Open Archive TOULOUSE Archive Ouverte (OATAO)

OATAO is an open access repository that collects the work of Toulouse researchers and makes it freely available over the web where possible.

This is an author-deposited version published in: http://oatao.univ-toulouse.fr/
Eprints ID: 16471

To link to this article: DOI: 10.1016/j.ces.2016.10.024
URL: http://dx.doi.org/10.1016/j.ces.2016.10.024

To cite this version: Bacchin, Patrice An energy map model for colloid transport. (2017) Chemical Engineering Science, 158. pp. 208-215. ISSN 0009-2509

Any correspondence concerning this service should be sent to the repository administrator: staff-oatao@listes-diff.inp-toulouse.fr
An energy map model for colloid transport

Patrice Bacchin
Laboratoire de Génie Chimique, Université de Toulouse, CNRS, INPT, UPS, Toulouse, France

A B S T R A C T

When dispersed colloids are flowing, they experience interactions with the fluid (friction) and with other colloids (surface interactions). These phenomena are usually taken into account through a Suspension Balance Model (SBM) that couples mass and momentum balances. However, in many applications, the dispersed particles flow close to an interface or inside a porous media. The flow in such a confined environment leads to significant particle-wall interactions. This paper puts forward an energy map model that accounts for these particle-wall interactions. A way to implement the energy map in the SBM is to introduce an interfacial pressure concept. The new possibilities opened up by the energy map that account for interfacial interaction in the SBM are analysed. A transient 1D case study for the transfer of colloids through a pore illustrates the potentialities of the Suspension Balance Model integrating an Energy Map (SBM-EM). The model enables the description of the transmission of the colloids through the energy map representing the membrane (mass balance) and the consequences in terms of an out-of-equilibrium counter pressure (momentum balance). The counter osmotic pressure is then explained by the interfacial interaction between the colloids and the interface; these interfacial interactions that prevent the colloids from leaving the bulk volume generate forces that are transmitted to the fluid (via the drag force), thus inducing osmosis. The energy map model can enable the incorporation of the physical and chemical heterogeneities of the interacting surfaces. It might be of interest to explore the transfer of colloids along or inside real surfaces (being a mosaic of nano- or micro-scale domains with specific interactions).

1. Introduction

The transport of colloids cannot be described only by classical diffusive and convective mass transport terms. The main reasons are the existence of both surface interactions between the colloids (or between a colloid and its surrounding interface) and hydrodynamic interactions between the particle and the fluid (interactions with the shear rate). These interactions that occur at a nano- or micro-scale are deeply modifying the way in which colloids are diffusing and/or being advected. For example, processes such as ultrafiltration, nanofiltration or reverse osmosis, which are classically used to purify, eliminate and concentrate colloids or nanoparticles, strongly depend on these interfacial phenomena. The level of fouling, its kinetics or even the way colloids build up (porosity, hydraulic resistance or accumulation reversibility) are driven by colloidal properties (Bacchin et al., 2011). Such an impact of surface interactions is also crucial during the transport of drug and carriers in the crowd environment of cells (Al-Obaidi and Florence, 2015); the nano-scale interactions playing a significant role on the hindered diffusion or advection towards cellular goals.

It is therefore necessary to establish experimental and theoretical connections between colloidal properties at a local (micro) scale and the efficiency of the mass and momentum transport phenomena; this knowledge is compulsory for the control of numerous processes that deal with concentrated colloids and/or colloids in confined situations.

