the system to facilitate design and upgrading.
of riser reactors.

EMMS-based computational works\textsuperscript{38,39} and filtered closures\textsuperscript{40} have been employed to characterize flow heterogeneities. In this way, heterogeneity is modeled in Euler-Euler simulations, which resolution scale can be significantly larger than the typical cluster size. For instance, EMMS schemes are based on energy minimization assumptions to model flow heterogeneous structures. Analogous EMMS/mass transfer methods can be employed to determine interphase mass transfer coefficients: from gas to particles of a dilute phase, from gas to particles of the dense phase and from gas to cluster surface.\textsuperscript{20,21} Other authors employed EMMS/mass transfer models to analyze the influence of particle heterogeneity over the riser performance.\textsuperscript{41–43} Although some of the EMMS-based computational studies account for clustering phenomena, there are simplifying assumptions made. Some of these are: Clusters have uniform density, equal to maximum packing fraction;\textsuperscript{44} Clusters are spherical structures that interact with a lean surrounding phase and are homogeneously dispersed in a control volume.\textsuperscript{45} Some of these computational studies have suggested that global scale mass transfer resistances are related to cluster formation.\textsuperscript{20,25,46} However, these findings are not explicitly supported with cluster scale level computations, such as the instantaneous gas-solid contact efficiency of particle clusters.

In riser flows, the definition of a gas solid contact efficiency has been employed to quantify the deviation of a riser flow from an idealized plug flow, where all catalyst particles are fully exposed to the bulk fluid phase. The contact efficiency is not only related to the exposed area of a single particle to the surrounding gas phase,\textsuperscript{47} but also related to the particle exposure to a reactant-rich gas phase that tends to circumvent dense particulate regions,\textsuperscript{48} which can retain gas pockets of highly depleted reactant. In experimental works,\textsuperscript{35,49–52} inefficient contacting was presumed to be due to cluster formation since gas dispersion effects could be discarded.\textsuperscript{16,50} In these cases, the quantification of gas-solid contact efficiency was computed from time-averaged properties (solids volume fraction, reactant concentration and gas velocity) without obtaining instantaneous data that confirmed these hypotheses. Previous
authors\textsuperscript{25,42} analyzed the influence of the reaction rate on the mass transfer rate, which is reported to be less affected at lower values of \(k_r\).\textsuperscript{42} The difficulty to measure experimental data at cluster level has not allowed yet to confirm these assumptions by a direct method.

CFD-DEM has been shown to be a suitable model to predict riser hydrodynamics and complex clustering phenomena,\textsuperscript{53} study the particles mixing and deactivation in a catalytic process, or even track the real-time particle activity.\textsuperscript{54,55} In this paper, different mass transfer mechanisms are isolated using ozone decomposition as a model reaction. The aim is to quantify their respective contribution to mass transfer-limited chemical processes, and obtain a close relation between cluster formation and gas-solid contact efficiency. Besides this, we quantify the instantaneous gas-solid contact efficiency by relating the ozone gas mass fraction inside the cluster and the bulk ozone gas mass fraction at the cross section where each detected cluster is located. These results illustrate the severe impact that clustering phenomena can have on gas-solid contacting and the estimation of global mass transfer coefficients. By means of a CFD-DEM we confirm show that clustering phenomena have more severe impact at higher reaction rates.

**Methodology**

A CFD-DEM has been utilized to resolve simulations of a pseudo-2D riser bed reactor. The governing equations for the gas phase have been solved using a semi-implicit finite difference technique.\textsuperscript{56} For the species field the gas convection-diffusion equation has been solved as well:

\[
\frac{\partial (\varepsilon \rho_g)}{\partial t} + \nabla \cdot (\varepsilon \rho_g \mathbf{u}) = 0
\]  
\[ (1) \]

\[
\frac{\partial (\varepsilon \rho_g \mathbf{u})}{\partial t} + \nabla \cdot (\varepsilon \rho_g \mathbf{u} \mathbf{u}) = -\varepsilon \nabla P - \nabla \cdot (\varepsilon \mathbf{\tau}) - \mathbf{S}_p + \varepsilon \rho_g \mathbf{g}
\]  
\[ (2) \]

\[
\frac{\partial w_{A,g}}{\partial t} + (\nabla w_{A,g} \cdot \mathbf{u}) = \nabla \cdot D_{A,B}^{eff} \cdot \nabla w_{A,g} + S_e
\]  
\[ (3) \]
The Zehner and Schlünder model is used to evaluate the effective diffusivity:

\[ D_{AB}^{eff} = \frac{1 - (\sqrt{1 - \varepsilon})}{\varepsilon} D_{AB} \]  

(4)

The gas and solids motion is coupled via a sink term that involves the computation of the interphase momentum transfer coefficient by means of the Beetstra drag correlation. Additionally, gas phase mass balances are coupled to the mass balance at the particle level by means of a sink term, where the mass transfer coefficient is computed by means of the Gunn correlation:

\[ S_p = \frac{1}{V_{cell}} \sum_{i=0}^{N_p} \beta V_p \left( u - v_p \right) D \left( r - r_p \right) \]  

(5)

\[ S_e = \frac{1}{V_{cell}} \sum_{i=0}^{N_p} k_{mt} \cdot A_p \left( w_{A,g} - w_{A,p} \right) D \left( r - r_p \right) \]  

(6)

where \( A_p \) and \( V_p \) are respectively the surface area and volume of a single particle, \( \beta \) is the interphase momentum transfer coefficient and \( k_{mt} \) is the computed particle-based mass transfer coefficient by Gunn correlation.

Two way coupling is performed by means of a regularized Dirac delta function \( D \left( r - r_p \right) \), which maps the gas phase variables from neighboring Cartesian nodes to the particle location to enable evaluation of the drag force and the interphase mass transfer coefficient. Moreover, changes in particle momentum are fed back from the particle position to the surrounding Eulerian nodes using the same regularized function.

The particle translational and rotational momentum is governed by the Newtonian equations of motion:

\[ m_p \frac{d^2 r_p}{dt^2} = -V_i \nabla P + \frac{\beta V_p}{1 - \varepsilon} \left( u - v_p \right) + m_p g + F_c \]  

(7)

\[ I_p \frac{d \omega_p}{dt} = T_p \]  

(8)
The particle collisional forces are deterministically computed, by means of a soft sphere model that was originally proposed by Cundall and Strack\textsuperscript{59} and first employed in a gas fluidized system simulation by Tsuji et al.\textsuperscript{60} Gas turbulence was assumed to be insignificant compared to the velocity fluctuations due to gas-particle interaction, following other authors.\textsuperscript{61,62} Thus, a subgrid turbulence model was not employed. Additionally, no mass transfer exchange was taken into account during particles collisions.

