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Coupling the fictitious domain and sharp interface methods for the simulation of convective mass transfer around reactive particles: Towards a reactive Sherwood number correlation for dilute systems

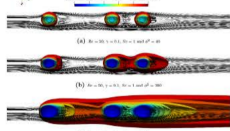
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- DLM/FD SIM numerical method for the simulation of convective mass transfer past reactive obstacles.
- Numerical method validated on multiple test cases.
- New reactive Sherwood number correlation based on external internal coupling.
- New correlation is shown to perform well on a flow configuration representative of a dilute regime.



(c) $Re = 50, \gamma = 0.1, Se = 1 \text{ and } d^2 = x$

We suggest a reactive Sherwood number model for convective mass transfer around reactive particles in a dilute regime. The model is constructed with a simple external internal coupling and is validated with Particle Resolved Simulation (PRS). The PRS of reactive particle fluid systems requires numerical meth ods able to handle efficiently sharp gradients of concentration and potential discontinuities of gradient concentrations at the fluid particle interface. To simulate mass transfer from reactive catalyst beads immersed in a fluid flow, we coupled the Sharp Interface Method (SIM) to a Distributed Lagrange Multiplier/Fictious Domain (DLM/FD) two phase flow solver. We evaluate the accuracy of our numerical method by comparison to analytic solutions and to generic test cases fully resolved by boundary fitted simulations. A previous theoretical model that couples the internal diffusion reaction problem with the external advection diffusion mass transfer in the fluid phase is extended to the configuration of three aligned spherical particles representative of a dilute particle laden flow. Predictions of surface concentra tion, mass transfer coefficient and chemical effectiveness factor of catalyst particles are validated by DLM FD/SIM simulations. We show that the model captures properly the effect of an internal first order chemical reaction on the overall respective reactive Sherwood number of each sphere depending on their relative positions. The proposed correlation for the reactive Sherwood number is based on an existing non reactive Sherwood number correlation. The model can be later used in Euler/Lagrange or Euler/ Euler modelling of dilute reactive particle laden flows.

Keywords: Sharp Interface Method Catalyst particle Mass transfer Sherwood number Chemical reaction Thiele modulus

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1. Introduction

Systems involving interactions of a dispersed solid phase with a continuous fluid phase through momentum, heat and mass trans fer are ubiquitous in a wide range of industrial and energetic pro cesses. A classical process in the chemical engineering industry is catalytic cracking in a reactor. If the dispersed solid phase, com monly referred to as particles, is fixed in the reactor, the system is known as a fixed bed and the flow is relevant of the flow through a porous medium represented by the network of randomly stacked catalytic particles (Furuta et al., 2006). If particles are mobile, the system is known as a fluidized bed (Gidaspow, 1994; Montero et al., 2018). In both reactors, the fluid enters the system with an imposed concentration of reactants and reactants are transferred from fluid phase bulk to catalytic particles bulk through diffusion, where chemical reactions take place in the form of heteroge neously catalyzed gas or liquid reaction. The modeling, operation, design and optimization of these systems necessitate an advanced comprehension of the coupling among the dominant transfer phe nomena, namely momentum, heat, and mass transfer, that are usu ally associated to the presence of chemical reactions. For decades, operating experimental setups and deriving simplified analytical solutions were the two only ways to improve the comprehension of these systems. With the emergence of robust, accurate and com putationally efficient numerical approaches/methods, we can com plement and extend our comprehension with, e.g., reliable information about the micro scale interactions in these systems that are not accessible through experiments or theory.

Over the past two decades, diverse Computational Fluid Dynamics (CFD) approaches/methods for the simulation of systems involving fluid/solid interactions have been developed. Combined to the increasing power of supercomputers that now enables one to perform Direct Numerical Simulations (DNS) at the scale of par ticles, CFD tools are now capable of supplying reliable and high quality detailed data in the flow. In this class of highly resolved CFD methods, Particle Resolved Simulation (PRS) has arisen as a mature method able to provide reliable local information about momentum, heat and mass transfer at the particle scale in partic ulate flows (Sun et al., 2016 among many others). PRS methods can be classified into two categories:

- (i) The first category comprises body fitted mesh methods. The advantage of this type of method relies on the ease to enforce boundary conditions at the particle surface, i.e., at the fluid/particle interface. This technique has been used to investigate transport properties in a bed made of multiple fixed particles by Romkes et al. (2003) and Augier et al. (2010a) and convective heat/mass transfer over a single par ticle by Feng and Michaelides (2000). It has also been employed for moving particles by Hu et al. (2001) and mov ing boundaries by Duarte et al. (2004).
- (ii) The second category comprises fixed mesh methods. The challenge of this type of method is the difficulty in enforcing the correct boundary conditions at the particle surface while the main advantage is the use of a regular Cartesian grid. The Immersed Boundary Method (IBM) uses Lagrangian markers at the particle surface to impose boundary conditions and introduces an additional forcing term (Uhlmann, 2005). Xia et al. (2014) applied IBM to study convective heat/mass transfer over a single particle. IBM was also used to evaluate the heat transfer Nusselt number in dense particulate flow systems by Deen et al. (2014) and Sun et al. (2015). Both studies compared their results to the pioneering experimen tal work of Gunn (1978) and proposed corrections of Gunn's correlation based on their data sets. Recently, IBM has been used by Lu et al. (2018) to examine mass transfer in partic

ulate flows with surface reaction. The Lattice Boltzmann Method (LBM) is another computational method that uses a fixed mesh. LBM has also been applied to particulate flows with heat transfer by Khiabani et al. (2010) and Kruggel Emden et al. (2016). Unlike conventional discretization schemes that solve the classical conservation equations, LBM solves convection collision steps of probability density functions. Finally, the Distributed Lagrange Multiplier/Ficti tious Domain Method (DLM/FD), firstly introduced by Glowinski et al. (2001), combines the particle and fluid equations of motion into a single, weak, and general equa tion of motion called combined momentum equation. The combined equations are derived through the combined velocity space incorporating the rigid body motion (no slip) in the particle. The DLM/FD method has been extended to treat heat/mass transfer by Yu et al. (2006), Dan and Wachs (2010) and Wachs (2011).

Apart from Yu et al. (2006) that considered diffusion inside par ticles, the common feature of the aforementioned fixed mesh methods is that they solve the convection diffusion equation by enforcing a uniform temperature (or concentration) in the whole solid particle volume. This type of method is inadequate for the treatment of systems where temperature/concentration gradients are prominent within the particle. In this case, the convection diffusion (or convection diffusion reaction) equation must be solved in both domains, i.e., in the solid domain and in the fluid domain. Augier et al. (2010b) used a Volume of Fluid method (VOF) to study the efficiency of partially wetted stacked catalyst particles. Catalyst efficiency was studied for different particles shapes as a function of Thiele modulus. Augier et al. (2010b) solved the diffusion reaction in the solid domain only with different Dirichlet boundary conditions for the concentration on the particle surface depending on whether the surface is locally wetted or non wetted (as given by the VOF field). Haroun et al. (2010) employed a VOF method to study interfacial reactive mass transfer in two phase flows. The VOF field, i.e., the phase concentration field, was then used in the Finite Volume solution of the concentration equa tion such that properties in cells cut by the interface are simple phase concentration weighted averages of the properties in the two phases. Although efficient, the Finite Volume method with simple phase concentration Weighted Average of properties, thereafter called FVWA, necessitates a highly refined mesh at the interface in order to accurately enforce the correct boundary con ditions, which may render, for a fixed Cartesian mesh, the method computationally inefficient due to the huge computational cost. The main drawback of the FVWA method that it does not capture discontinuities but instead smoothes them out.

The Sharp Interface Method (SIM), also referred to as Ghost Fluid Method (GFM), is a fixed mesh numerical method used to accurately capture boundary conditions with discontinuities along embedded interfaces (Shi et al., 2011). In the SIM, jump conditions are incorporated in the discretization of the differential operators on the Cartesian grid in the vicinity of the interface. SIM was firstly introduced by Fedkiw et al. (1999) and employed to impose boundary conditions at a contact discontinuity in the inviscid Euler equations. SIM was later extended to treat more general disconti nuities by Liu et al. (2000). In their work, Liu et al. (2000) devel oped a version of SIM to address the problem of inhomogeneous Poisson equation in the presence of interfaces. The method is easy to implement in three dimensions and the matrix associated to the discrete Laplacian operator remains symmetric, allowing to use 'black box' scientific libraries to solve the corresponding linear sys tem. Gibou et al. (2002) considered the Poisson equation with a non uniform coefficient and Dirichlet boundary conditions on an irregular domain and showed that a second order accuracy can be obtained with a simple discretization scheme that also pre serves the matrix symmetry. Udaykumar and Mao (2002) and Gibou et al. (2003) employed the SIM to track the evolution of solidification fronts in the presence of heat and solute transport in the dendritic solidification of aqueous salt solutions. Kang et al. (2000) extended the SIM to treat multiphase incompressible flows including effects of viscosity, surface tension and gravity and applied the method to two phase water air mixtures. Marella et al. (2005) used a SIM to simulate immersed boundary problems while Liu et al. (2005) simulated droplet interactions with objects of dif ferent shapes using a SIM that accounts for surface tension and vis cosity jumps. Kapahi et al. (2013) used a modified SIM to treat interfaces of embedded objects, with an application to shock wave particulate flows. Finally, to end this non exhaustive list of SIM related works, Shao et al. (2012) combined the SIM with a Fic titious Domain method to simulate heat transfer in particulate flows with heat diffusion inside particles.

PRS is very powerful to supply high quality data inside the flow but is still limited to up to a few thousands, at best a few tens of thousands, of particles due to the large computing cost of these simulations. Finely resolved simulations as PRS can easily comprise hundreds of millions, and even a few billions, of grid cells, that rep resent highly challenging parallel computing problems, whether on multi CPU or the emerging GPU/multi GPU technology. Thereby, from a modeling viewpoint, there has been a sustained appeal to combine these particle scale models to larger scale mod els, namely Euler/Lagrange at the so called meso scale and Eulere/ Euler at the so called macro scale. The conceptual features of a fully integrated multi scale modeling of particle laden flows is comprehensively described by Deen et al. (2014). The vast majority of the multi scale approaches suggested in the field of particle laden flows assume a bottom up strategy or phrased in a more emphatic way an upwards cascade of knowledge. In short, what is learnt through highly resolved simulations at the micro scale on small representative systems is meant to be transferred to higher scale models and to contribute to a deeper understanding of the particle laden flow dynamics. Among the assorted ways to transfer knowledge, the most popular way has undoubtedly been over the last 15 years to enhance existing correlations for dimen sionless numbers representative of momentum, heat or mass transfer. Among many others, Deen et al. (2014) and Sun et al. (2015) suggested corrections to enhance the Nusselt number cor relation suggested by Gunn (1978) 30 years earlier. Our objective in this paper follows the same line, i.e., to suggest a new correlation or to enhance an existing correlation derived from micro scale PRS that can be later used in meso scale Euler/Lagrange and macro scale Euler/Euler modelling.