In a sheared flow, the colloids are submitted to hydrodynamic interactions (due to the fluid velocity-drag force and to the velocity gradient-shear induced diffusion or lateral migration). Additionally, in a concentrated flow, colloids experience multi-body surface interaction (i.e. DLVO forces, etc.). In these flows, it is crucial to account for the momentum coupling or exchange between the fluid and the particle phase. These interactions (and their coupling) can be taken into account by the Lagrangian approach (like the Force Coupling Method or the Monte Carlo procedure) or by the Eulerian approaches (two fluid model, mixture models, suspension balance model). Multiple inter-particle DLVO interactions have been implemented in the Force Coupling Method in order to depict the collective effect induced by the filtration through a pore (Agbangla et al., 2014). However, this method, based on the tracking of individual particles (around 1 µm), remains impossible to apply for describing the process scale (around 1 m). For this reason, the Eulerian approach that considers the variation of spatial averaged variables, is more adapted for the
description of the transport of concentrated colloidal dispersion. The different hydrodynamic and colloidal forces can be accounted for with two coupled momentum balances for both the particle phase and the fluid phase (called for this reason the two fluid model (Noetinger, 1989)). These momentum balances are coupled by considering momentum exchanges. According to this formulation, the equation can be written for the whole mixture (particles and fluid). It is then called the suspension balance model (Nott and Brady, 1994) or the mixture model (Jackson, 2000, 1997). Both the momentum exchange between the particle and the fluid phases and the slip velocity between the particles and the fluid have to be introduced as closure relationships in order to fully describe the problem. This last Eulerian approach will be introduced in the background section of this paper.

However, the picture is even more complex when the flow of particles takes place in confined conditions, where interactions with the walls are occurring. Furthermore, real surfaces are often chemically heterogeneous on a micro- or a nano-scale (like a biological membrane composed of lipid bilayers with inclusions) and can present local morphological heterogeneity (for example asperities) that can induce different local interaction energies when an object approaches the surface. To account for this complexity, the particle-wall interactions can be accounted for through an energy map. Several authors have defined interactions maps to characterise the approach of colloids near a surface. Interaction maps allows for example, the effect of the roughness, through DLVO calculations (Hoek et al., 2003), to be described. These maps have been used to determine the local equilibrium position that is due to both lateral and normal components of the DLVO force (Kemps and Bhattacharjee, 2005). Comparing the hydrodynamic forces with a DLVO energy map can then help to have a better evaluation of the interactions between colloids and heterogeneous surfaces (Shen et al., 2012). However, this energy map should be integrated in a full transport model in order to account for the coupling with diffusion, advection and hydrodynamic or colloidal interactions.

The aim of this paper is to propose a model that describes the transport of colloids in (or close to) porous media and thus to integrate the description of the transport of concentrated colloidal dispersion. The shear-induced migration was depicted by several authors (arising from the differences in velocity between the particles and the fluid phases), and a stress-induced force, \( \Sigma_m \) (arising from the gradient in the field of velocity). A review of the mixture models for shear-induced migration in flowing, viscous and concentrated particle suspensions have highlighted the possibility of describing the non-equilibrium osmotic pressure and shear-induced diffusion coefficients in the same model formulation (Vollebregt et al., 2010). All these recent developments have been integrated in a revised form of the Suspension Balance Model (Nott et al., 2011) that will be the starting point of the analysis done in the paper.

2. Theoretical background

2.1. The suspension balance model (SBM)

The SBM is based on solving field equations written from the volumic averaging of the governing equations (local momentum and mass balances) on the two phases. These field equations resulting from momentum and mass balances, are written below for the fluid phase, the dispersed phase and the mixture (the balance for the mixture being the sum of the two phases):

**Momentum balance**

For the dispersed phase

\[
\phi \eta \nabla \cdot \vec{F} + \nabla \cdot \left( \Sigma_f \right) = 0
\]

For the fluid

\[
(1 - \phi) \eta \nabla \cdot \vec{F} + (1 - \phi) \nabla \cdot \left( \phi \Sigma_f \right) = 0
\]

For the mixture

\[
(1 - \phi) \nabla \cdot \vec{F} + (1 - \phi) \nabla \cdot \left( \phi \left( \nabla \cdot \vec{F} + \nabla \cdot \left( \Sigma_f \right) \right) \right) = 0
\]

**Mass balance**

For the dispersed phase

\[
\frac{\partial \phi}{\partial t} = -\nabla \cdot (\phi \vec{m}_f)
\]

For the fluid

\[
\frac{\partial (1 - \phi)}{\partial t} = -\nabla \cdot ((1 - \phi) \vec{m}_f)
\]