A first order irreversible reaction under riser flow conditions was included in the CFD-DEM to simulate ozone decomposition:

$$2\text{O}_3 \rightarrow 3\text{O}_2$$

Pure ozone was fed from the bottom of the riser. It was assumed that ozone reacts at the particle surface without taking into consideration internal mass transfer effects throughout the particle volume:

$$\frac{dw_{A,p}}{dt} = \left( \frac{6}{d_p} \text{Sh} \left( \frac{D_{AB}}{d_p} (w_{A,g} - w_{A,p}) - k_r w_{A,p} \right) \right)$$ \hspace{1cm} (9)

After integration, the particle ozone mass fraction can be expressed as follows:

$$w_{A,p}(t) = \frac{6}{d_p} \frac{\text{Sh} D_{AB} w_{A,g}}{6/d_p \text{Sh} D_{AB} + k_r d_p} \left[ 1 - \exp \left( -\frac{6}{d_p} \left( \text{Sh} \left( \frac{D_{AB}}{d_p} + k_r \right) t \right) \right) \right]$$ \hspace{1cm} (10)

If we compute the characteristic time scale of the exponential term, assuming a 0.85 mm glass bead with a slip velocity of 1 m/s and $k_r = 10$ s\textsuperscript{-1}, we obtain that

$$t = \frac{1}{\frac{6}{d_p} \left( \text{Sh} \left( \frac{D_{AB}}{d_p} + k_r \right) \right)} = 1.4 \cdot 10^{-5} \text{seconds}$$ \hspace{1cm} (11)

which is lower than the computational gas time step of our simulations ($5 \cdot 10^{-5}$ s). It is noted that the computed characteristic time scale is an upper bound (when $k_r = 10$ s\textsuperscript{-1}) in our set of simulations. So, for all other values of $k_r$ considered in this work, this parameter is even
lower. Since this reaction is fast compared to the gas residence time in the system, the mass balance at the particle level is well approximated by:

\[
\frac{6}{d_p} \text{Sh} \frac{D_{AB}}{d_p} (w_{A,g} - w_{A,p}) - k_r w_{A,p} = 0
\]  

(12)

where the Sherwood number was computed by means of the Gunn correlation.\textsuperscript{8} As for momentum, the ozone mass fraction is computed for all particles in the system. The gas concentration \(w_{A,g}\), is interpolated from the neighboring Cartesian nodes to the particles locations, which are represented as points and act as ozone sink sources. As a result, the particle mass fraction \(w_{A,p}\) is computed by means of equation 12 and the gas mass fraction is updated via two-way coupling.\textsuperscript{56}

Thus, the source term (equation 6) can be expressed as follows:

\[
S_e = \frac{1}{V_{cell}} \sum_{i=0}^{N_p} k_r k_{nt} \cdot \frac{A_p w_{A,g} D (r - r_p)}{d_p^2 k_{nt} + k_r}
\]  

(13)

Simulation conditions.

Figure 1 shows the simulation domain used. This consisted of a pseudo-2D riser of \(1.57 \times 0.07 \times 0.006\) meters. Particles were fixed at the top of the simulation domain to mimic the lateral curved outlet of an experimental unit published elsewhere.\textsuperscript{53} In the simulations, a system with a highly diluted ozone content is simulated. Thus, physical properties of air gas are used in the simulations, and corrections for molecular counter diffusion of reaction products are not needed. Besides, the reaction is assumed to be equimolar and no heat effects are considered. More details about the simulation settings and physical parameters used are listed in Tables 1 and 2. During the whole simulation Geldart D particles were placed at random x-y coordinates near the bottom of the riser at a height of 2-3 times the particle radius above \(z = 0\). The insertion was only accepted if there was no particle overlap. The simulations were performed under fast fluidization regime at several gas superficial velocities.
(\(U = 5.55, 5.95, 6.35\) and \(6.74\) m/s) and a fixed solids mass flux rate of 32 kg/(m\(^2\)s). The particles were inserted at random positions of the bottom X-Y plane of the simulation domain at a velocity that was nearly zero (0.01 m/s). And the particles that reached the lateral outlet, left the simulation domain.

At the top, front, back and right walls, no-slip boundary conditions were applied. With a prescribed inflow axial velocity equal to \(U\), gas was supplied at the bottom of the domain. The left side wall (\(x=0\)) was subdivided into two regions: for the top-left outflow region of 0.07 m the pressure \(P_0\) is described and Neumann conditions were applied for the species field. Below this region no-slip and no-flux boundary conditions were applied for the gas momentum and ozone mass fraction respectively, as Figure 1 illustrates.
Table 1: Simulation conditions.

| Variable | Value |
|----------|-------|
| NX | 28 |
| $d_p$ (mm) | 0.85 |
| NY | 5 |
| $\rho_s$ (kg/m$^3$) | 2500 |
| NZ | 628 |
| $\mu_g$ (kg/m·s) | 2.0·10$^{-5}$ |
| X (m) | 0.07 |
| T (K) | 298 |
| Y (m) | 0.006 |
| $e_{p-p}$ | 0.96 |
| Z (m) | 1.57 |
| $e_t$ | 0.33 |
| $\Delta t_{\text{Gas}}$ (s) | 5.0·10$^{-5}$ |
| $\Delta t_{\text{DEM}}$ (s) | 5.0·10$^{-6}$ |
| $G_s$ kg/(m$^2$·s) | 5.0·10$^{-5}$ |
| U (m/s) | 5.16-6.74 |
| P | 1 atm |

Table 2: Characteristic dimensionless numbers.

| U (m/s) | $\langle \varphi_s \rangle$ | Re | Pe$_p$ | St |
|---------|-----------------|-----|-------|-----|
| 5.16 | 0.0567 | 1858 | 274 | 28 |
| 5.55 | 0.0410 | 1998 | 295 | 31 |
| 5.95 | 0.0228 | 2142 | 316 | 33 |
| 6.35 | 0.0147 | 2286 | 337 | 35 |
| 6.74 | 0.0094 | 2426 | 358 | 37 |