In the present study, we combine a Fictitious Domain method and a first order Sharp Interface method to investigate mass trans fer in particulate flows in the presence of a first order reaction inside particles. The combined numerical methods constitute a general framework capable of simulating fixed and moving parti cles. The present study is a part of a series of consecutive compu tational studies of reactive particulate flows. We investigated the problem of a single fixed particle where we used a boundary fitted method in Sulaiman et al. (2018), now examine in this paper the problem of multiple aligned particles with a non boundary fitted method, and then will use this non boundary fitted method in future works to examine a random array of fixed particles and freely moving particles in a fluidized bed. The 3D simulation results supplied by this computational method in a flow configura tion representative of dilute particle laden flows are used to assess the validity of a reactive Sherwood number correlation that we already suggested for the case of a single sphere in an unbounded domain in Sulaiman et al. (2018) and that we revisit later in this work. We intend to show that a reactive Sherwood number corre

lation can be constructed based on simple external internal cou pling. The obtained model relies on any non reactive Sherwood number correlation available in the literature and is deemed to perform well. The rest of the paper is organized as follows. We shortly summarized in Section 2 the equations and the correspond ing dimensionless numbers governing the problem. Section 3 elab orates on the features of our numerical method that combines a DLM/FD method to compute the flow field and a SIM to compute the chemical species concentration field. Then, we present in Sec tion 4 a series of validation tests of growing complexity involving a single sphere (Remark: readers familiar with the adopted numeri cal methods and the classical validation tests presented in Sections 3 and 4, respectively, may safely skip these two sections and move from Section 2 right away to Section 5). We start with diffusion, then move on with diffusion reaction, convection diffusion, and finally consider convection diffusion reaction. In the last case, and due to the lack of analytical solutions and previously estab lished correlations to compare our numerical results to, we per form a comparison between SIM and a body fitted method that fully resolves the interface to investigate the efficiency of our SIM method and determine its limits of validity in terms of range of dimensionless parameters considered and grid size, through a series of convergence tests. We also show, through comparison in a diffusion reaction case, the advantage of SIM over FVWA. Finally, and this is the core of this work, we investigate in Section 5 the problem of three interacting spheres, firstly presented by Ramachandran et al. (1989) for heat transfer without diffusion and chemical reaction inside the spheres, with mass transfer cou pled to diffusion and chemical reaction inside the spheres. We test our reactive Sherwood number correlation on this configuration. The validity of the model and the steps forward in extending/ improving our reactive Sherwood number correlation are dis cussed in Section 6.

2. Governing equations

We aim at solving the time dependent and incompressible flow of a Newtonian fluid past multiple fixed obstacles with mass trans fer between the fluid and the solid obstacles. We define the full flow domain as Ω , the part of Ω occupied by the solid obstacles as *P* and the part of Ω occupied by the fluid as $\Omega \setminus P$. The problem is governed by the following conservation equations: fluid mass conservation, fluid momentum conservation and chemical species conservation. Here we assume a single chemical species C at a low concentration in the fluid such that it does not affect the constant density and viscosity of the fluid. Dimensional quantities are dis tinguished from dimensionless quantities by a "*" superscript. We denote u^* the fluid velocity, p^* the fluid pressure, C_f^* the chem ical species concentration in the fluid and C_s^* the chemical species concentration in the solid. The chemical species is assumed to undergo a first order reaction in the solid obstacles. With appropri ate initial conditions in Ω on $(\boldsymbol{u}^*, C_f^*, C_s^*)$ and boundary conditions on $\partial \Omega$, the boundary of Ω , on \boldsymbol{u}^* (and potentially on p^*), the set of conservation equations together with fluid/solid interface condi tions is written as follows:

• in the fluid

$$\rho_f^* \left(\frac{\partial \boldsymbol{u}^*}{\partial t^*} + (\boldsymbol{u}^* \cdot \nabla) \boldsymbol{u}^* \right) \quad \eta^* \nabla^2 \boldsymbol{u}^* + \nabla p^* \quad \mathbf{0}, \tag{1}$$

$$\nabla \cdot \boldsymbol{u}^* = \boldsymbol{0}, \tag{2}$$

$$\frac{\partial C_f^*}{\partial t^*} + \boldsymbol{u}^* \cdot \nabla C_f^* \quad \nabla \cdot \left(D_f^* \nabla C_f^* \right) \quad \boldsymbol{0}, \tag{3}$$

where ρ_f^* denotes the fluid density, η^* the fluid viscosity and D_f^* the chemical species diffusion coefficient in the fluid.

• in the solid

$$\boldsymbol{u}^* \quad \boldsymbol{0}, \tag{4}$$

$$\frac{\partial C_s^*}{\partial t^*} \quad \nabla \cdot \left(D_s^* \nabla C_s^* \right) \qquad k_s^* C_s^*, \tag{5}$$

where D_s^* denotes chemical species effective diffusion coefficient in the solid and k_s^* the effective first order reaction constant in the solid.

• at the fluid/solid interface ∂P

$$\boldsymbol{u}^* \quad \boldsymbol{0}, \tag{6}$$

$$C_s^* \quad C_f^*, \tag{7}$$

$$D_s^* \frac{\partial C_s^*}{\partial \mathbf{n}} = D_f^* \frac{\partial C_f^*}{\partial \mathbf{n}},$$
 (8)

where \boldsymbol{n} denotes the unit normal vector at the fluid/solid interface.

Governing equations are made dimensionless by introducing a characteristic length L_c^* , a characteristic velocity U_c^* and a charac teristic convective time $T_c^* = L_c^*/U_c^*$. In the various problems exam ined thereafter, solid obstacles are spheres, hence an obvious choice for L_c^* is the particle diameter d_p^* . When the problem is not purely diffusive, an obvious choice for U_c^* is the far field inlet veloc ity U_{in}^* . Also, by normalizing the chemical species concentration between 0 and 1 and introducing the chemical species diffusion coefficient ratio $\gamma = \frac{D_s^*}{D_s^*}$ conservation Eqs. (3) and (5) together with interface conditions Eqs. (7) and (8), i.e., continuity of chemical species concentration and continuity of chemical species normal flux, can be recast into a single dimensionless conservation equa tion for the chemical species *C* with appropriate no jump condi tions at the fluid/solid interface on the chemical species concentration and on its normal flux. The set of dimensionless equations eventually reads as follows:

$$\frac{\partial \boldsymbol{u}}{\partial t} + (\boldsymbol{u} \cdot \nabla)\boldsymbol{u} \quad \frac{1}{\mathrm{R}\boldsymbol{e}} \nabla^2 \boldsymbol{u} + \nabla \boldsymbol{p} \quad \boldsymbol{0} \quad \text{in} \quad \Omega \setminus \boldsymbol{P}, \tag{9}$$

$$\nabla \cdot \boldsymbol{u} \quad \mathbf{0} \quad \text{in} \quad \Omega \setminus P, \tag{10}$$

$$\boldsymbol{u} \quad \boldsymbol{0} \quad \text{in} \quad \boldsymbol{P} \cup \partial \boldsymbol{P}, \tag{11}$$

$$\frac{\partial C}{\partial t} + \boldsymbol{u} \cdot \nabla C \quad \nabla \cdot \left(\frac{h(\gamma)}{Pe} \nabla C\right) + \frac{h(\gamma)g(\phi^2)}{Pe}C \quad 0 \quad \text{in} \quad \Omega,$$
$$[C]_{\partial P} \quad 0, \quad \left[h(\gamma)\frac{\partial C}{\partial \boldsymbol{n}}\right]_{\partial P} \quad 0, \tag{12}$$

where $[]_{\partial P}$ represents the jump condition across the fluid/solid interface. The dimensionless numbers introduced above are defined as follows:

- Reynolds number: $\mathbb{R}e^{-\frac{\rho_{f}^{*}U_{c}^{*}L_{c}^{*}}{n^{*}}}$,
- Peclet number: $Pe = \frac{U_c^* L_c^*}{D_c^*}$,
- Damkohler number: $Da \qquad \phi^2 \qquad \frac{k_s^* L_c^*}{D^*}$

where $\phi = \sqrt{Da}$ is the Thiele modulus, and the functions $h(\gamma)$ and $g(\phi^2)$ are simple Heavyside like functions defined as:

$$h(\gamma) \begin{cases} 1 & \text{in } \Omega \setminus P, \\ \gamma & \text{in } P. \end{cases}$$
$$g(\phi^2) \begin{cases} 0 & \text{in } \Omega \setminus P, \\ \phi^2 & \text{in } P. \end{cases}$$

As usual, we can also introduce a Schmidt number $Sc = \frac{\eta^*}{\rho_j^* D_j^*}$ such that Pe = Re Sc. Hence the flow is equivalently characterized by the pair (Re, Pe) or the pair (Re, Sc).

3. Numerical model

The chemical species problem is one way coupled only to the fluid problem through the velocity field **u**. At each discrete time t^{n+1} , n > 0 being the time index and t^0 being the initial time, we solve the full problem as a sequence of the fluid problem followed by the chemical species problem using the computed velocity field u^{n+1} . The solution algorithm is hence of the 1st order in time weak coupling type. In the next subsections, we shortly elaborate on the strategy adopted to solve each sub problem.

3.1. Solution to the fluid problem: flow around fixed obstacles

We use our well validated Finite Volume/Staggered Grid DLM/ FD solver implemented in our in house code PeliGRIFF. The whole method is fully detailed in Wachs et al. (2015), Rahmani and Wachs (2014) for freely moving particles and in Dorai et al. (2015) for fixed obstacles and was pioneered by Glowinski et al. (1999) in a Finite Element context. To summarize, we use a Carte sian structured mesh of constant grid size around obstacles, we solve the fluid conservation Eqs. (9) and (10) everywhere in the domain (not only in $\Omega \setminus P$ but in the entire Ω) and we enforce the rigid body motion (motionless in the particular case of the applica tion treated in this paper) in the region (filled with fictitious fluid) occupied by the obstacles and represented by (11) using a dis tributed Lagrange multiplier field. Our Finite Volume/Staggered Grid DLM/FD method also involves an implicit solution of the resulting DLM/FD saddle point problem by a Uzawa algorithm, a collocation point method to discretize the solid obstacles on the fluid mesh and a second order interpolation of the fluid velocity at the particle boundary (Wachs et al., 2015; Rahmani and Wachs, 2014; Dorai et al., 2015). In our Finite Volume/Staggered Grid discretization method, the diffusive term is discretized with a 2nd order accurate centered scheme while the advective term is discretized with a 2nd order accurate TVD (Total Variation Diminishing)/Superbee limiter scheme. Finally, the diffusive term is treated implicitly in time with a 2nd order Crank Nicholson scheme while the advective term is treated explicitly in time with a 2nd order Adams Bashforth scheme. The strength of our method is that it does not require any kind of hydrodynamic radius calibra tion (see Wachs et al., 2015 for more detail about the problem of hydrodynamic radius calibration). The overall spatial accuracy of the discretization scheme is however not fully 2nd order due to the non boundary fitted feature of the mesh around the solid obstacles. The dimensionless mesh size Δx is related to the number of points per sphere diameter through $N_p = 1/\Delta x$.

The solution algorithm for the fluid problem is of the 1st order operator splitting type and comprises two stages as follows:

 A classical L2 projection scheme for the solution of the Navier & Stokes problem: find u^{n+1/2} and pⁿ⁺¹ such that

$$\frac{\tilde{\boldsymbol{u}} \quad \boldsymbol{u}^{n}}{\Delta t} \quad \frac{1}{2\mathrm{R}e} \nabla^{2} \tilde{\boldsymbol{u}} \qquad \nabla p^{n+1} + \frac{1}{2\mathrm{R}e} \nabla^{2} \boldsymbol{u}^{n} \\ \qquad \qquad \frac{1}{2} \left(3\boldsymbol{u}^{n} \cdot \nabla \boldsymbol{u}^{n} \quad \boldsymbol{u}^{n-1} \cdot \nabla \boldsymbol{u}^{n-1} \right) \quad \boldsymbol{\lambda}^{n},$$
(13)

$$\nabla^2 \psi \quad \frac{1}{\Delta t} \nabla \cdot \tilde{\boldsymbol{u}}, \quad \frac{\partial \psi}{\partial n} \quad 0 \text{ or } \psi \quad 0 \text{ on } \partial \Omega, \tag{14}$$

$$\begin{aligned} \mathbf{u}^{n+1/2} & \tilde{\mathbf{u}} & \Delta t \nabla \psi, \\ p^{n+1} & p^n + \psi & \frac{\Delta t}{2\pi e} \nabla^2 \psi. \end{aligned}$$
 (15)

2. A fictitious domain problem: find \boldsymbol{u}^{n+1} and $\boldsymbol{\lambda}^{n+1}$ such that

$$\frac{\boldsymbol{u}^{n+1} \quad \boldsymbol{u}^{n+1/2}}{\Delta t} + \boldsymbol{\lambda}^{n+1} \quad \boldsymbol{\lambda}^n, \tag{16}$$

$$u^{n+1}$$
 0 in *P*. (17)

where $\mathbf{u}, \lambda, \psi$ and Δt denote the non divergence free predicted fluid velocity vector, DLM/FD Lagrange multiplier to relax the constraint (17), pseudo pressure field and time step, respectively.