2.2. A set of closure relationships for colloids

Closure relationships are necessary to close the problem and to be able to determine the fluid properties (the velocity and the volume fraction) from the previous set of equations (Eqs. (1)–(6)). A first closure relationship expresses the drag force as a function of the slip velocity between the particle phase, \( \vec{u}_p \), and the mixture phase, \( \vec{u}_m \):

\[
\vec{F}_{\text{drag}} = -\phi \frac{\vec{u}_p - \vec{u}_m}{V_p (m(\phi))}
\]

where \( m(\phi) \) is the mobility of the particles accounting for the effect of the volume fraction, i.e. \( K(\phi) / \delta_{\text{hs} \phi} \) where \( K(\phi) \) is the hindered settling coefficient.

The writing of the stresses \( \Sigma_f \) and \( \Sigma_m \) is more controversial and a different set of closure relationships have been proposed (as reviewed in Vollebregt et al. (2010)). As underlined by Lhuillier (2009), some of these sets of closure presents some inconsistencies. Clausen (2013) proposes a more consistent formulation: this set of modified closure relationship will be the starting point of the one proposed in this paper.

For low Péclet numbers, the particle-phase stress, \( \Sigma_p \), can be written by considering only the normal stress (NS) contribution (Clausen, 2013). Furthermore, a reasonable premise for colloidal particles at moderate shear rates is to consider the stress as isotropic (Hallez et al.,
The mechanical particle pressure, $\Pi_{c, m} (\phi)$, is the contribution of the particle pressure due to mechanical contact between colloids (inelastic collisions, friction) or between colloids and the fluid (frictional drag on the walls). This contribution can be composed of the shear-induced thermodynamic property (elastic contribution) when the dispersive forces (entropic or electrostatic) overcome the attraction i.e. if $\phi_{\text{crit}} > 0$. If this last derivative is negative (for example when high concentration lead colloids to interact at shorter distances with attractive interactions), a spinodal decomposition occurs. Colloids are no longer spontaneously thermodynamically dispersed: colloids can interact with mechanical interactions and the particle pressure is no longer osmotic (when osmotic is defined as an idealized, spontaneous reversible and non-dissipative process, constituted of a continuous sequence of equilibrium states).

The mechanical particle pressure, $\Pi_{c, m} (\phi, \tau)$, is the contribution of the particle pressure to the viscosity, $\eta$ ($\phi$), and of the thermodynamic osmotic pressure, $\Pi_{m} (\phi)$:

$$\Sigma_{f} = 2\eta_{f} \nu_{f} < \phi > + \Pi_{c, m} (\phi) \psi$$

The particle contribution to the viscosity combines with $2\eta_{f} \nu_{f} < \phi >$ to represent the shear viscosity of the mixture: $\eta_{f} (\phi) = \eta_{f} (1 + \eta_{m} (\phi))$. The thermodynamic osmotic pressure is here accounted as an exchange between the particle and the fluid phases, as proposed by Lhuillier (2011) for the thermodynamic forces within the interphase force.

The addition of Eqs. (8) and (9) (also defined by Batchelor’s, $\Sigma_{f}$, the solid-phase stress) defines the total stress for the mixture (Eq. (3)), thus accounting for a viscos term and a particle pressure term: $\Sigma_{p} + \Sigma_{f} = 2\eta_{f} \nu_{f} < \phi > - \Pi_{c, m} (\phi, \tau) I$

This writing is consistent with the most commonly form (Miller et al., 2009; Morris and Boulay, 1999) for non-Brownian suspensions, accounting for a non-Brownian particle pressure contribution when assuming no significant normal-stress difference.

### 2.3. The set of Eulerian equations to solve

Combining the field equations (Section 2.1) and the modified rheological model enables the set of the Eulerian equation to solve to be defined.