The Damköhler number for all simulations presented in this work ranges between 0.014 (U = 6.74 m/s and $k_r = 100$ s$^{-1}$) and 13.5 (U = 5.95 m/s and $k_r = 1000$ s$^{-1}$). These parameters have been computed assuming an effective molecular diffusivity of 1.6·10$^{-5}$ m$^2$/s and a characteristic hydraulic diameter equal to the depth of the system (6 mm).
Gas-solid contact efficiency

In order to quantify the gas-solid contacting it is necessary to provide a parameter that captures this effect, e.g. a contact efficiency. Otherwise, by assuming a homogeneous system, e.g. ideal plug flow, overestimation of the conversion rate in heterogeneous systems is likely to result. The contact efficiency could be determined by assuming a riser as a steady state plug flow reactor:

\[ U \frac{dw_{A,g}}{dz} = -K_{ov}w_{A,g}\varphi_s \]  

with a solution:

\[ \frac{w_{A,g,z+\Delta z}}{w_{A,g,z}} = \exp\left( -\frac{K_{ov}\varphi_s \cdot \Delta z}{U} \right) = \exp\left( -\gamma_{pf} \cdot Da \cdot \frac{\Delta z}{L_{riser}} \right) \]  

where \( \varphi_s \) is the averaged solids volume fraction in a slice of thickness \( \Delta z \), \( U \) is the gas superficial velocity, \( K_{ov} \) is the apparent volumetric reaction rate constant, \( \gamma_{pf} \) is a gas-solid contact efficiency and \( Da \) is the Damköhler number defined as:

\[ \gamma_{pf} = \frac{K_{ov}}{k_r} \cdot Da = \frac{k_r\varphi_s \cdot L_{riser}}{U} \]  

The gas-solid contact efficiency \( \gamma_{pf} \) could be computed at different axial increments of \( \Delta z \), by quantifying the contact efficiency as the ratio between the apparent conversion rate and the conversion obtained when a 1D plug flow model is assumed, ignoring any heterogeneity.

In this paper, we perform CFD-DEM simulations to quantify the instantaneous cluster-level gas-solid contact efficiency, which is the ratio between the gas ozone mass fraction inside a cluster region and an average gas mass fraction that will be precisely defined below.

As previously reported clusters are defined as connected regions with local solids fractions exceeding 0.2 everywhere that have a minimum (projected) area of 60 mm\(^2\) and a dense core with at least one grid cell with \( \varphi_s > 0.4 \). The minimum area requirement limits the amount of noise in our measurements that would be caused by the frequent appearance and
disappearance of small clusters. The area of 60 mm$^2$ corresponds to an equivalent circle diameter of 8 mm. The detection of clusters was performed by post-processing simulation data by means of a Matlab® script. In Figure 2, we show a snapshot of some particle clusters obtained from a typical CFD-DEM simulation. The ozone mass fraction of red-colored cells are averaged in order to compute the ozone mass fraction of a cluster as follows:

$$w_{A,cluster} = \frac{\sum_{n=1}^{N} (1 - \varphi_{s,n}) \cdot w_{A,n}}{\sum_{n=1}^{N} (1 - \varphi_{s,n})}$$ (17)

where $w_{A,n}$ is the ozone mass fraction in cell $n$ that is part of the cluster under consideration, $\varphi_{s,n}$ is the solids volume fraction in that cell and $N$ is the total number of cells that are occupied by that particular cluster.

This average concentration inside the cluster is compared to:

$$\bar{w}_A = \frac{\sum_{k=1}^{K} n_k \cdot \bar{w}_{A,k}}{\sum_{k=1}^{K} n_k}$$ (18)

where $\bar{w}_{A,k}$ is the cross-sectional-averaged ozone mass fraction of slice $k$, where $n_k$ is the
number of cells of the $k^{th}$ slice that are occupied by the cluster under consideration. The cross-sectional averaged ozone mass fraction, $\bar{w}_{A,k}$, is computed by excluding cells that are identified as part of a cluster. These cross-sectional-averages are thus weighted by the number of cells that the cluster occupies at each cell row. Thus if a cluster consist of 3 cells at $k^{th}$ row and 1 cell at $(k-1)^{th}$ row, the bulk gas mass fraction (averaged value of those cells that are not occupied by a cluster at that particular row of cells) of cell row $k^{th}$ is weighted 3 times over the bulk gas mass fraction of cell row $(k-1)^{th}$, producing a unique value of the bulk gas ozone mass fraction.

The contact efficiency is defined as the ratio between the gas ozone mass fraction inside the cluster $w_{A,cluster}$ and the average cross-sectional gas ozone mass fraction of the bulk gas, $\bar{w}_A$:

$$\gamma_{cl} = \frac{w_{A,cluster}}{\bar{w}_A} \quad (19)$$

Efficiency values very close to 1 can then be interpreted as highly efficient contacting where all catalyst particles are fully exposed to the bulk gas concentrations and gas diffusion through the particle cluster is much faster than the intrinsic reaction rate. Conversely, numbers very close to zero indicate poor gas solid contacting either due to diffusional limitations or gas bypassing around the clusters. It is worth to mention that $\gamma_{pf}$ term can be interpreted as the ratio of external surface area of the catalyst that is exposed to the gas phase; while $\gamma_{cl}$ represents the ratio of external surface area of cluster particles that are exposed to the bulk gas phase. However, these two terms are not comparable since they belong to two different interpretations. The gas-solid contact efficiency $\gamma_{cl}$, is a parameter that quantifies the instantaneous gas bypassing around particle clusters and differs from $\gamma_{pf}$, which measures the deviation from a steady state 1D ideal plug flow model. Thus, if the dominant mass transfer resistances are found to be at the particle level, it is expected that $\gamma_{pf} \ll 1$, while $\gamma_{cl} \approx 1$. The cluster-level contact efficiency can then employed to identify well the level at which the mass transfer resistance lies.
Results and discussion

In this section, we first show some results of mass transfer coefficients at different operating conditions and values of $k_r$. In this case, a 1D plug flow model is assumed to compute mass transfer coefficients from time-averaged ozone gas mass fraction profiles using the CFD-DEM generated axial solids distribution. The aim is to show that low Sherwood numbers are obtained when these assumptions are made for riser flows.