3.2. Solution to the chemical species problem: Sharp Interface method

The conservation equation in problem (12) is discretized in time with a 1st order scheme. The diffusive term is treated implicitly in time with a 1st order Backward Euler scheme and the advective term is discretized explicitly in time with a 2nd order Adams Bashforth scheme. Since the reactive term is linear with *C*, it can be easily treated implicitly too. The discrete in time version of the conservation equation in problem (12) reads as follows:

$$\frac{C^{n+1} \quad C^{n}}{\Delta t} \quad \nabla \cdot \left(\frac{h(\gamma)}{Pe} \nabla C^{n+1}\right) + \frac{h(\gamma)g(\phi^{2})}{Pe} C^{n+1}$$
$$\frac{1}{2} \left(3\boldsymbol{u}^{n+1} \cdot \nabla C^{n} \quad \boldsymbol{u}^{n} \cdot \nabla C^{n-1}\right)$$
(18)

The primary difficulty in the spatial discretization of (18) is how to discretize the diffusive operator on a non boundary fitted mesh and to account for the continuity of *C* and of its normal flux across the fluid/solid interface through the 2 no jump conditions $[C]_{\partial P} = 0$ and $[h(\gamma) \frac{\partial C}{\partial n}]_{\partial P} = 0$, respectively. This is achieved with a Sharp Inter face Method (Fedkiw et al., 1999; Liu et al., 2000; Shao et al., 2012). We use the original version suggested by Fedkiw et al. (1999). The method is only 1st order accurate in space but is discontinuity cap turing and easy to implement. The core of the method is to incorporate the jump conditions into the discrete in space diffusive term. The other terms in (18) are discretized in a classical way. Since the method is well established, we simply shortly summarized its construction in 1D and the extension to 3D suggested by Shao et al. (2012) in the special case of spherical solid bodies.

3.2.1. Description of the Sharp Interface Method in one dimension

To ease notation, let's rewrite $\frac{h(\gamma)}{Pe}$ as β . β is hence a diffusion coefficient with a discontinuity across the fluid/solid interface. We shortly elaborate below on the discretization of the diffusive term ∇ ($\beta\nabla C$) in 1D, i.e., (βC_x)_x, with $x = \frac{\partial}{\partial x}$.

We consider general jump conditions on *C* and on its flux defined at the interface ∂P as:

$$[C]_{\partial P} \quad C^+_{\partial P} \quad C^-_{\partial P} \quad a_{\partial P} \tag{19}$$

$$[\beta C_x]_{\partial P} \quad (\beta C_x)_{\partial P}^+ \quad (\beta C_x)_{\partial P} \quad b_{\partial P} \tag{20}$$

where Ω denotes the part of the domain on one side of the inter face (e.g., Ω $\Omega \setminus P$ is the fluid domain) and Ω^+ denotes the part of the domain on the other side of the interface (e.g., $\Omega^+ P$ is the solid domain). We assume that the 1D space is discretized uni formly with a constant grid size Δx and that the interface ∂P lies between two grid points $i \in \Omega$ and $i + 1 \in \Omega^+$. Following Fedkiw et al. (1999), Liu et al. (2000), the diffusive term $(\beta C_x)_x$ for point iis discretized in a Finite Difference fashion as follows:

$$(\beta C_x)_x = \frac{1}{\Delta x} \left[\hat{\beta} \left(\frac{C_{i+1} - C_i}{\Delta x} \right) - \beta \left(\frac{C_i - C_{i-1}}{\Delta x} \right) \right] = \frac{\hat{\beta} a_{\partial P}}{\Delta x^2} = \frac{\hat{\beta} b_{\partial P} (1 - \zeta)}{\beta^+ \Delta x}$$
(21)

where the coefficient $\hat{\beta}$ is calculated as follows:

$$\hat{\beta} = \frac{\beta^+ \beta}{\beta^+ \zeta + \beta \ (1 - \zeta)}$$
(22)

and ζ is the cell fraction, calculated through the level set function χ based on the interface location.

$$\zeta \quad \frac{|\chi_i|}{|\chi_i| + |\chi_{i+1}|} \tag{23}$$

In our case, we have neither a jump for *C* nor for its flux βC_x , which translates into $a_{\partial P}$ 0 and $b_{\partial P}$ 0, so (21) simplifies to:

$$(\beta C_x)_x \quad \frac{1}{\Delta x} \left[\hat{\beta} \left(\frac{C_{i+1} \quad C_i}{\Delta x} \right) \quad \beta \quad \left(\frac{C_i \quad C_{i-1}}{\Delta x} \right) \right]$$
(24)

3.2.2. Extension to three dimensions

The discretization of the diffusive term $\nabla (\beta \nabla C)$ in the vicinity of an interface ∂P in 3D is a bit trickier as jump conditions exist only for *C* and its normal flux $[\beta C_n]_{\partial P}$ 0, but nothing is specified about the tangential flux across ∂P . Liu et al. (2000) suggested a dimension by dimension application of the 1D discretization method that we adopt here. However, Liu et al. (2000) also empha sized that a simple projection of the normal flux jump condition on the Cartesian coordinate axis leads to the right jump condition in the normal direction but also imposes an artificial and essentially physically wrong additional condition on the tangential flux across ∂P of the form $[\beta C_t]_{\partial P}$ 0, while the right physical condition is sim ply $[C_t]_{\partial P}$ 0.

Since our solid obstacles are all spheres, we follow the approach suggested by Shao et al. (2012) that involves changing coordinates from Cartesian to spherical and writing the jump conditions in Cartesian coordinates as a function of the jump conditions in spherical coordinates in a way that the right jump conditions are imposed.

Cartesian and spherical coordinate systems are related to each other through:

- $x \quad r\sin(\theta)\cos(\varphi) \tag{25}$
- $y \quad r\sin(\theta)\sin(\varphi) \tag{26}$
- $z \quad r\cos(\theta) \tag{27}$

with $\theta \in [0, 2\pi]$ and $\varphi \in [\pi/2, +\pi/2]$. The gradient of *C* in the Cartesian coordinate system is related to the gradient of *C* in the spherical coordinate system through:

 $C_{x} = C_{r}\sin(\theta)\cos(\varphi) + C_{t_{\theta}}\cos(\theta)\cos(\varphi) = C_{t_{\varphi}}\sin(\varphi)$ (28)

$$C_y = C_r \sin(\theta) \sin(\varphi) + C_{t_\theta} \cos(\theta) \sin(\varphi) + C_{t_\varphi} \cos(\theta)$$
(29)

$$C_z = C_r \cos(\varphi) = C_{t_{\varphi}} \sin(\varphi)$$
 (30)

where t_{θ} is the unit tangential vector in the θ direction and t_{φ} is the unit tangential vector in the φ direction. Obviously on a sphere sur face we have $C_r = C_n$. Multiplying the above equations by β we can write the jump conditions in the Cartesian directions x, y and z as a function of the jump conditions in the normal and tangential directions as:

$$[\beta C_{x}]_{\partial P} \quad [\beta C_{n}]_{\partial P} \sin(\theta) \cos(\varphi) + [\beta C_{t_{\theta}}]_{\partial P} \cos(\theta) \cos(\varphi) \quad [\beta C_{t_{\varphi}}]_{\partial P} \sin(\theta)$$
(31)

$$[\beta C_z]_{\partial P} \quad [\beta C_n]_{\partial P} \cos(\varphi) \quad [\beta C_{t_\theta}]_{\partial P} \sin(\varphi) \tag{33}$$

In our problem, we have $[C]_{\partial P} = 0, [\beta C_n]_{\partial P} = 0, [C_{t_{\theta}}] = 0$ and $[C_{t_{\varphi}}] = 0$. However, $[\beta C_{t_{\theta}}]_{\partial P} = [\beta]_{\partial P} C_{t_{\theta}}$ and $[\beta C_{t_{\varphi}}]_{\partial P} = [\beta]_{\partial P} C_{t_{\varphi}}$ are non zero because $[\beta]_{\partial P} \neq 0$, i.e., the diffusion coefficient β is discon tinuous across the interface ∂P . To calculate these 2 terms, we need to calculate the tangential derivatives $C_{t_{\theta}}$ and $C_{t_{\varphi}}$ at the particle sur face. At the discrete level, these 2 tangential derivatives are approx imated for each point on the particle surface using a central difference scheme that involves the values of *C* at two adjacent points. As *C* is not known at the time level t^{n+1} , we use *C* from the previous time level, i.e., t^n , to compute $C_{t_{\theta}}$ and $C_{t_{\varphi}}$ as suggested by Shao et al. (2012). Eventually, the 3 jump conditions Eqs. (31) (33) are added to the right hand side of the conservation equation as in a 1D case.

4. Validation

We verify here our implementation of the SIM and assess its capability to resolve the different flow problems we are interested in. We perform a step by step validation of the computed solution in problems of growing complexity. For diffusion and diffusion reaction problems, we compare our SIM results to existing analyt ical solutions. For convection diffusion problems, we compare our SIM results to existing correlations. Finally, for convection diffusion reaction problems, there does not exist any correlation or analytical solution, to the best of our knowledge. For this reason, we compare our SIM results to results computed with a boundary fitted method with local mesh refinement that fully resolves the gradients at the particle interface. We test the limitations of SIM for a wide range of dimensionless numbers. In all the cases exam ined from now on, we are interested in the steady state solution only. However, the steady state solution is computed by our tran sient algorithm as the solution obtained when time derivatives are negligibly small. All transient computations are run with 10³. Δt

4.1. Pure diffusion in a finite domain

4.1.1. Steady state diffusion from a single particle

The first validation test is performed in a purely diffusive regime. A spherical particle of radius $r_p^* \quad d_p^*/2$ is placed at the center of a cubic domain $\Omega \quad L_x^* \times L_y^* \times L_z^*$ and a zero concentration $C_s \quad 0$ is imposed at the particle boundary and inside the particle. This condition is enforced through an infinitely fast reaction $\phi^2 \to \infty$ and a large diffusion coefficient ratio γ that makes the problem mass transfer controlled. A fixed concentration C_∞ is imposed at the boundary $\partial\Omega$ of the domain so that the concentration difference (or driving force) is constant. When the problem is posed in a spherical domain Ω of finite radius r_∞^* , we can solve the 1D problem and derive an expression for the Sherwood number (see Appendix A for the details) as follows:

sh
$$\frac{k^* d_p^*}{D_f^*} = \frac{2r_{\infty}^*}{r_{\infty}^* - r_p^*}$$
 (34)

where k^* denotes the mass transfer coefficient defined as:

$$k^* = \frac{D_f^*}{4\pi r_p^{*2}(C_\infty - C_s)} \int_{\pi/2}^{\pi/2} \int_0^{2\pi} \frac{\partial C(r^*)}{\partial r^*} \bigg|_{r^* - r_p^*} r_p^{*2} \sin(\theta) d\varphi d\theta \qquad (35)$$

We set the dimensionless box size to $L_x = L_y = L_z = 10$. The par ticle dimensionless radius is $r_p = 0.5$. The analytical value of the Sherwood number in a spherical domain of finite radius $r_m = 5$ is Sh_m 20/9 2.22. We assume that this solution is a good approx imation of the solution posed in a cubic domain L_x L_y L_z $2r_{\infty}$. We examine the convergence of the method by computing Sh as a function of the grid size $\Delta x = 1/N_p$. We then compute the relative error with respect to Sh_{an} as $\epsilon = \frac{Sh(N_p) - Sh_{an}}{Sh_{an}}$ and plot the convergence of ϵ with N_p in Fig. 1. The solution follows an expected first order spatial convergence. In this test, the error originates from two contributions. The first contribution is related to the spatial discretization of the problem and the contribution we are inter ested in. The second contribution is due to the fact that the analyt ical solution is derived in a spherical domain and compared to the computed solution in a cubic box. For the range of N_n considered, it is clear in Fig. 1 that the second contribution is negligible and that provided $r_{\infty} = L_x/2$ is chosen large enough, the analytical solution in a spherical domain does not differ much from the solution in a cubic domain.