#### Momentum balance

For the dispersed phase

$$\phi (\vec{p}_{f} + \vec{F}_{\text{drag}}) - \nabla \cdot \Pi_{c, m} I - \nabla \cdot \Pi_{c, m} I = 0$$

For the fluid

$$\phi \vec{p}_{f} - \nabla \cdot \Pi_{c, m} I = 0$$

For the mixture

$$\frac{\partial (1-\phi) \vec{p}_{f}}{\partial t} = \nabla \cdot (\phi \vec{v}_{f} - \nabla \cdot \Pi_{c, m} I)$$

The momentum balance for the particle phase (Eq. (11)) permit the drag force to be expressed. The drag force being linked to the slipping velocity (Eq. (7)), it is possible to express the particle velocity as follows:

$$\vec{v}_{p} = \nabla \cdot (\phi \vec{v}_{f} - \nabla \cdot \Pi_{c, m} I)$$

The particle velocity can be implemented in the mass balance for the dispersed phase (Eq. (14)) leading to Eq. (20). This equation has to be solved together with the Eqs. (18) and (19) (the mixture mass balance and the mixture momentum balance respectively) to have the full set of the 3 SBM equations:

$$\nabla \cdot \vec{u}_{f} = 0$$

$$\mu_{f} \vec{v}_{f} - \nabla \cdot \Pi_{c, m} I = 0$$

$$\frac{\partial \phi}{\partial t} = \nabla \cdot (\phi \vec{v}_{f} - \nabla \cdot \Pi_{c, m} I)$$

The solving of these 3 equations then enables the identification of the mixture velocity, $\vec{u}_{f}$, the fluid pressure $p$, and the volume fraction of colloids, $\phi$. The other variable can easily be determined from this 3 variables; for example the particle velocity can be determined from Eq. (17).

#### 2.4. Physical consistencies of the set of equations

It has to be noted that the gradient of the complete particle mixture, $\nabla \phi_{m}$ (combining thermodynamic and mechanic contributions) acts in the mass transport equation (Eq. (20)) as proposed in the classical diffusive flux model (Deboeuf et al., 2009; Leighton and Arrivos, 1987). However, the momentum, due to the gradient of the thermodynamic contribution of the solid pressure, is released in the fluid (Eq. (12)). Consequently, only the gradient of the mechanical part of the particle pressure, $\nabla \Pi_{c, m}$, plays a role on the momentum balance of the mixture (Eq. (13)). This differentiation of the solid pressure action emphasizes the dual behavior of colloids that are both exchanging “thermodynamical” energy with the fluid, due to the Brownian motion (collisions with the liquid molecules) and dissipate “mechanical” energy in the mixture (collisions and friction with the particles). Such a way to write models enables the description of the equilibrium between the static pressure and the osmotic pressure (Eq. (12) leads to $\nabla \cdot (\phi \vec{v}_{f} = \nabla \Pi_{c, m} I)$ when the drag force is zero (at equilibrium). Furthermore, for non-equilibrium conditions, the Eq. (11) describes the equivalence between drag force and the gradient in osmotic pressure as already discussed by Wijmans et al. (1985) and Elimelech and Bhattacharjee (1998) for polarization concentration.

At the end, as schematized in Fig. 1, in this rheological model for the phase stresses, the presence of colloidal particles under a given load
(under a shear rate and under a concentration gradient) contributes to the modification of the momentum exchanges with:

- a dissipative contribution in the fluid phase, \(2V\nabla \Pi \phi (\phi) < \varepsilon > \) due to the particle phase. Such a term contributes to the mixture effective viscosity in the Stokes equation.

- a storage contribution in the solid phase through the gradient in particle pressure, \(-V\cdot \Pi_0 (\phi, \gamma) I\). The particle pressure here accounts for the storage of the energy in concentrated or high-sheared zone. The thermodynamic contribution to this energy \(V\cdot \Pi_{cc} (\phi) I\) can be later released in the fluid phase. The storage is then an elastic contribution (occurring only when a force is applied i.e. non-permanently). The similarity between osmotic pressure and storage modulus of viscoelastic dispersion has been experimentally evidenced (Mason et al., 1997). The other part of the energy storage, due to mechanical interaction \(-V\cdot \Pi_{ff} (\phi, \gamma) I\), can be released in the mixture, where it will be dissipated.