In the next subsections, one of our main objectives is to identify and quantify the influence of different mass transfer mechanisms on the performance of a riser reactor when clusters are present. The influence of reduced slip velocity, axial gas dispersion and gas bypassing are evaluated. In addition, the influence of the reaction rate and cluster phenomena on the riser performance are quantified and analyzed.

Concentration profiles - 1D Plug flow model

In Figure 3, time-averaged ozone gas mass fraction profiles at several gas superficial velocities are shown. It can be observed that at the bottom region of the riser, higher conversion rates are found. This is because a dense bottom region exists. It has to be noted that in riser flows, there is a trade-off between catalyst holdup and cluster formation. Although high gas superficial velocities can lead to more homogeneous systems (less clustering) both the solids inventory and the gas phase residence time drop. This gives a lower conversion rate at higher superficial velocities as we can see in Figure 3.

To compute a global mass transfer coefficient for each one of these cases, a plug flow model can be assumed (see eqn. 14). The values of $K_{ov}$ were solved through linear regressions above heights of $z = 0.2$ m, where a constant decaying trend of the ozone gas mass fraction profiles was obtained. The values are provided in Table 3.

When one assumes that there is external mass transfer resistance only at the particle level, a global resistance analysis can be used to decompose an overall mass transfer coefficient for
The computed overall mass transfer coefficients are listed in Table 3 for different values of the superficial gas velocity. It can be seen that the overall mass transfer coefficient increases with increasing gas superficial velocity. It can be noticed that at $U$ values exceeding 5.95 m/s, mass transfer rates ($k_{mt} \cdot a_v$) are of similar order of magnitude than the reaction rate ($k_r = 100 \text{ s}^{-1}$). It is clear that the hydrodynamic resistances play an important role even at high superficial velocities. At higher superficial velocities the mass transfer rates increase. This is consistent with an assumption of external mass transfer limitations at the particle level. Note, however, that with increasing superficial velocity also the size and amount of particle clusters change. In our previous study, it was shown that the formation of clusters is highly influenced by the operating conditions, as well as cluster-related properties such as size and aspect ratio.\textsuperscript{53} Therefore (part of) the dependency of $K_{ov}$ versus superficial velocity

$$\frac{1}{K_{ov}} = \frac{1}{k_{mt}a_v} + \frac{1}{k_r}$$

where $a_v$ is the specific particle surface area $a_v = 6/d_p$.

Figure 3: Axial profiles of time-averaged ozone gas mass fraction at $k_r=100 \text{ s}^{-1}$. 

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\textsuperscript{53}
might actually be indirect, i.e., due to changing characteristics of clusters.

Table 3: Mass transfer coefficients. $U$ influence at $k_r = 100 \, \text{s}^{-1}$.

| $U$ (m/s) | $K_{ov} (\text{s}^{-1})$ | $\gamma_{pf} = K_{ov}/k_r$ | $k_{mt} \cdot a_v (\text{s}^{-1})$ | $k_{mt} (\text{m/s})$ |
|-----------|--------------------------|-----------------------------|-----------------------------------|-----------------------|
| 5.16      | 35.48                    | 0.35                        | 56.0                              | 0.0079                |
| 5.55      | 42.69                    | 0.43                        | 74.5                              | 0.0106                |
| 5.95      | 65.09                    | 0.65                        | 186.5                             | 0.0264                |
| 6.35      | 81.18                    | 0.81                        | 431.6                             | 0.0611                |
| 6.74      | 85.75                    | 0.86                        | 601.8                             | 0.0853                |

Figure 4: Axial profiles of time-averaged ozone gas mass fraction at $U = 5.95 \, \text{m/s}$.

In Figure 4, the axial profiles of the time-averaged ozone gas mass fraction profiles at different reaction rates are shown. In Table 4, overall mass transfer coefficients obtained from these profiles at different reaction rates are shown. As expected, the conversion rate increases at higher values of $k_r$.

The ‘plug flow contact efficiency’, $\gamma_{pf}$, decreases at higher reaction rates. As expected, the kinetic resistance decreases, with increasing reaction rates, while the hydrodynamic resistance becomes more dominant. However, the dependence of $k_{mt}a_v$ on the reaction rate,
Table 4: Mass transfer coefficient. Kinetic constant influence at $U = 5.95 \text{ m/s}$.

| $k_r \,(\text{s}^{-1})$ | $K_{ov} \,(\text{s}^{-1})$ | $\gamma_{pf} = K_{ov}/k_r$ | $k_{mt} \cdot a_v \,(\text{s}^{-1})$ | $k_{mt} \,(\text{m/s})$ |
|-------------------------|-------------------------|-------------------------|-------------------------------|-------------------|
| 10                      | 8.04                    | 0.80                    | 41.2                          | 0.0058            |
| 50                      | 36.17                   | 0.72                    | 130.9                         | 0.0185            |
| 100                     | 65.09                   | 0.65                    | 186.5                         | 0.0264            |
| 200                     | 115.94                  | 0.58                    | 275.9                         | 0.0391            |
| 400                     | 196.67                  | 0.49                    | 386.9                         | 0.0548            |
| 1000                    | 367.58                  | 0.37                    | 581.2                         | 0.0823            |

$k_r$, is inconsistent with the assumption of external mass transfer limitations at the particle level. If $k_{mt}a_v$ represented the external mass transfer limitation at the particle level one would expect it to remain constant when only $k_r$ is changed. One can draw a more general conclusion, namely, that the resistances in series analysis of eqn. (17) where $k_{mt}a_v$ is an external mass transfer resistance at whatever level is not valid here.

There can be several mass transfer mechanisms that cause the under-performance of the riser reactors. Falling clusters close to the walls may reduce local slip velocities, leading to lower particle-based Reynolds numbers and thus decreasing local mass transfer coefficients (through Gunn’s correlation). Or maybe the bulk gas stream may circumvent dense particle regions (clusters), leaving the system without contacting with all particles. Gas back-mixing effects can also impede the reactor performance. All these hypotheses are assessed in the next sections.