4.1.2. Steady state diffusion from a single particle in a gradient of concentration

We test our SIM in another diffusive configuration previously examined by Shao et al. (2012). A solid particle of diffusion coefficient D_s^* is immersed at the center of a cubic domain Ω $L_x^* \times L_y^* \times L_z^*$ of fluid at rest of diffusion coefficient D_f^* . Dirichlet boundary conditions C C_1 1 at the top wall and C C_2 0 at the bottom wall are imposed to generate a concentration gradient in the *z* direction. Zero normal flux boundary conditions are imposed on the 4 lateral walls. The average Sherwood number cor responding to the flux through a horizontal *xy* plane is defined as:

Sh
$$\frac{L_z^*}{C_2 C_1} \cdot \frac{1}{L_x^* L_y^*} \int_0^{L_x^*} \int_0^{L_y^*} \left(\frac{\partial C}{\partial Z^*}\right) dy^* dx^*$$
 (36)

The average Sherwood number can be analytically predicted based on the analogy with Maxwell Garnett electric conductivity (Maxwell and Garnett, 2005) as:

$$sh \quad 1 + \frac{3\alpha(\gamma - 1)}{\gamma + 2} \tag{37}$$

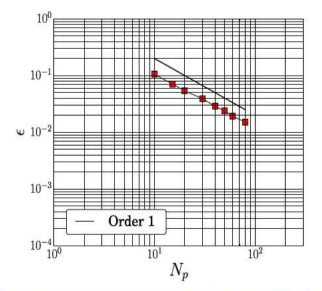


Fig. 1. Steady state diffusion from a single particle in a finite domain L_x L_y L_z 10: spatial convergence of the error on the Sherwood number computed with SIM compared to the analytical solution.

where α is the solid volume fraction. Following Shao et al. (2012), we simulate two cases with two different solid volume fractions. box size We keep the dimensional constant to L_{v}^{*} L_{v}^{*} L_{z}^{*} 10 mm and vary the solid volume fraction α through selecting 2 different particle radii r_p^* 1.25 mm and r_p^* 2 mm, i.e., $r_p^*/L_x^* = 0.125$ and $r_p^*/L_x^* = 0.2$, respectively. We select the diffusion ratio γ to span the interval $\begin{bmatrix} 10^{-2}, 10^{2} \end{bmatrix}$. For each value of γ we com pute the average Sherwood number and compare its value to the theoretical prediction and the numerical results of Shao et al. (2012). Fig. 2 exhibits a very satisfactory agreement of our results with both the analytical prediction and numerical results of Shao et al. (2012). We compute an additional case for r_p^* 1.5 mm, i.e., $r_n^*/L_x^* = 0.15$, and plot the convergence towards the analytical solu tion in Fig. 3 for the three solid volume fractions corresponding to $r_n^*/L_x^* = 0.1, 0.15, 0.2$. As expected, a first order spatial convergence is obtained. The error is shown to increase with the increase of α . The concentration contours in a xz vertical cut plane containing the sphere center are illustrated in Fig. 4 for the case of $r_{\rm p}^*/L_{\rm x}^*$ 0.2 and γ 0.1.

4.2. Steady state internal diffusion and chemical reaction in a single particle

We consider a spherical porous catalyst particle of diameter d_p^* and diffusion coefficient D_s^* immersed at the center of a cubic domain $\Omega = L_x^* \times L_y^* \times L_z^*$. The particle is assumed to undergo a first order chemical reaction controlled by a constant k_s^* such that the chemical species concentration *C* inside the particle satisfies (5). If the surface concentration denoted C_s is known and assumed to be constant over the sphere surface, the problem becomes 1D in space and we can easily write the steady state solution of (5) as follows:

$$C(r) \quad C_s \frac{\sinh(\phi r)}{2r \sinh(\phi/2)} \tag{38}$$

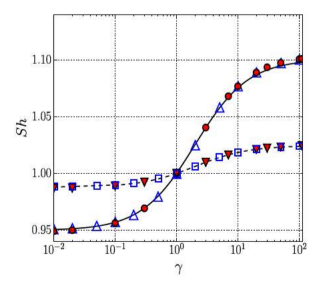


Fig. 2. Steady state diffusion from a single particle in a gradient of concentration: comparison of average Sherwood number computed with SIM to the analytical solution and to the previous study of Shao et al. (2012). Case 1, r_p^*/L_x^* 0.2: continuous black line corresponds to the analytical solution, open triangles correspond to the results of Shao et al. (2012), and circles correspond to our simulation results. Case 2, r_p^*/L_x^* 0.125: dashed black line corresponds to the analytical solution, open squares correspond to the results of Shao et al. (2012), and circles corresponds to the analytical solution, open squares correspond to the results of Shao et al. (2012) and triangles correspond to our simulation results.

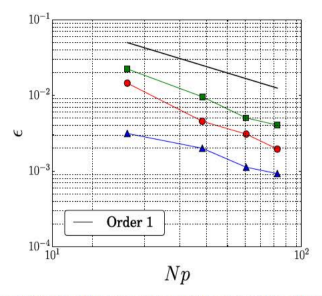


Fig. 3. Steady state diffusion from a single particle in a gradient of concentration: spatial convergence of the relative error on the average Sherwood number computed with SIM. Triangles correspond to $r_p^*/L_x^* = 0.1$, circles correspond to $r_p^*/L_x^* = 0.125$, and squares correspond $r_p^*/L_x^* = 0.2$.

where as usual $r = r^*/d_p^*$ is the dimensionless radial position and $\phi = d_p^* \sqrt{\frac{k_s^*}{D_s^*}}$ is the Thiele modulus. The derivation of (38) can be found in many textbooks.

Dirichlet boundary conditions *C* 1 are imposed on the 6 walls of the cubic domain. We impose C_s 1 through an infinitely large diffusion coefficient D_f^* in fluid phase, i.e., γ is chosen asymptoti cally small. The domain size does not matter here and only needs to be chosen dimensionlessly larger than 1. We compare our SIM concentration profile inside the particle to the analytical solution (38) in Fig. 5 for N_p 20. The agreement is once again very satis factory. The numerical method is capable of capturing the steep concentration gradients at the particle surface even for a modest resolution. The corresponding concentration iso countours are shown in Fig. 6 for N_p 80.

Based on the concentration profile, the effectiveness factor η for a catalyst particle can be defined as the ratio of the overall internal reaction rate in the particle to the reaction rate that can be attained in the absence of diffusion limitations. For a spherical catalyst par ticle with a fixed surface concentration C_{s_1} we have:

$$\eta = \frac{\int_{0}^{2\pi} \int_{\pi/2}^{\pi/2} \int_{0}^{r_{p}^{*}} \frac{k_{s}^{*} C(r^{*}) \sin(\theta) r^{*2} d\phi d\theta dr^{*}}{k_{s}^{*} \frac{4\pi}{3} r_{p}^{*3} C_{s}}}{\frac{\overline{C_{\nu}}}{C_{s}} - \frac{6}{\phi} \left(\frac{1}{\tan h(\phi/2)} - \frac{2}{\phi}\right)}$$
(39)

where $C(r^*)$ $C_s \frac{d_p^* \sinh(\phi r^*/d_p^*)}{2r^* \sinh(\phi/2)}$ is the Thiele concentration profile given by (38) in a dimensional form for a given surface concentration C_s and $\overline{C_p}$ is the average volume concentration in the particle.

Depending on reaction kinetics and assuming, e.g., $\gamma \in [10^2, 10^2]$, i.e., γ is not asymptotically small/large, the effec tiveness factor η exhibits two asymptotic limits. When the reaction rate is very low compared to diffusion, $\phi \ll 1$, the system is con trolled by kinetics and the catalyst surface concentration C_s is equal to fluid concentration, and so $\eta \rightarrow 1$. When the reaction rate is high compared to diffusion, i.e. $\phi \gg 1$, the system is limited by diffusion and the surface concentration C_s approaches zero, there fore $\eta \rightarrow 6/\phi$.

In the following test, we once again impose $C_s = 1$ through an asymptotically small γ . We vary the Thiele modulus ϕ in the range

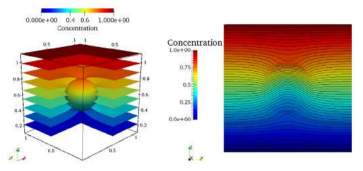


Fig. 4. Steady state diffusion from a single particle in a gradient of concentration: concentration iso-countours for r_p^*/L_x^* 0.2 and γ 0.1 in a 3D view (left) and in a xz vertical cut plane containing the sphere center (right).

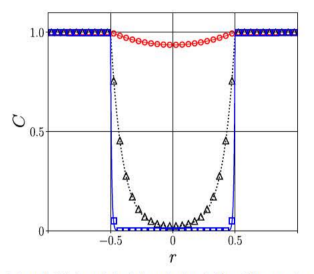


Fig. 5. Internal diffusion and chemical reaction in a single particle: comparison of concentration profiles computed with SIM to analytical profiles for different Thiele moduli. Lines correspond to analytical solutions and markers correspond to our simulation results. Red line and red circles for ϕ^2 1.6. Black doted line and open triangles for ϕ^2 160. Blue line and open squares for ϕ^2 16,000. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

[0.2, 120] by varying the reaction rate constant k_s^* . For each value of ϕ , we calculate numerically the effectiveness factor η and examine the spatial convergence of the solution computed with our SIM and a standard FVWA method. Hence we compute SIM solutions for N_p 10,40,80 and FVWA solutions for N_p 40,80. Fig. 7 shows η as a function of ϕ for N_p 10,80 with SIM and N_p 40 with FVWA. Results obtained with SIM and FVWA are further compared

to each other and to the analytical solution in Figs. 8 and 9 for var ious values of ϕ and N_p . We can make the 3 following comments: (ii) both methods show a first order spatial convergence, (ii) FVWA with a grid size 4 times smaller than SIM gives approximately the same computed solution, and (iii) when N_p 80, SIM predictions are very close to the analytical solution with a slightly growing deviation for $\phi > 20$. The superior accuracy of the solution com puted with SIM compared to that computed with FVWA is further emphasized in Fig. 8 where we plot the error to the analytical solu tion as a function of ϕ for N_p 80. Finally, Fig. 9 also highlights that fact that the magnitude of the error increases with increasing ϕ in relation to the concentration gradients becoming steeper in the vicinity of the particle surface as ϕ increases, i.e., the internal mass boundary layer gets thinner as ϕ increases.