A key feature of this proposed set of modified closure relationships for colloids is the differentiation between the thermodynamic and the mechanic parts of the particle pressure that are acting on the fluid and mixture momentum balances, respectively.

3. Implementation of an energy map in SBM: the SBM-EM model

When colloids flow close to an interface or inside a porous material, each particle experiences an additional force that can result in a force toward the bulk if the interactions are repulsive or toward the surface if the interactions are attractive. It is possible to access this force by performing the derivative of the interaction energy. The colloid/interfacial potential interaction energy (a free energy), \(V_c\), can be calculated for each point of the fluid where the particle can flow, then, constituting an energy landscape: the energy map, \(V_c (x, y, z)\). This energy map can also be associated to a pressure, considering \(V_c = -\Pi_0\), where \(\Pi_0 (x, y, z)\) is the interfacial pressure (that can also be the mapped parameter). The associated force per unit of particle volume is given by the interfacial pressure gradient \(\nabla \Pi_0\). The additional force per unit of volume of fluid is then the product of the volume fraction, multiplied by the gradient of the interfacial pressure:

\[\phi \nabla \Pi_0\]

This term is implemented in the dispersed phase momentum balance (Eq. (11)). The term \(-\phi \nabla \Pi_0 (x, y, z)\) can also be written as \(-\nabla \Pi_{cc} (x, y, z, \phi)\) where \(\Pi_{cc}\) is the contribution of the interface to the colloid bulk pressure, \(\Pi_{cc} (\phi)\).

The new set of equations is given in Supplementary information (S3). When the SBM equation is combined together with the closure relationship (as previously discussed in Section 2.3), a set of three equations (later called SBM-EM for Suspension Balance Model with Energy Mapping) has to be solved.

Conservation of the velocity of the mixture:

\[\nabla \cdot \mathbf{v}_m = 0\]  \[(22)\]

Momentum balance

\[\rho \ddot{\mathbf{v}}_m - \nabla (\Pi_m (\phi)) + 2\nabla \cdot (\phi \mathbf{v}_m (\phi)) \nabla (\Pi_m (\phi)) - \phi \nabla \Pi_m (x, y, z) = 0\]  \[(23)\]

Mass balance

\[\frac{\partial \phi}{\partial t} = -\nabla \cdot (\phi \mathbf{v}_m) - \nabla \cdot (\phi \mathbf{v}_m (\phi)) \nabla \Pi_m (\phi) - \phi \nabla \Pi_m (x, y, z)\]  \[(24)\]

It has to be noted, that even if the physical context is different, this set of equations can show some similarities with the ones used by Jacazio et al. (1972). These describe electro-osmosis where the Nernst-Planck equation is combined with the Navier Stokes equation that includes an electrokinetics term for the exclusion of ions by the stationary phase.

4. The energy map: an ingredient to understand complex mechanisms

In a general way, the interfacial energy introduced in the energy map, \(V_c (\phi, \gamma)\), can be defined as the energy needed to bring the dispersed phase close to the interface. It has to be noted that this interfacial pressure presents some similarities with a disjoining pressure (Derjaguin and Churaev, 1974). This energy can take different forms and thus describe different mechanisms. In the most conventional way, this interaction energy can represent the potential interaction energy between the particle and the interface (for example through particle/wall surface interactions: electrostatic interaction, van der Waals, polymer brush, etc.). However, this energy can also reflect the potential energy that an object must acquire in order to be transported in the map. For example, this energy can represent the stretching energy that is required to deform a plasmid DNA and, then, to enable its penetration inside a channel (Li et al., 2015). This energy can also represent the energy needed for a molecule to be solubilized in a material (as in the diffusion/solubilization model in reverse osmosis). This colloid/interface energy can then help to take into account various types of interaction in a global way: the interaction of the colloid with the interface but also the internal interaction within the colloid (change in configuration or phase change) needed to be transferred.