**Slip velocity**

We will first focus on the particle-level mass transfer resistance. The particle-level mass transfer coefficient depends on the slip velocity of the particle with respect to the surrounding gas. For larger slip velocities the mass boundary layers around particles are thinner which leads to increased mass transfer rates. This dependence is captured by mass transfer
correlations such as the one reported by Gunn, where increased slip velocities lead to larger particle Reynolds numbers and consequently larger interphase mass transfer coefficients. As Helland et al. (2000) suggested, a falling cluster exerts a local reaction force (via two-way coupling) on the gas phase, decelerating the gas motion, such that it could even follow the cluster trajectory. This phenomenon leads to a local drop of the slip velocity and therefore a lower local mass transfer coefficient.

In order to analyze the influence of the slip velocity on the computed local mass transfer coefficient (via the Gunn correlation), the particle-averaged mass transfer coefficient is computed in each computational cell. In this way, we can evaluate whether the drop in the computed particle mass transfer coefficient is the main source of lowered riser reactor performance.

Figure 5: Particle-averaged mass transfer coefficient from a CFD-DEM simulation of 40 seconds. $U = 5.95$ m/s.

In Figure 5, it can be seen that the particle-averaged mass transfer coefficient is significantly lower close to the walls than in the core of the riser. This confirms that cluster
formation leads to lower particle-level mass transfer rates due to a drop in the particle-based Reynolds number.

In Figure 6, the probability distribution function (pdf) of the instantaneous particle Reynolds at different gas superficial velocities is shown. It can be seen that these profiles describe bimodal data distributions. There is a high-peak at rather small Reynolds numbers (5-20), while at high Reynolds numbers a peak appears which grows and shifts to the right with increasing superficial gas velocity. These profiles are consistent with a “slow-moving” solid phase, which can be characterized by particles that are immersed in dense areas where the local slip velocities are low; and a “fast-moving” solid phase that could be characterized by particles located in dilute areas where the slip velocities are relatively large.

This pattern can also be observed in Figure 7, where the probability density distribution of the particle-based mass transfer coefficient at the same operating conditions are plotted. We see in Figure 7, that the pdf’s of the instantaneous particle mass transfer coefficient describe bimodal data distributions as well. It should be noted that the Gunn correlation has a strong dependence on the particle-based Reynolds number and the solids volume fraction. At similar values of the particle-based Reynolds number, dilute areas acquire lower mass transfer coefficients than dense regions. At higher gas superficial velocities, clusters are less likely to form and slip velocities increase (see Figure 6). It is then expected that the occurrence probability of the dense phase decreases as well. So, looking at Figure 7, we can state that dense particle regions are characterized by slow motion and low local mass transfer coefficients, while the dilute solid phase is characterized by high slip velocities and relatively high mass transfer coefficients.

In this subsection, we have confirmed that cluster formation leads to a lower particle-level mass transfer coefficient. As can be seen from Figure 7 shows, the mean particle mass transfer coefficient of a dense solid phase ranges between 0.22 and 0.27 m/s, while the mean value of the same property for the dilute solid phase ranges between 0.43 and 0.49 m/s.

When comparing the particle-level mass transfer coefficient measured here with the val-
Figure 6: Probability density distribution of instantaneous particle-based mass transfer coefficient.

Figure 7: Probability density distribution of instantaneous particle-based mass transfer coefficient.

ues reported in Tables 3 and 4 we clearly see that the actual particle-level mass transfer coefficients are much larger than those in the tables. The values reported in the tables are obtained by assuming that the dominant mass transfer resistance is at the particle-level. The
disagreement shows that this assumption is incorrect. We conclude that, while the presence of clusters significantly influences the particle-level mass transfer coefficients, the dominant mass transfer resistance is not at the particle level.

**Axial gas dispersion coefficient**

Axial dispersion might lead to a lower apparent $K_{ov}$ when experiments are interpreted using a plug-flow model without axial dispersion. Some of the measured low Sherwood numbers might be explained by this type of ‘misinterpretation’. Therefore the influence of the gas axial dispersion is evaluated in this subsection. In order to analyze whether gas dispersion effects play a major role in these deviations, a 1D convection dispersion equation can be employed to compute the apparent reaction rate in the riser reactor. Given the time-averaged axial ozone mass fraction profile, the influence of the axial gas dispersion coefficient can be determined. Changes in $D_{ax}$ are evaluated in order to analyze the deviations between the attained mass fraction profiles and those obtained by assuming a steady state axially dispersed plug flow model.

\[
0 = -\frac{\partial U w_{A,g}}{\partial z} + D_{ax} \frac{\partial^2 w_{A,g}}{\partial z^2} - K_{ov} \langle \phi_s \rangle w_{A,g} \tag{21}
\]

For $D_{ax} \ll U^2 / (K_{ov} \langle \phi_s \rangle)$ it has solutions: $\lambda_1 \approx -(K_{ov} \langle \phi_s \rangle)/U$, $\lambda_2 \approx U / D_{ax}$. Since $\lambda_2 \gg -\lambda_1$, $\lambda_2$ corresponds to a shorter length scale. In fact, this second solution (with positive exponential factor $\lambda_2$) only influences the mass fraction near the exit of the column. That is, the contribution of $\exp(\lambda_2 z)$ will be only significant near the exit and is sensitive to the outlet boundary condition. Away from the exit only a single exponent solution, namely $\exp(\lambda_1 z)$, is relevant and therefore $w_{A,g}$ is expected to decay exponentially in this region. This means that $\lambda_1 z$ can be locally estimated using: $\lambda_1 \Delta z = \ln\langle w_{A,z+\Delta z} \rangle / \langle w_{A,z} \rangle$, such that $K_{ov}$ can be obtained from the characteristic equation (22) as:

\[
K_{ov}(z) = -\frac{1}{\langle \phi_s \rangle \Delta z} \ln \left( \frac{\langle w_{A,z+\Delta z} \rangle}{\langle w_{A,z} \rangle} \right) \left[ U - \frac{D_{ax}}{\Delta z} \ln \frac{\langle w_{A,z+\Delta z} \rangle}{\langle w_{A,z} \rangle} \right] \tag{22}
\]
The axial gas dispersion coefficient has been determined in CFD-DEM, by injecting a pure ozone gas pulse of 0.01 seconds over steady state simulations in an inert environment (no chemical reaction). The gas velocity fluctuations are expected to be larger in simulations with higher degree of clustering (low $U$). In Figure 8, the obtained gas residence time, $E(t)$, for $U = 5.16$ m/s is plotted. The axial dispersion coefficient was computed by means of equation (23):

$$D_{ax} = \frac{U \cdot L \int_0^\infty E(t) (t - t_m)^2 \, dt}{2t_m^2 \epsilon_{bed}}$$

where $t_m = \int_0^\infty t \cdot E(t) \, dt$ is the mean gas residence time, $L$ is the riser length, $\epsilon_{bed}$ is the bed porosity which amounted to 0.943, and $U$ is the gas superficial velocity.