4.3. Steady state convection diffusion in the flow past a single sphere: external mass transfer problem

We now validate our SIM in the case of external mass transfer in the flow past a spherical solid particle in an unbounded domain. A spherical particle is placed in a box of size $L_x \times L_y \times L_z$ $5 \times 5 \times 15$. The fluid enters the flow domain on the left boundary with an imposed fluid velocity **u** (0,0,1) and a concentration $C \quad C_\infty$ 1. The particle is centered in the *x* and *y* directions. Peri odic boundary conditions are imposed in *x* and *y* directions while a classical outflow boundary condition $\frac{\partial u}{\partial z} = \frac{\partial C}{\partial z}$ 0 and $p \quad p_{ref}$ 0 is imposed at the outlet boundary. Fig. 10 illustrates the flow config uration in a *xz* cut plane containing the sphere center. The concen tration *C* in the particle is imposed to 0 by selecting an extremely large value of ϕ . The problem is hence controlled by Sc and Re.

To illustrate that our SIM predicts the right external mass trans fer, we set Sc 1 and vary Re in the range [10,200]. We compute the steady state Sherwood number Sh for N_p 80 and compare its value to literature correlations in Fig. 11. Overall, we observe a very

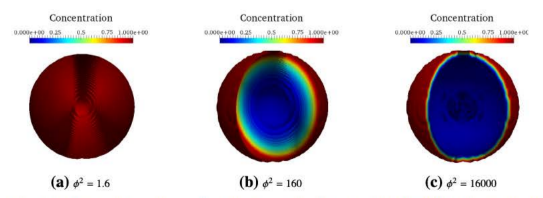


Fig. 6. Internal diffusion and chemical reaction in a single particle: concentration iso-surfaces inside the particle for different reaction rates.

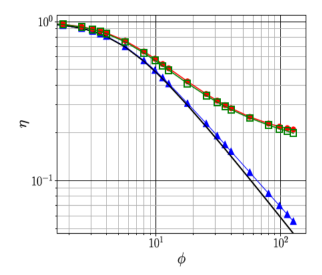


Fig. 7. Internal diffusion and chemical reaction in a single particle: comparison of effectiveness factor computed with FVWA and SIM as a function of ϕ . Red color corresponds to FVWA obtained with N_p 40 and green color corresponds to SIM obtained with N_p 10. Blue color corresponds to SIM with N_p 80 and black line corresponds to the analytical solution. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

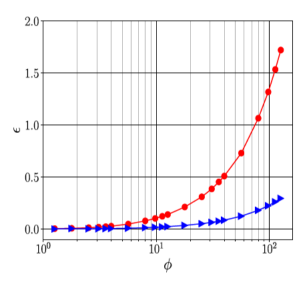


Fig. 8. Internal diffusion and chemical reaction in a single particle: relative error on the effectiveness factor as a function of ϕ for N_p 80. Red color corresponds to FVWA and blue color corresponds to SIM. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

satisfactory agreement between our SIM results and literature cor relations. The correlation of Feng and Michaelides (2000) seems to be the closest to our SIM results. A plausible reason is that Feng and Michaelides (2000) derived their correlation numerically through PRS as we did while other correlations as, e.g., the correla tion suggested by Ranz and Marshall (1952), were obtained experimentally.

4.4. Steady state convection diffusion in the flow past a single reactive sphere: external internal mass transfer

We now consider a similar problem to the one in Section 4.3 but with a first order reaction inside the particle. To validate our SIM, we compare its computed results to results computed by a highly accurate boundary fitted method implemented in the JADIM code.

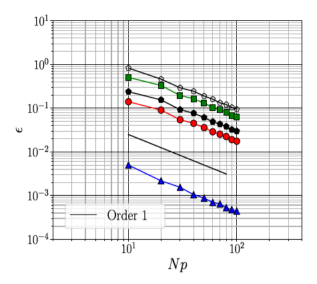


Fig. 9. Internal diffusion and chemical reaction in a single particle: spatial convergence of relative error on effectiveness factor computed with SIM and FVWA for various Thiele moduli: ϕ 1.2, blue triangles with SIM, ϕ 1.2, red circles with SIM, ϕ 20, black filled pentagons with SIM, black open pentagons with FVWA, ϕ 120, green squares with SIM. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

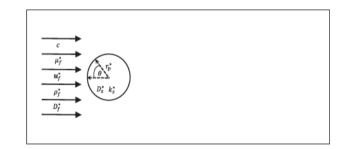


Fig. 10. Geometric configuration for the problem of steady state convectiondiffusion in the flow past a single sphere with or without reaction inside the particle: view in a *xz* cut plane containing the sphere center (simulation domain is $5d_p^* \times 5d_p^* \times 15d_p^*$).

JADIM solves the same system of Eqs. (9) (12) but with a different discretization scheme in space and in time. The high accuracy of solutions computed with JADIM derives both from its boundary fit ted spatial discretization scheme and the ability to locally refine the mesh in both the internal and external mass boundary layers.

The numerical methods used in JADIM have been thoroughly described by Magnaudet et al. (1995) and Calmet and Magnaudet (1997). Consequently they are only quickly summarized here. The JADIM code solves the incompressible Navier Stokes equations and the concentration equation in general orthogonal curvilinear coordinates which are boundary fitted to the particle surface. Equations are integrated in space using a finite volume method in which advective and diffusive terms are evaluated with second order accurate centered schemes. The solution is advanced in time by means of a three step Runge Kutta time stepping proce dure in which advective terms are computed explicitly while diffu sive terms are treated by a semi implicit Crank Nicholson scheme. Incompressibility is satisfied after the third intermediate time step by solving a Poisson equation for an auxiliary potential from which the true pressure is deduced, similarly to Eqs. (13) (15). The com plete algorithm is second order accurate in both space and time.

The mesh grid used in the present work is sketched in Fig. 12. The orthogonal axisymmetric mapping is obtained by using the streamlines and the equipotential lines of the potential flow

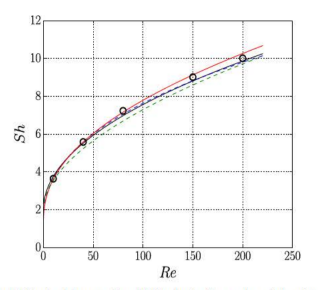


Fig. 11. Steady state convection-diffusion in the flow past a single sphere: comparison of computed Sherwood number with literature correlations at Sc 1. Our SIM results are represented by circles while correlations are as follows: red line for Feng and Michaelides (2000), black line for Ranz and Marshall (1952), green-doted line for Whitaker (1972) and blue-dashed line for Clift et al. (2005). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

around a circular cylinder. The mesh is stretched in order to have at least four points in the external mass boundary layer that scales as $Pe^{-1/3}$. Simulations are performed in a 2D axisymmetric configura tion which reproduces the geometry of a spherical particle. The fluid computational domain is limited by the particle surface and by external boundaries on which inflow, free stream, axial symme try, and outflow boundary conditions are imposed. The equations are solved inside the particle over a polar mesh adjusted to the fluid mesh at the particle surface. The internal mass boundary layer thickness reduces when the kinetics of the chemical reaction, i.e., ϕ , increases. The mesh inside the particle is thus refined close to the particle surface following the scaling of the boundary layer as ϕ^{-1} . At least four grid points stand within the internal boundary layer in order to compute properly the internal concentration gra

dient at the particle surface. A particle of radius r_p^* is placed in a domain with a spatial extension of at least r_{∞}^* 100 r_p^* , so that the assumption of infinite domain is physically valid.

All simulations are performed with N_p 80 unless spatial convergence is investigated. The DLM FD/SIM simulation domain is $5 \times 5 \times 15$. We set γ 10, ϕ 2 and Sc 1 and vary Re in the range [10, 200]. Our SIM results plotted in Fig. 13 agree well with the JADIM results and the model for the mean surface concentration $\overline{C_s}$ suggested in Sulaiman et al. (2018). Fig. 13 shows that the mean surface concentration $\overline{C_s}$ increases with the increase of Re. Then we set γ 10, ϕ 150 and Sc 1 and vary ϕ in the range [0, 40]. Once again a good agreement between our SIM results, the JADIM results and the model for the mean surface concentration $\overline{C_s}$ suggested in Sulaiman et al. (2018) is highlighted in Fig. 14, with a small deviation of our SIM results from the JADIM results and the model for the JADIM results and the model for the JADIM results and the model for the mean surface concentration $\overline{C_s}$ suggested in Sulaiman et al. (2018) is highlighted in Fig. 14, with a small deviation of our SIM results from the JADIM results and the model predictions that grows as ϕ increases due to the internal boundary layer getting thinner as $1/\phi$ and culminates at about 12% for the highest ϕ 4000.

We now compare concentration profiles computed with our SIM and JADIM. We set y 10, Re 10, and Sc 1, select two Thiele moduli ϕ 4 and ϕ 10 and plot the concentration profile obtained with each method in Fig. 15. The agreement is visually very good. The actual error between SIM and JADIM (not shown here for the sake of conciseness) increases for ϕ 10 compared to ϕ 4, in line with what we observe for $\overline{C_s}$. We run another set of simulations with γ 10, ϕ 10, Sc 1 and two Reynolds num 10 and Re 100. Concentration profiles plotted in Fig. 16 bers Re once again in a cross section normal to the inlet flow, correspond 90°. They highlight a satisfactory agreement between ing to θ our SIM results and the JADIM results. The error between SIM and JADIM (not shown here for the sake of conciseness) only mildly increases with Re. The numerical accuracy is consequently mostly controlled by the internal mass boundary layer thickness.

Finally, we examine the spatial convergence of the SIM computed solution to the JADIM reference solution. We set $\gamma = 0.1, \phi^2 = 40$ and Re = 150, select Sc = 0.1, Sc = 1 and Sc = 10 and N_p in the range [10, 80]. We then compute and plot the error ϵ on the mean surface concentration $\overline{C_s}$ between SIM and JADIM. Fig. 17 shows that ϵ increases slightly only with the increase of Sc. This behavior is similar to the behavior of the solution with increasing Re at constant ϕ , that shows that ϵ is mainly

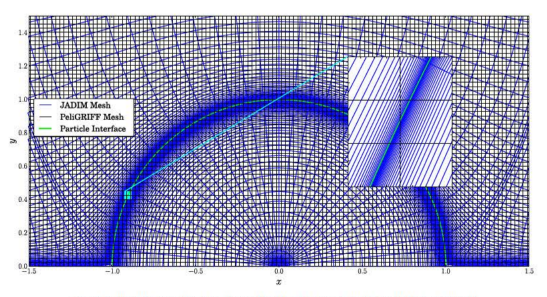


Fig. 12. Comparison of the finest meshes used by JADIM (boundary fitted) and SIM (Cartesian grid).

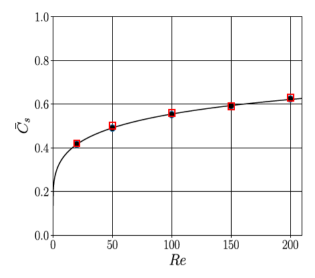


Fig. 13. Steady state convection-diffusion in the flow past a single reactive sphere: comparison of the mean surface concentration $\overline{C_s}$ at $\phi = 2$, $\gamma = 10$ and Sc = 1 as function of Re, computed with SIM (red squares), computed with JADIM (black circles), and predicted by the model of Sulaiman et al. (2018) (black line). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

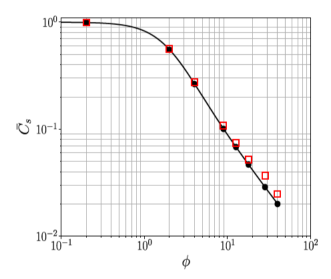


Fig. 14. Steady state convection-diffusion in the flow past a single reactive sphere: comparison of mean surface concentration $\overline{C_s}$ at Re 150, γ 10 and Sc 1 as function of ϕ , computed with SIM (red squares), computed with JADIM (black circles), and predicted by the model of Sulaiman et al. (2018) (black line). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

controlled by the reaction rate and the internal mass boundary layer thickness. To investigate the effect of the reaction rate on the solution, we perform a final set of simulations with γ 0.1, Sc 10, Re 150 and ϕ^2 varying in the range [40, 4000]. Fig. 18 reveals that ϵ increases significantly with the increase of ϕ^2 . For ϕ^2 40, even a mesh resolution N_p 20 is sufficient to sup ply a computed solution with a relative error of 3%. At ϕ^2 200, a grid resolution N_p 60 is needed to reach the same accuracy. For ϕ^2 400, a grid resolution with more than N_p 80 is needed to reach an accuracy corresponding to a relative error of less than 4%. And finally at ϕ^2 4000, even the finest grid resolution consid ered here N_p 80 supplies a computed solution that is still 11% off from the reference solution.