The consequences of the introduction of the energy map in Eqs. (23) and (24) can first be investigated by analysing the contribution of each term of the SBM-EM model and secondly, by considering the mechanisms that can be described when coupling two of these terms:

- classically, the coupling of the convective term \(u_m \phi\) and the diffusive mass flux \(m(\phi) \nabla \Pi_m\) describes the polarization layer mechanism that can develop in a boundary layer. The effect of the multibody interaction and of the shear induced diffusion can be accounted through the \(\Pi_{cc}\) and \(\Pi_{ff}\) contributions respectively.

- the coupling of the diffusive mass flux \(\nabla \Pi_m\) with the interfacial pressure \(\phi \nabla \Pi_m\) describes the distribution of the concentration with the interfacial pressure: an exclusion if the interfacial pressure is positive (repulsion) or an accumulation if the interfacial pressure is negative (attraction). When considering the distribution of the colloids between two phases having a difference of interfacial pressure \(\Pi\), a partition coefficient, \(K\), can be defined as the ratio between the concentration in the phase (where it exists \(\Pi\)) and the concentration in the bulk phase (where \(\Pi = 0\)). With the limits of an
ideal non-interacting dispersion (if the osmotic pressure follows the Van’t Hoff law), this equilibrium leads to a Boltzmann distribution:

\[
\nabla \Pi_c - \phi \nabla \Pi_l = 0 \Leftrightarrow K = e^{-\frac{\Pi_i}{kT}}
\]

(25)

– the coupling of the convection \( u \phi \) and the interfacial pressure term \( m(\phi) V \nabla \Pi_c \) represents the balance between the hydrodynamic drag force and the particle-wall force

– in the momentum balance, the coupling between the hydrostatic pressure, \(-V(1-\phi)\rho \) and the interfacial pressure, \(-\phi \nabla \Pi_l \), permits the effect of the osmotic pressure on the driving pressure to be described. It will be shown in the next section that this term allows to express the counter pressure due to osmosis in a reverse osmosis process.

The model can also describe the phase transitions that occur when attractive interaction predominate over repulsions (characterized with a zero derivative of the osmotic or the interfacial pressure). A phase transition can occur for a critical volume fraction that corresponds to the zero derivative of the “bulk” osmotic pressure then representing a homogeneous aggregation of particles. For the interfacial pressure, the phase transition is represented by critical coordinates in the energy map that will conduct to a heterogeneous aggregation with particle/wall contact. The model can then describe the critical fluxes (or velocities) conditions leading to these heterogeneous (Bacchin et al., 1995) or homogeneous phase transitions (Bacchin et al., 2011, 2002).

The full SBM-EM model is then able to describe the effect of the multi-body interactions (particle/particle and particle/wall) on both the mass and momentum transports. It enables the local analysis of the dynamic transport of colloidal fluid that account for nanoscale interactions; the energy map enables the description of these interactions but also their possible patchy character (when interactions change along a surface). The tuning of this interacting architecture can be the key point for facilitated transport mechanisms or for non-adhesive surface design (Jiang and Cao, 2010). The model enables the investigation of the complex coupling between the flow and the multi-body surface interactions (in the bulk and the wall) that occur during the flow of concentrated dispersions in confined environment. These potentialities are illustrated in the next section through a simplified case study for the uniaxial transfer of colloids through a membrane.

5. Case study: transient transfer of a colloid through a membrane

The SBM-EM model enable the 2D simulation of the transfer of colloids inside a membrane by accounting the specific interactions between the pore wall and the colloids (Fig. 3 left part). However, in this paper, the simplest case study will be considered. The case study represents the transfer of a solute along the pore axis or through a dense membrane. The problem will then be considered in 1D (the \( z \) direction normal to the membrane surface) with no shear (\( \tau_{xy}(\phi) = 0 \)), and when the particle pressure has only a thermodynamical contribution (\( \Pi_{c,n} = 0 \) therefore corresponds to the absence of deposition). Eqs. (23) and (24) are simplified to the partial differential equations:

\[
-\frac{d}{dz}(1-\phi) + \phi \frac{d}{dz} \Pi_l(z) = 0
\]

(26)

\[
\frac{d\phi}{dt} - u \phi \frac{d}{dz} \left( m(\phi) V \frac{d\Pi_c(z)}{dz} \right) = 0
\]

(27)
where $u_m$ represents the permeate flux and is constant along $z$ (according to Eq. (22)). The interfacial pressure will then be a function of $z$: the non-dimensional distance on the axis normal to the membrane surface.