![Figure 8: Gas residence time distribution at $U = 5.16$ m/s.](image)

The mean gas residence time was around $0.307 \pm 0.11$ seconds and the axial dispersion coefficient amounted to $D_{ax} = 0.527$ m$^2$/s. By feeding this input parameter into the previous convection-dispersion equation we can obtain the order of magnitude of the deviation in $K_{ov}$ ($k_r = 1000$ s$^{-1}$) when axial gas dispersion effects are accounted for in the interpretation model. In Figure 9, it can be seen that the change due to axial dispersion is small. So, discarding gas axial dispersion in an interpretation model is not a major cause of overesti-
mation of the conversion rate. For the case that dispersion only has a limited influence on the
determined mass transfer coefficient we find that the relative contribution is approximately
$(D_{ax}\langle \phi_s \rangle K_{ov})/U^2$. For the measured dispersion coefficient this gives a 13% deviation (with
$\langle \phi_s \rangle=0.0535$). So in this case, gas backmixing is not a major cause for (apparent) mass
transfer limitations on riser reactor performance.

**Gas solid contact efficiency**

Gas solid contact efficiency is related to gas bypassing: some of the reactant will have an
intimate contact with the catalyst particles and the rest may leave the system chemically
unchanged due to a very poor exposure to the particulate phase. In Figure 10, we present
an illustrative snapshot when gas bypassing occurs.

On the left-hand side of Figure 10, particles are colored according to their respective
ozone mass fraction. These values can be assumed as the ozone mass fraction at the surface
of each particle. On the right-hand side of the figure, the gas velocity field is shown. It
can be seen that the gas flows at high velocities in the core of the pseudo 2D riser. Red-
colored particles (rich in ozone content) are mainly encountered in exposed regions to the main bulk stream, where the velocities are higher. We can see that cluster regions are mostly composed of blue-colored particles that possess low ozone content presumably due to higher gas residence times or trapped gas pockets inside the clusters. Actually, Ouyang et al.\textsuperscript{51} suggested that falling particle clusters could capture and retain gas and these observations confirm this suggestion.

The gas solid contact efficiencies have been computed for several values of the reaction constant, $k_r$, at the same operating conditions ($U = 5.95$ m/s) and for different gas superficial velocities at a fixed $k_r = 100$ s$^{-1}$ to analyze the influence of cluster characteristics on the gas solid contacting.
Reaction rate effect

In Figure 11, the pdf’s of the cluster contact efficiency of all these simulation cases are shown. It can be noticed that at lowest kinetic constant $k_r = 10 \text{ s}^{-1}$, CFD-DEM predicts the major part of occurrences have contact efficiency values ranged between 0.8 and 1. So, assuming ideal plug flow would in this case be a reasonable assumption if we want to estimate the riser reactor performance. However, when the reaction rate is increased, larger errors result. For instance, if a significant improvement is made on a catalyst by increasing its activity with corresponding change in $k_r$ from 10 to 1000 $\text{s}^{-1}$, a plug flow model assumption can lead to larger over-estimations, since the gas-solid contact efficiency would be much lower than at low reaction rates (see Figure 11).

Figure 11: Probability density distribution of gas-solid contact efficiency at different $k_r$.

In Figure 12, the cluster-averaged contact efficiency for each simulation is plotted, where the error flags represent the confidence intervals of the 68.2 % ($\gamma_{cl} \pm \sigma = \sqrt{(\bar{\gamma}_{cl}-\gamma_{cl})^2} / N_{clusters}$) of the cluster contact efficiency data. It can be seen that it drops at higher values of the kinetic constant as previously stated by other authors.$^{16,42}$
In a previous study\textsuperscript{53} it was shown that complex clustering phenomena can be well predicted by means of CFD-DEM. In risers, the total cluster population increases at low gas superficial velocities.\textsuperscript{64} Larger populations of falling clusters close to the walls can retain gas pockets of highly depleted reactant,\textsuperscript{51} leading to inefficient gas-solid contacting. At higher gas superficial velocities, the system becomes more dilute and the particle shielding effect does not become that influential as Figure 13 reveals (see $U = 6.74$ m/s line). It is noticed then that clustering and consequently operating conditions play a major role on the performance of a riser reactor (see Figure 13).

These results show that clustering phenomena are a major cause of inefficient contacting. From Figure 14, it can be noticed that the cluster-averaged contact efficiency significantly increases at higher gas superficial velocities. Thus, the measurement of global mass transfer coefficients requires and accurate estimate of cluster-related properties. This seems to be the cause of so much disagreement between global Sherwood number data. In systems, where clustering and particle shielding phenomena are very pronounced; or in systems in which the reaction rate is very high; the global Sherwood number will tend to zero.
Figure 13: Probability density distribution of cluster contact efficiency at several gas superficial velocities at $k_r = 100 \text{ s}^{-1}$.

Figure 14: Cluster-averaged contact efficiency at several gas superficial velocities at $k_r = 100 \text{ s}^{-1}$.

**Influence of dilution ratio**

In this subsection, we present gas-solid contact efficiency results of CFD-DEM simulation at different dilution ratios of active particles (number of active over total number of parti-
Diluted fluidized systems have been employed in the past to measure mass transfer coefficients. Active spheres can be mixed with inert ones to experimentally measure mass transfer coefficients. In CFD-DEM all particles are numbered and tracked. By means of a simple algorithm, a fixed number of particles could be labelled as active or inert. Each particle label was permanent for the whole simulation and the particles were assumed to be homogeneously mixed in the system.

![Figure 15: Contact efficiency pdf at $U = 5.55 \text{ m/s}$ and $k_r = 100 \text{ s}^{-1}$.](image)

It should be noted that the Gunn correlation was also utilized in these simulations to compute the particle-based Sherwood number. Mass transfer correlations for dilute particle systems and Gunn correlation differ in the asymptotic behavior at low Reynolds number ($2 \cdot \epsilon/\tau$ and 2, respectively). These differences in the diffusional contribution of the Sh number were negligible for this set of simulations. The reasons are that first the mass transfer at the particle level is not limiting especially for particles inside clusters. Second, for particles outside clusters the higher Reynolds number contribution to the particle-level Sherwood number is relevant.