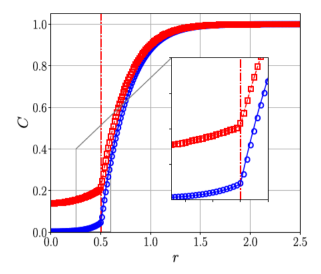


Fig. 15. Steady state convection-diffusion in the flow past a single reactive sphere: comparison of concentration profile in the direction normal to the flow ($\theta = 90^\circ$), at Re 10, $\gamma = 10$ and So = 1 for two Thiele moduli: $\phi = 4$, red color, and $\phi = 10$, blue color. Continuous line correspond to JADIM and markers correspond to SIM. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

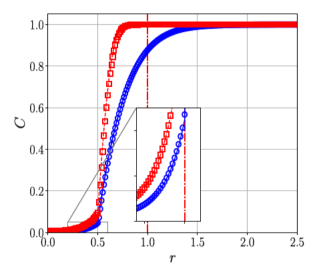


Fig. 16. Steady state convection-diffusion in the flow past a single reactive sphere: comparison of concentration profile in the direction normal to the flow (θ 90°), for ϕ 4, γ 10 and Sc 1 and two Reynolds numbers: Re 100, red color, and Re 10, blue color. Continuous lines correspond to JADIM and markers correspond to SIM. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

5. Interacting spheres

We examine here the external forced convection diffusion on a sequence of interacting spheres undergoing an internal first order irreversible chemical reaction. The non reactive problem without diffusion inside solid bodies was firstly introduced by Ramachandran et al. (1989) for heat transfer. As in Section 4, we are interested in the steady state solution only and all transient computations leading to steady state are run with $\Delta t = 10^{-3}$. Ramachandran et al. (1989) suggested empirical corrective terms to relate the non reactive Sherwood number of a single sphere to the non reactive Sherwood numbers of interacting spheres. The corrective terms account for separation distances effect between the particles. The aim of this work is to include the effect of a

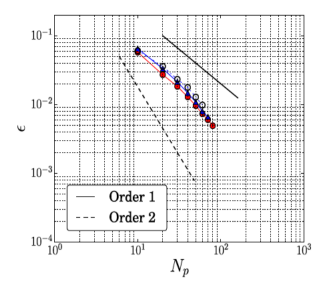


Fig. 17. Steady state convection-diffusion in the flow past a single reactive sphere: spatial convergence of relative error on mean surface concentration $\overline{C_s}$ computed with SIM at $\mathbb{R}e_{-}150, \phi^2_{-}40$, and $\gamma_{-}0.1$ for $\mathbb{S}e_{-}0.1$, red circles, $\mathbb{S}e_{-}1$, black triangles, and $\mathbb{S}e_{-}10$, black open circles. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

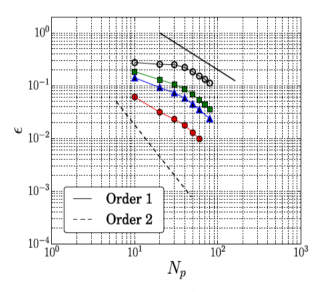


Fig. 18. Steady state convection-diffusion in the flow past a single reactive sphere: spatial convergence of relative error on mean surface concentration $\overline{C_s}$ computed with SIM at $\mathbb{R}e_150$, $\mathbb{S}c_10$, and γ_10 , $10 \text{ er}\phi^2_1$ 40, red circles, ϕ^2_2 200, blue triangles, ϕ^2_1 400, green squares, and ϕ^2_2 4000, black open circles. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

chemical reaction inside the solid bodies and to introduce, based on internal external coupling, a model for the reactive Sherwood number for each particle, that accounts for the effects of diffusion, convection, and reaction. We aim at extending the model we sug gested for a single sphere in (Sulaiman et al., 2018) to more con centrated particle laden flows and consider the flow configuration investigated here as a proper toy model for mass transfer with reaction in dilute particle laden flows. The core aspect of our model is that it is possible to construct a reactive Sherwood number formula based on an existing non reactive Sher wood number formula. We show thereafter that this approach is still valid in dilute particle laden systems experiencing hydrody namic interactions.

5.1. Problem definition

We consider a system composed of three aligned spherical cat alyst particles along z in a box of size $L_x \times L_y \times L_z$ $5 \times 5 \times 15$. The fluid enters the flow domain on the left boundary with an imposed fluid velocity **u** (0,0,1) and concentration C 1. The three parti cles are centered in the x and y directions. Periodic boundary con ditions are imposed in x and y directions while a classical outflow boundary condition $\frac{\partial u}{\partial z} = \frac{\partial C}{\partial z}$ 0 and p p_{ref} 0 is imposed at the outlet boundary. The separation distances between the first and second particles and between the second and third particles are d_{12} and d_{23} , respectively, as shown in Fig. 19. The problem is con trolled by the Reynolds number Re, the Schmidt number Sc, the diffusion coefficient ratio γ and the Damkohler ϕ^2 . All computa tions are performed with N_p 70.

We approach the problem through the external internal coupling based on the continuity of *C* and of its normal flux at the fluid/solid interface ∂P . The normal flux density at the particle sur face in the solid phase $N_{s,\partial P}^*$ is given by:

$$N_{s,\partial P}^* = D_s^* \frac{dC}{dr^*} \Big|_{r^* = r_p^*}$$

$$\tag{40}$$

The concentration profile in the solid phase is given by (38). Calculating the radial derivative at r^* r_p^* (or r = 0.5), the flux in the solid phase can be written as:

$$N_{s,\partial P}^{*} = \frac{D_{s}^{*}C_{s}}{d_{p}^{*}} \left(\frac{\phi}{tanh(\phi/2)} - 2\right)$$
(41)

The normal local flux density at the particle surface in the fluid phase $N_{f,\partial P}^*$ can be written as:

$$N_{f,\partial P}^* = k_f^*(C_s - C_\infty) \tag{42}$$

where k_f^* is the external local mass transfer coefficient in the fluid phase, analytically unknown in case of external convection. The continuity of the normal flux at ∂P , i.e., $N_{s,\partial P}^* = N_{f,\partial P}^*$, leads to:

$$k_f^*(C_s \quad C_\infty) \quad D_s^* \frac{C_s}{d_p^*} \left(\frac{\phi}{tanh(\phi/2)} \quad 2 \right)$$
(43)

And we can hence deduce the expression of the surface concentration C_s :

$$C_s = \frac{C_{\infty}}{1 + \frac{D_s^*}{d_p^* k_f^*} \left(\frac{\phi}{tanh(\phi/2)} - 2\right)}$$
(44)

In order to determine C_s , we need to evaluate the external mass transfer coefficient k_f^* . Here, we assume that the internal problem is only affecting the external problem through the concentration gradient. Thus, the external Sherwood number is assumed indepen dent of the reaction rate. We hence evaluate the average external

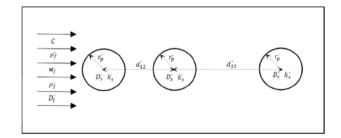


Fig. 19. Geometric configuration for the problem of steady state convectiondiffusion in the flow past 3 aligned reactive spheres: view in a xz cut plane containing the sphere center (simulation domain is $5d_p^* \times 5d_p^* \times 15d_p^*$).

mass transfer coefficient from the external Sherwood number Sh, i.e., a non reactive Sherwood number, as $k_f^* = Sh D_f^*/d_p^*$. Then, any appropriate empirical correlation available in the literature for Sh can be used. For instance in (Sulaiman et al., 2018), we used for a single sphere the correlation defined by Feng and Michaelides (2000):

Sh
$$0.922 + \operatorname{R}e^{1/3}\operatorname{Sc}^{1/3} + 0.1\operatorname{R}e^{2/3}\operatorname{Sc}^{1/3}$$
 (45)

For a series of three interacting spheres, Ramachandran et al. (1989) suggested to account for the interactions between the spheres through corrective coefficients for the Sherwood number of each of the three spheres. These corrective coefficients, mainly empirical, are established based on computational data, and related to the correlation for a single sphere. The individual Sher wood number reads:

$$\mathrm{S}h_j \quad \mathrm{S}h \cdot \beta_j \quad j \quad 1, 2, 3$$

$$\tag{46}$$

where *j* is the sphere number and the β_j , j = 1, 2, 3 are the empirical corrective terms defined by Ramachandran et al. (1989) as:

$$\beta_1 = 0.9555 \operatorname{Sc}^{0.0276} \operatorname{R} e^{0.0108} d_{12}^{0.113/[1+1.5ln(10Pe)]} e^{-0.02113/d_{23}}$$
(47)

$$\beta_{2} = 1 = 0.0697(1 + ln(\text{Re}))^{0.767} [1 + ln(10\text{ReSc})]^{0.095} d_{23}^{-0.13}$$

$$4.807(1 + ln(\text{Re}))^{0.012} d_{12}^{-5.28}$$
(48)

$$\beta_3 = 1 = 0.532 \operatorname{Sc}^{-0.019} \operatorname{R} e^{0.032} e^{-0.0146/d_{12}} \left(\frac{1}{d_{23}}\right)^{3.044/[1+1.1ln(10Pe)]}$$
(49)

The average surface concentration $\overline{C_{sj}}$ for sphere *j* with correspond ing Sherwood number Sh_j , can be written as follows:

$$\overline{C_{s,j}} = \frac{C_{\infty}}{1 + \frac{2\gamma}{sh_j} \left(\frac{\phi/2}{tanh(\phi/2)} - 1\right)} \quad j = 1, 2, 3$$
(50)

The formulation of a 'reactive' Sherwood number that accounts for convection, diffusion and reaction is based on two concentra tion gradients instead of a single concentration gradient in the external non reactive case. The first gradient is the external gradi ent involving $(\overline{C}_s \quad C_{\infty})$ and the second gradient is the internal gradient involving $(\overline{C_v} \quad \overline{C_s})$, where $\overline{C_v}$ is the mean volume con centration of *C* in each sphere. To derive an expression for $\overline{C_v}$, we integrate the 1D concentration profile given by (38), assuming C_s is a function of the position on the sphere and the radial depen dence is still reasonably valid, over the particle volume as

$$\overline{C_{\nu}} \simeq \frac{3}{4\pi r_{p}^{*3}} \int_{0}^{2\pi} \int_{-\pi/2}^{+\pi/2} \int_{0}^{r_{p}} C_{s}(\phi,\theta) \\
\times \frac{r_{p}^{*} \sinh(\phi r^{*})}{r^{*} \sinh(\phi r_{p}^{*})} \sin(\theta) r^{*2} d\varphi d\theta dr^{*}$$
(51)

and further approximate this integral as:

$$\overline{C_{\nu}} \simeq \frac{3\overline{C_{s}}}{4\pi r_{p}^{*3}} \int_{0}^{2\pi} \int_{-\pi/2}^{+\pi/2} \int_{0}^{r_{p}^{*}} \frac{r_{p}^{*}\sinh(\phi r^{*})}{r^{*}\sinh(\phi r_{p}^{*})} \sin(\theta) r^{*2} d\varphi d\theta dr^{*}$$

$$\frac{6\overline{C_{s}}}{\phi} \left(\frac{1}{\tanh(\phi/2)} - \frac{2}{\phi}\right)$$
(52)