In Figure 15, the probability density distribution of cluster gas-solid contact efficiency at different dilution ratios are shown. We can see that the gas-solid contact efficiency is higher
Figure 16: Cluster-averaged contact efficiency.

at increasing dilution ratios. At a fixed catalyst activity ($k_r = 100 \text{ s}^{-1}$), lower dilution rates (more active spheres) will lead to more severe particle shielding effects when clusters are formed.\textsuperscript{16}

Table 5: Mass transfer coefficient. Plug flow model. Kinetic constant influence at $U = 5.95 \text{ m/s}$.

| Simulation | $k_r$ (s$^{-1}$) | % active particles | $U$ (m/s) |
|------------|---------------|-------------------|---------|
| 1          | 100           | 10                | 5.55    |
| 2          | 10            | 100               | 5.55    |
| 3          | 100           | 50                | 5.55    |
| 4          | 50            | 100               | 5.55    |
| 5          | 100           | 90                | 5.55    |
| 6          | 90            | 100               | 5.55    |

As expected, Figure 16 shows that the cluster-averaged contact efficiency drops at decreasing dilution ratio.

It can be seen that an increase of the dilution ratio effect is comparable to an increased
catalyst activity (see Figure 11). The gas inside the cluster becomes more depleted of reactant, consequently the average gas reactant concentration is lower, leading to poorer gas-solid contact efficiencies.

In diluted systems, if the active particles are homogeneously mixed, the performance of a system where a fraction, $\varphi_{\text{active}}$, of the particles are active and $k_r = 100 \, \text{s}^{-1}$ is expected to be similar to that one with $k_r = \varphi_{\text{active}} \cdot 100 \, \text{s}^{-1}$ where all particles are active. To prove this statement, we ran simulations at $U = 5.55 \, \text{m/s}$ at equivalent $k_r$ values, where all the particles are active (see Table 5).

In Figure 17, the contact efficiency pdf of simulations 1 to 6 are plotted (see Table 5). If we compare pdf profiles of simulations 1 and 2, it can be noticed that the effect of the particle dilution ratio is equivalent to the effect of the catalyst activity. The same trend is shown for the remaining simulation pairs. Thus, we confirm previous author’s observations,\textsuperscript{18} namely that mass transfer coefficients obtained from diluted systems should not be comparable to undiluted fluidized systems.

![Figure 17: Contact efficiency pdf at $U = 5.55 \, \text{m/s}$.](image)
Mass transport inside clusters

The situation of mass transfer resistance inside a cluster of reacting particles is qualitatively analogous to internal mass transfer limitation inside a catalytic porous particle. If the analogy also holds beyond a qualitative similarity a type of Thiele modulus could be applicable in order to determine the reaction effectiveness inside clusters and hence permit the development of a cluster-based mass transfer model. In this case a correlation between cluster size and gas-solid contact efficiency should be obtained, regardless of the gas phase velocity. In this section, we will see that this is quite challenging due to the large data scattering that such a correlation shows.

The scatter plot of cluster contact efficiency, $\gamma_{cl}$, against the equivalent cluster diameter $(\frac{2}{\pi} \sqrt{A_{\text{cluster}}})$ in Figure 18 shows no clear correlation between the two quantities. If the gas solid contact efficiency was assumed to depend only on the cluster size, a trend should be visible. Moreover, the same master curve would be expected at different gas superficial velocities. However, it is observed that there is a no clear correlation, especially in denser systems (i.e., at lower $U$ values) where clustering phenomena are more intense.

At higher gas superficial velocities ($U = 6.35$ and $6.74$ m/s) the systems are rather dilute. Here clusters are less likely to interact with each other, and this might be the cause of less data scattering (although this is still quite large as evident from Figure 18).

Thus, it is worthwhile to show the causes of this scattering, and why cluster-based mass transfer models should not merely depend on the equivalent cluster diameter.

In Figure 19, we present a snapshot sequence of two clusters in a relatively dilute region of the riser domain. The gas velocity vector field is superimposed on the porosity field. It can be observed that in the first snapshot, the gas stream does penetrate into the cluster wake of the cluster located at the left. Although this riser section is quite dilute, we can observe (if we follow the sequence) how the gas passes through the smallest cluster located at the top as well. Van der Ham et al. suggested that the increase in the gas solid contact efficiency could be due to the breakup of cluster structures. Although cluster formation leads
Figure 18: Gas solid contact efficiency versus equivalent cluster diameter of 500 random clusters at $k_r = 100 \, \text{s}^{-1}$ a) $U = 5.16 \, \text{m/s}$ b) 5.55 m/s c) 5.95 m/s d) 6.35 m/s e) 6.74 m/s.

To poor gas-solid contacting, we also see that the gas can pass through the cluster structure causing only a change/orientation of the cluster shape without destroying it. We observe that the cluster structure is quite dynamic and can adopt different shapes and orientations in time that can be more susceptible to gas permeation.

In Figure 20, another sequence in a denser region of the riser is shown. In this figure, it can be observed more clearly how the gas stream accelerates due to the high cluster content at the bottom of the riser. Denser regions will not only lead to enhanced bypassing, but also the formation of gas streams with large velocities that can eventually pass through some of the clusters. This phenomenon causes that cluster particles found upstream are more easily accessible to the gas phase and can experience a more efficient gas-particle contact.
Figure 19: Porosity field with the gas velocity vector field superimposed in a dilute region of the riser domain. $U = 5.16$ m/s. Snapshots time frame of 0.01 s.

In general, we observe in our simulations a rather chaotic behavior of particle clusters. They not only form, grow, break up or merge, but also they can adopt different shapes, densities, aspect ratios and orientations. All these phenomena have an effect on the gas-solid efficiency of the cluster itself or/and other neighboring clusters that are found upstream. Although it seems clear that clustering phenomena enhance gas bypassing and poor gas-solid contacting, these phenomena feature such a broad variety of structures that it remains very challenging to develop closures for cluster-based mass transfer models.