 $\overline{C_s}$ is estimated by model (50) and we eventually obtain an expres sion for the mean volume concentration $\overline{C_{vj}}$ for sphere *j* that reads as follows:

$$\overline{C_{\nu j}} \quad \frac{6C_{\infty}}{1 + \frac{2\gamma}{Sh_j} \left(\frac{\phi/2}{tanh(\phi/2)} - 1\right)} \left(\frac{1}{\phi tanh(\phi/2)} - \frac{2}{\phi^2}\right) \quad j = 1, 2, 3$$
(53)

The aim is to establish a model for a Sherwood number that accounts for the effect of a first order irreversible reaction along with convection and diffusion. The model depends on mean vol ume concentration instead of mean surface concentration, and can be used as a closure law in Euler/Lagrange meso scale simula tions, where mean volume quantities are known at particle scale. The mass flux balance in this case is written as:

$$\frac{\pi d_p^{*3}}{6} \frac{d\overline{C}_\nu}{dt^*} \qquad 4\pi r_p^{*2} h^* \left(\overline{C}_\nu \quad C_\infty\right) \quad \frac{\pi d_p^{*3}}{6} k_s^* \overline{C}_\nu \tag{54}$$

Mass conservation yields:

$$4\pi r_p^{*2} h^* \left(\overline{C}_v \quad C_\infty\right) \quad 4\pi r_p^{*2} k_f^* \left(\overline{C_s} \quad C_\infty\right) \tag{55}$$

where k_f^* is the external mass transfer coefficient referring to con centration difference $\Delta \overline{C}_s$ ($\overline{C_s}$ C_∞), and h^* is the overall mass transfer coefficient that accounts for internal and external effects and refers to the mean catalyst particle concentration, $\Delta \overline{C}_v$ (\overline{C}_v C_∞). Thus, the equivalent Sherwood number including the effect of chemical reaction can be obtained as:

$$\widetilde{\mathrm{Sh}} \quad \frac{h^* d_p^*}{D_f^*} \quad \mathrm{Sh}(\mathrm{R}e, \mathrm{Se}) \frac{\Delta \overline{C}_s}{\Delta \overline{C}_v}$$
(56)

The conjugate mass transfer problem can also be formulated with the additivity rule. The additivity rule states that the overall resistance to mass transfer in the system is the sum of two resis tances. The resistance is related to the inverse of the mass transfer coefficients. We hence define the two Sherwood numbers as:

• internal Sherwood number Shin

$$Sh_{in} = \frac{N_{s,\partial P}^* d_p^*}{D_s^* (\overline{C_s} - \overline{C_v})} = \frac{k_{in}^* d_p^*}{D_s^*}$$
(57)

where k_{in}^* is the internal mass transfer coefficient. • external Sherwood number $\mathbb{S}h$

$$Sh = \frac{N_{f,\partial P}^* d_p^*}{D_f^* \left(C_{\infty} - \overline{C_s}\right)} = \frac{k_f^* d_p^*}{D_f^*}$$
(58)

We now introduce the total mass transfer coefficient k^* and write the additivity rule as:

$$\frac{1}{\tilde{k}^*} = \frac{1}{k_{in}^*} + \frac{1}{k_f^*}$$
(59)

(59) can be reformulated in terms of Sherwood numbers. Intro ducing the reactive (i.e. total) Sherwood number $\widetilde{Sh} = \frac{k^* d_p^*}{D_j^*}$, we get:

$$\frac{1}{\widetilde{sh}} = \frac{1}{\gamma sh_{in}} + \frac{1}{sh}$$
(60)

Finally, using the continuity of the flux density at ∂P , we can obtain Sh_{in} as a function of Sh, insert this in (60) and after some simple algebra get the following expression:

$$\widetilde{\mathrm{Sh}} \quad \mathrm{Sh}(\mathrm{R}e,\mathrm{Sc})\frac{C_{\infty}}{C_{\infty}} \quad \overline{C_{s}} \quad \mathrm{Sh}(\mathrm{R}e,\mathrm{Sc})\frac{\Delta\overline{C_{s}}}{\Delta\overline{C_{v}}}$$
(61)

As expected, (61) is similar to (56). Finally, using (50) and (52), we establish the expression of the reactive Sherwood number \widetilde{Sh} of each sphere as:

$$\widetilde{Sh_{j}}(\mathbb{R}e, \mathbb{S}c, \phi, \gamma) = \frac{Sh_{j}(\mathbb{R}e, \mathbb{S}c)}{\frac{Sh_{j}(\mathbb{R}e, \mathbb{S}c)}{2\gamma} \left[\frac{tanh(\phi/2)}{\phi/2 \ tanh(\phi/2)} - \frac{12}{\phi^{2}}\right] + 1} \qquad j = 1, 2, 3$$
(62)

 $Sh_j(Re, Sc)$ in the above expression is the individual Sherwood number for a sphere *j* in the convective diffusive problem (without reaction) where the particle internal concentration is uniform, and the mass transfer is only controlled by the Reynolds number Reand the Schmidt number Sc. The 'reactive' Sherwood number, $\widetilde{Sh_j}(Re, Sc, \phi, \gamma)$, also depends on the Damkohler number $Da \quad \phi^2$ and the diffusion coefficient ratio γ , in addition to the first two dimensionless numbers for convection diffusion. (62) can further be written in an explicit form:

$$\frac{1}{\widetilde{\mathrm{Sh}}_{i}} = \frac{1}{2\gamma} \left[\frac{\tanh(\phi/2)}{\phi/2} \frac{12}{\tanh(\phi/2)} - \frac{12}{\phi^{2}} \right] + \frac{1}{\mathrm{Sh}_{i}}$$
(63)

(63) has two asymptotic limits. When the reaction rate is very fast, i.e., the Thiele modulus $\phi \to \infty$, the term $\begin{bmatrix} \tanh(\phi/2) & \frac{12}{\phi^2} & 0 \end{bmatrix} \to 0$ and so $\widetilde{Sh_j} \to Sh_j$. In this case the system is limited by diffusion, i.e., con trolled by mass transfer. When the reaction rate is very slow, i.e. the Thiele modulus $\phi \to 0$, the term $\begin{bmatrix} \tanh(\phi/2) & \frac{12}{\phi^2} & 0 \end{bmatrix} \to \infty$, $\widetilde{Sh_j} \to 0$. In this case the system is controlled by internal reaction kinetics.

5.2. Model validation

We examine here how the model performs in the 3 aligned sphere configuration by spanning ranges of the 4 governing dimen sionless numbers $\operatorname{Re}, \operatorname{Sc}, \gamma$ and ϕ^2 , as well as assorted interparticle distances. The mean surface concentration $\overline{C_s}$, given by (50), is the main unknown in the problem. Therefore, we primarily assess the validity of our proposed model by comparing $\overline{C_s}$ as computed by our DLM/FD SIM numerical method to $\overline{C_s}$ as predicted by our model.

We vary the dimensionless parameters in the ranges: $\operatorname{Re} \in [10, 100], \phi^2 \in [0, \infty], \text{Sc} \in [0.5, 10] \text{ and } \gamma \in [10^{-2}, 10^2] \text{ and select the following three different geometric configurations:}$

We first examine the influence of the geometric configuration. To do this, we set Re = 50, $\gamma = 0.1$, and Se = 1 and keep them con stant while we vary the Damkohler number ϕ^2 in the range [0,400] and compare our model to our numerical results for the three geo

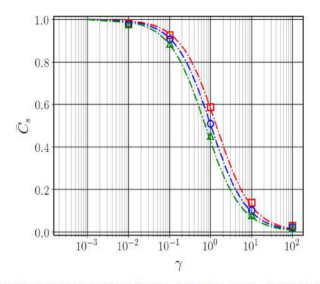


Fig. 21. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: mean surface concentration $\overline{C_s}$ as a function of γ in configuration A at Re = 50, Sc = 1 and $\phi^2 = 40$. Lines correspond to model and markers correspond to SIM. Red color corresponds to first sphere, blue color to second sphere and green color corresponds to third sphere. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

metric configurations A, B, and C in Fig. 20a, b and c, respectively. Overall, $\overline{C_s}$ as predicted by the model show a good agreement with computed $\overline{C_s}$ for the three geometric cases. Essentially, the error increases with the increase of ϕ^2 as the internal concentration boundary layer thickness decreases with the increase of ϕ^2 as $1/\phi$. However, the inter particle distances have literally no impact on the magnitude of the differences between model predictions and numerical results. Even in configuration C where the 3 parti cles are the closest to each other, no significant difference is visible between Fig. 20c, and Figs. 20a and b. For the 3 configurations, the system is still dilute enough such that the first sphere behaves sim ilarly, and quite close to a single isolated sphere.

From now on, we consider the geometric configuration A in the rest of this subsection. We fix Re 50, Sc 1, and ϕ^2 40 and we vary the diffusion ratio γ in the range $\left[10^{-2}, 10^2\right]$. Once again, we compute numerically the mean surface concentration $\overline{C_s}$ and we compare it to the value predicted by the model for the three particles in Fig. 21. Results show again a good agreement between model predictions and numerical results.

Finally, we examine the influence of Sc. To do this, we fix Re 50, γ 0.1, and ϕ^2 40 and we vary Sc \in [0.5, 10], which

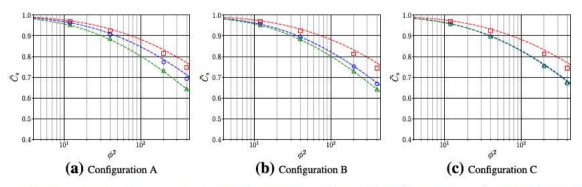


Fig. 20. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: mean surface concentration $\overline{C_s}$ as a function of ϕ^2 in the three configurations (a) A, (b) B and (c) C at Re = 50, Sc = 1 and $\gamma = 0.1$. Lines correspond to model and markers correspond to SIM. Red color corresponds to first sphere, blue color to second sphere and green color corresponds to third sphere. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

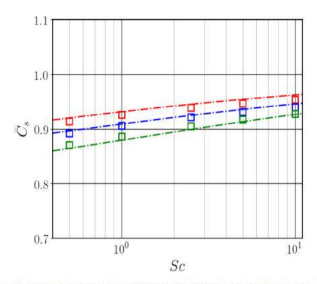


Fig. 22. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: mean surface concentration $\overline{C_s}$ as a function of S₀ in configuration A at $\phi^2 = 40$, Re = 50 and $\gamma = 0.1$. Lines correspond to model and markers correspond to SIM. Red color corresponds to first sphere, blue color to second sphere and green color corresponds to third sphere. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

consequently varies $Pe \in [25, 500]$. Computed $\overline{C_s}$ is compared to $\overline{C_s}$ predicted by the model for the three particles in Fig. 22. The agree ment is deemed to be satisfactory. As a side comment, we addressed in the above gas solid reactive particulate flow prob lems in which the Schmidt number Sc is of the order of 1. For liquid solid systems Sc is (potentially much) larger than 1. The accuracy of the numerical solution with respect to Sc is deter mined by the number of points across the external boundary layer. The external boundary layer thickness for a fixed Re varies as Sc $\frac{1}{3}$, and so for a given numerical precision the mesh size should be accordingly reduced by a factor Sc $\frac{1}{3}$. This limits the range of Sc that can be examined with a constant mesh size approach as our DLM FD/SIM method.