Conclusions

In this work we have performed CFD-DEM simulations in order to generate more insight about mass transfer mechanisms that take place under riser flow conditions. The instantaneous cluster-level contact efficiency between the gas phase and cluster particles have been computed at several reaction rates and gas superficial velocities. This work explicitly con-
firms and corroborates suggestions made by other authors,\textsuperscript{16,20,46,51,63,67} namely that particle clusters have a large influence on the gas-solid contact efficiency and on global mass transfer phenomena. We clearly showed that for the system studied here the increased mass transfer resistance is due to the presence of particle clusters and not due to axial dispersion effects, or changes of the particle-level mass transfer coefficient. Moreover, we showed that a decreasing gas superficial velocity leads to lower $\gamma_{cl}$ values. At lower $U$ values, the fact that clusters are larger and a less intense convective mass transfer exists inside these particle structures seem to be the main causes of obtaining such pattern. Besides, increasing reaction rate showed to decrease $\gamma_{cl}$, thus increasing the influence of hydrodynamic resistances at cluster level as other authors suggested.\textsuperscript{21,42} Diluted fluidized systems were found to lead to higher gas-solid contacting rates.\textsuperscript{16,18} For the system studied here it was proved that the dilution rate effect is equivalent to reaction rate effects as Venderbosch et al. suggested.\textsuperscript{16} So, the effect of dilution by inactive particles can be easily understood in terms of an equivalent decrease of the
reaction rate.

Although in literature there is general agreement about the relevance of clusters, it is less clear whether the mass transfer resistance lies in the external mass transfer to the cluster surface;\textsuperscript{23–25} or whether clusters can be assumed as large porous spheres where only diffusional transport takes place.\textsuperscript{20,42} In this work, we have shown that cluster contact efficiency does not correlate well with the cluster size since there is large data scattering. Thus, clusters cannot be assumed as large porous particles, where effective molecular diffusivity is the only mass transport mechanism. Besides, convective mass transfer can play an important role when high $\gamma_{cl}$ values are attained. Convective mass exchange between dilute phase and dense phase exists and it could be enhanced by the formation of gas jets that pass through the cluster structures.

Particle clusters are transient entities that show a broad variety of shapes, sizes and orientations as other authors suggested.\textsuperscript{16,64,68} The large amount of properties that characterize particle clusters, altogether with their location, local density and proximity of high-velocity gas streams can cause interactions of very diverse nature. The scattering pattern of gas solid contact efficiency data, suggests that convective mass transfer inside clusters can be enhanced or limited by all cluster properties previously mentioned, obtaining quite unpredictable behavior if we only analyze a single parameter, e.g. cluster size.

Although we have shown that clusters enhance gas bypassing and result in poor gas-solid contacting, we find it challenging to develop mass transfer closures at cluster scale level. It seems very hard to capture the influence of a cluster using simple parameters such as cluster size. The main reason is that the cluster contact efficiency is very much influenced by convection through the cluster and that this convection depends on the configuration of clusters downstream. Our tentative conclusion is that accurate coarse-graining of the influence of particle clusters is difficult and in fact CFD-DEM is the tool to predict the performance best. Related to the particle-based closures used in CFD-DEM we would like to point out the following behavior. Most mass transfer seems to take place at the boundaries of clusters, where
flow can still partly penetrate the clusters. At these locations the solids volume fractions quickly change. However, the particle-level mass transfer correlations used in CFD-DEM were developed for (locally) homogeneous systems. This raises the question whether the particle-level correlations are accurate enough. Therefore we recommend performing Direct Numerical Simulations of freely evolving clusters to validate local particle-level Sherwood correlations of heterogeneous particle structures.
List of Symbols

Roman symbols

\(A_{cluster}\)  cluster area, m\(^2\)
\(a_p\)  particle surface area, 1/m
\(D\)  Damkohler number, -
\(D_{ax}\)  axial dispersion coefficient, m\(^2\)/s
\(D_{AB}\)  ozone molecular diffusivity in air medium, m\(^2\)/s
\(D_{eff}\)  effective molecular diffusivity, m\(^2\)/s
\(d_p\)  particle diameter, m
\(d_{cl}\)  cluster diameter, m
\(e_{p-p}\)  particle-particle normal restitution coefficient, -
\(e_t\)  particle tangential restitution coefficient, -
\(e_{p-w}\)  particle-wall normal restitution coefficient, -
\(F_c\)  particle collision force, N
\(\langle G_s \rangle\)  time-averaged solids mass flux vector, kg/(m\(^2\)-s)
\(I_p\)  moment of inertia, N-m
\(k_{int}\)  particle-based mass transfer coefficient, (m/s)
\(k_n\)  particle spring stiffness, (N/m)
\(k_r\)  kinetic constant, s\(^{-1}\)
\(m\)  mass, kg
\(P\)  pressure, Pa
\(r\)  position vector, m
\(S_{hp}\)  particle Sherwood number
\(s_p\)  solids displacement vector, m
\(S_p\)  momentum source term, N/m\(^3\)
\(T_p\)  torque, N-m
\(u\)  gas velocity vector, m/s
\(U\)  gas superficial velocity, m/s
\(V_p\)  particle volume, m\(^3\)
\(v_p\)  grid-averaged particle velocity vector, m/s
\(w_{A,g}\)  ozone gas mass fraction at bulk gas phase, kg ozone/kg gas
Greek symbols

\( \gamma_{pf} \)  Gas-solid contact efficiency in a 1D plug flow model, -
\( \gamma_{cl} \)  Instantaneous gas-solid contact efficiency, -
\( \beta \)  interphase momentum transfer coefficient, kg/(m\(^3\).s)
\( \Delta t_{\text{Gas}} \)  gas phase time step, s
\( \Delta t_{\text{DEM}} \)  particle phase time step, s
\( \epsilon \)  porosity m\(^3\) voidage/m\(^3\) reactor
\( \rho_g \)  gas density, kg/m\(^3\)
\( \rho_s \)  solids density, kg/m\(^3\)
\( \mu \)  dynamic viscosity, kg/(m\(\cdot\)s)
\( \mu_{fr} \)  particle friction coefficient, -
\( \varphi_{cl} \)  cluster phase holdup, m\(^3\)cluster/m\(^3\) reactor
\( \varphi_s \)  solids volume fraction, m\(^3\)solid/m\(^3\) reactor
\( \tau \)  stress tensor, Pa
\( \omega_p \)  particle rotational velocity, 1/s

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Supporting Information

Supporting information is available on the model verification of the convection-diffusion equation and on additional literature references regarding studies related to mass transfer in riser flows.
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