We now plot $\overline{C_s}$ as a function of $\mathbb{R}e \in [10, 100]$ and $\phi^2 \in [0, \infty]$ per particle for γ 0.1 and Sc 1. We compare the computed val ues and values predicted by the model in Fig. 23a for the first sphere, Fig. 23b for the second sphere and Fig. 23c for the third sphere. We also compare the mean volume concentration $\overline{C_v}$ in Fig. 24a for the first sphere, Fig. 24b for the second sphere and Fig. 24c for the third sphere. In general, the agreement between computed values and values predicted by the model is once again deemed to be satisfactory. The Damkohler number ϕ^2 has the most significant impact on the observed difference between computed values and model predictions. For $\phi^2 < 200$, the agreement in very satisfactory for all spheres and the observed difference is very small. Then for larger $\phi^2 > 200$, the observed difference, though still limited, increases with ϕ^2 . The maximum difference of the order of 10% is attained for ϕ^2 4000. As pointed out in the above, this trend is related to the internal concentration boundary layer that decreases with the increase of the Thiele modulus as $1/\phi$. The same comments apply to the plots of the reactive Sherwood numbers given by (62) for the 3 spheres in Fig. 26a for the first sphere, Fig. 26b for the second sphere and Fig. 26c for the third sphere.

The observed differences between computed values of $\overline{C_s}, \overline{C_v}$ or \widetilde{sh} and the values of the same quantities predicted by the model have 3 sources:

- 1. approximations and assumptions adopted to derive the model,
- 2. not fully converged in space computed solutions for large values of ϕ^2 ,
- 3. correlations proposed for the coefficients β_j by Ramachandran et al. (1989) and established by a least square regression with a maximum error of 2.5% for β_1 , 4.6% for β_2 and 10% for β_3 .

In the case of the 3 aligned spheres, the equivalent solid volume fraction around the spheres is low and the system is representative of a dilute regime. The interactions between the 3 spherical obsta cles, both in terms of momentum and mass transfer, are limited, though far from negligible. As a result, the chemical concentration does not vary much along each sphere surface as shown in Fig. 25. Consequently, the approximations (51) and (52) to calculate the mean volume concentration, i.e., assuming that C is a function of the radial coordinate only and $C_s(\theta, \phi)$ does not vary too much around $\overline{C_s}$, are valid. The model is assumed to perform well in this flow configuration and this range of dimensionless parameters. The model validity has been thoroughly checked for a single particle system in our previous study (Sulaiman et al., 2018) where we plotted the internal concentration profiles for many angles and we showed that when concentration profiles are normalized by their local surface concentration, they agree well with the analyti cal solution (38) of the Thiele problem. So we believe that source 1 does not contribution much to the observed differences. The increase of the observed differences as a function of the Damkolher number ϕ^2 is certainly a sign that for high ϕ^2 , N_p 70 points per diameter might not yet be enough to yield fully spatially converged computed solutions as the internal boundary layer is getting thin ner with the increase of ϕ^2 , as suggested by Figs. 7 and 14. Source 2 is thereby a significant contribution to the observed differences at

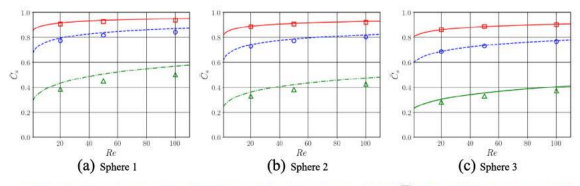


Fig. 23. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: mean surface concentration $\overline{C_s}$ of (a) sphere 1, (b) sphere 2 and (c) sphere 3 as a function of Re in configuration A at γ 0.1 and Sc 1. Lines correspond to model and markers correspond to numerical simulation. Red color corresponds to ϕ^2 40, blue color to ϕ^2 200 and green color corresponds to ϕ^2 4000. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

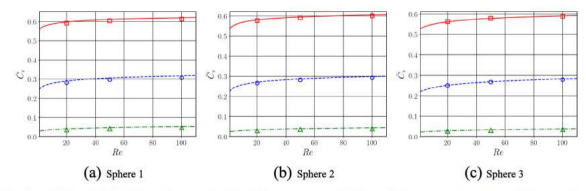


Fig. 24. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: mean volume concentration $\overline{C_{\nu}}$ of (a) sphere 1, (b) sphere 2 and (c) sphere 3 as a function of Re in configuration A at γ 0.1 and Sc 1. Lines correspond to model and markers correspond to numerical simulation. Red color corresponds to ϕ^2 40, blue color to ϕ^2 200 and green color corresponds to ϕ^2 4000. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

high ϕ^2 . Finally, the magnitude of the error on the coefficients β_j as reported by Ramachandran et al. (1989) is of the same order as the observed differences, so we cannot rule out the contribution of source 3. Overall, the model performs well. The difference between computed values and model predictions increases with the Dam kohler number ϕ but is capped to 10% in the range of dimension less parameters investigated.

6. Discussion and perspectives

We presented a numerical framework that couples a Sharp Interface Method (SIM) for the convection diffusion reaction con servation equation of the chemical species to a DLM/FD method to solve the incompressible Navier Stokes equations with fixed obstacles. We presented multiple validation tests of growing com plexity to ensure that our implementation of our DLM/FD SIM method supplies computed solution with the expected accuracy. We have shown through various convergence tests, comparisons to analytical solutions, comparisons to correlations and compar isons to body fitted simulations that our numerical tool indeed supplies reliable computed solutions. We then used our numerical tool to examine the problem of convective diffusive mass transfer in the flow past 3 reactive spheres.

The primary objective of the work was to show that a reactive Sherwood number correlation can be constructed in a dilute sys tem on the basis on a simple external internal coupling and the additivity rule, in a similar way as we constructed a reactive Sher wood number correlation for a single isolated sphere in <u>Sulaiman</u> et al. (2018). Model predictions show a satisfactory agreement

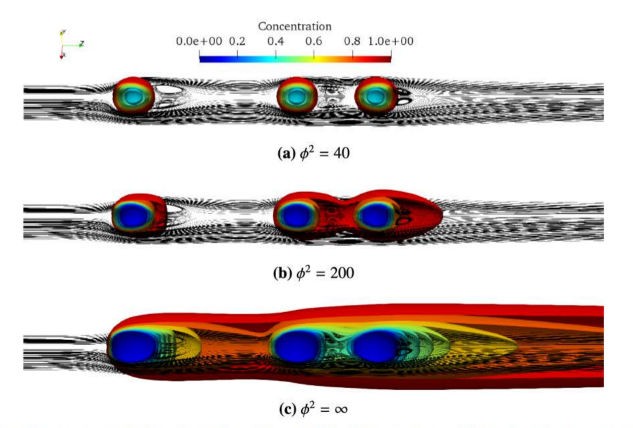


Fig. 25. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: concentration iso-surfaces in configuration A at $Re = 50, \gamma = 0.1, Se = 1$ and different ϕ^2 .

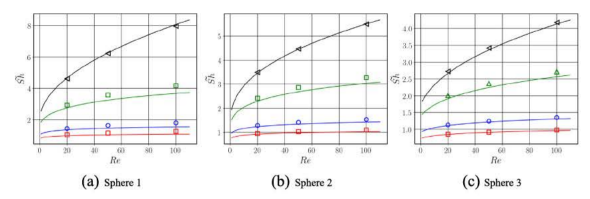


Fig. 26. Steady state convection-diffusion in the flow past 3 aligned reactive spheres: reactive Sherwood number \widehat{Sh} of (a) sphere 1, (b) sphere 2 and (c) sphere 3 as a function of Re in configuration A at γ 0.1 and Sc 1. Lines correspond to model and markers correspond to numerical simulation. Red color corresponds to ϕ^2 40, blue color to ϕ^2 200, green color correspond to ϕ^2 4000 and black color corresponds to $\phi^2 \infty$. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

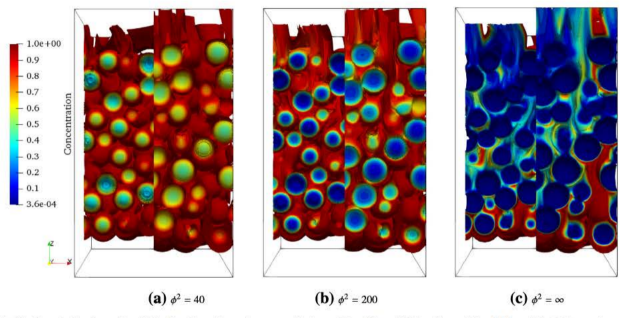


Fig. 27. Concentration iso-surfaces in the flow through a random array of spheres at Re = 25, $\gamma = 0.1$, Se = 1, $\alpha_s = 0.5$ and different Damkohler numbers ϕ^2 .

with our DLM/FD SIM numerical results for wide ranges of the 4 governing parameters in the problem of convective diffusive mass transfer in the flow past 3 reactive spheres. We consider this prob lem as an adequate toy model of a dilute particle laden system experiencing hydrodynamic interactions. The strength of our reac tive Sherwood number correlation is that it is based on any exist ing non reactive Sherwood number correlations. Here we used the non reactive Sherwood number correlations of Feng and Michaelides (2000) and Ramachandran et al. (1989), but this is not mandatory.

The ultimate objective is to extend the suggested reactive Sher wood number correlation to denser regime and eventually to use it in larger scale numerical models as Euler/Lagrange and Euler/Euler modelling of reactive particle laden flows. This can be achieved in two complementary ways. The former way involves investigating pairwise interactions in terms of reactive mass transfer. A model system would be the flow past two spheres not aligned with the flow and then investigate not only the effect of the inter particle distance but also of the relative angular position of the two parti cles, in a similar way to Akiki et al. (2017) for momentum transfer. The latter way involves computing the flow through a random array of reactive spheres up to a high solid volume fraction α_s close to packing, as illustrated in Fig. 27. This would require large com puting resources to span all parameter ranges but is feasible as our code is fully parallel and can run on large supercomputers with a satisfactory scalability. This work is currently under way.

Appendix A. Derivation of the diffusive Sherwood number in a finite spherical domain

We consider a sphere of radius r_p^* at concentration $C C_{s|_{r^*} r_p^*}$ at the center of a spherical domain of radius $r^* r_{\infty}^*$ filled with a qui escent fluid of diffusion coefficient D_f^* . At steady state, the concentration distribution C in the fluid is governed by:

$$D_f^* \nabla^2 C = 0 \tag{64}$$

(64) can easily be integrated with the two Dirichlet boundary con ditions $C \quad C_s|_{r^*} \stackrel{r}{}_{p}$ and $C \quad C_{\infty}|_{r^*} \stackrel{r}{}_{\infty}$. The solution, i.e., the concentra tion profile $C(r^*)$ in the fluid phase, reads:

$$C(r^*) = \frac{r_p^* r_\infty^*}{\left(r_p^* - r_\infty^*\right) r^*} (C_\infty - C_s) + \frac{C_s r_p^* - C_\infty r_\infty^*}{r_p^* - r_\infty^*}$$
(65)

The mass transfer coefficient k^* can be calculated through the calculation of the total flux through the sphere surface as:

$$k^{*} = \frac{D_{f}^{*}}{4\pi r_{p}^{*2}(C_{\infty} - C_{s})} \int_{0}^{2\pi} \int_{\pi/2}^{+\pi/2} \left. \frac{\partial C(r^{*})}{\partial r^{*}} \right|_{r^{*}} r_{p}^{*2} \sin(\theta) d\phi d\theta \qquad (66)$$

Substituting (65) into (66), the diffusive Sherwood number in a finite spherical domain Sh $\frac{k^*d_p^*}{D_*^*}$ can be written as:

Sh
$$\frac{k^* d_p^*}{D_f^*} = \frac{2r_{\infty}^*}{r_{\infty}^* - r_p^*}$$
 (67)

For an infinite domain, i.e., when $r_{\infty}^* \to \infty$, $Sh \to 2$. For a spher ical domain of finite size, Sh is larger than 2 and its value can be calculated by (67).

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