A continuous differentiable function is used to depict the variation of the interfacial potential along $z$ (the function is given in $S4$ of the Supplementary information). The non-dimensional maximum value of the function, $\frac{\Psi_{\text{max}}}{\Psi_{\text{inlet}}}$, is taken at $-\ln(0.1)$ which leads to a value of 0.1 for the partition coefficient according to the Eq. (25). The interfacial pressure variation allows then to define polarization layers, PL (where interfacial pressure is zero), the exclusion layer, EX (where the gradient of interfacial pressure is localized) and the membrane layer, MB (where the interfacial pressure is maximum). All of the simulation data, together with the osmotic pressure (fully described in Bacchin et al. (2006)) are given in the Supplementary information ($S4$).

To perform simulations, the boundary conditions are classically defined with a constant volume fraction at the inlet, $\phi_i = 0.001$, and a non-diffusive flux at the outlet. The initial condition is a nil volume fraction all along $z$. The simulations are coded with python language (Canopy Enthought package) by using the fipy partial differential equation solver (Guyer et al., 2009). The full code is available on request.

The results of the transient simulation for a mixture velocity, $u_m$ of $10^{-5}$ m/s are presented in Fig. 4. These conditions correspond to a Péctel number of 0.46, when defined as $u_m D_o / \delta_0$ with $\delta_0$ being the thickness of the boundary layer thickness ($10^{-6}$ m) and $D_o$, the diffusion coefficient in dilute conditions ($2.18 \times 10^{-11}$ m$^2$/s). The simulations are continuously depicting the polarization layer formation induced by the membrane surface exclusion, the convection-diffusion balance inside the membrane and the exclusion at the extremities of the membrane. At steady state, the volume fraction is, $\phi_f = 0.256$, in the permeate side leading to a solute retention, $R = 1 - \frac{\phi_f}{\phi_i} = 0.744$.

The description of the transfer through a membrane is classically described by an analytical relationship (see $S3$ of the Supplementary information), based on (1) the coupling of a convection-diffusion balance inside the membrane, and (2) the coupling in the boundary layer, and (3) a partition coefficient (with a concentration discontinuity) between the membrane and the outside solution. Simulations have been performed for different permeate velocities (or Péctel number) and the solute retention at steady state is compared to this analytical model in Fig. 5. The analytical model appears with two dashed lines due to the fact that the Pe numbers used in the analytical equation can be written differently depending on whether the exclusion layers are accounted for in the membrane thickness or in the boundary layer thickness. The simulation results are close to the result obtained with the analytical expression. The small discrepancies could result from the way the partition is accounted for: with a ramp of potential in the simulation and with a discontinuity in the concentration profile in the analytical expression. The effect of the slope of the ramp, that could explain the facilitated transfer observed when working with pores that exhibit a conic shape of an hour glass shape (Gravelle et al., 2013; Li et al., 2015), will be further investigated.

Solving of the mass balance (Eq. (18)) with the 1D energy map (Fig. 3) enables a description of the main transport phenomena that occurs through a membrane, with:

- the convection-diffusion balance inside the membrane that explains the increase in the retention for a small Péctel number, usually observed in reverse osmosis process (when the convection enables the reduction of the negative impact of the diffusion on the selectivity)
- the polarization layer in the boundary layer near the interface that leads to a decrease in retention for a higher Péctel number, usually observed in ultrafiltration (when the increase in concentration at the membrane interface favors the transmission of the solute in the permeate).

From the integration of the momentum balance (Eq. (20)), it is also possible to determine the counter pressure that will be opposed to the difference in static pressure. The counter pressure, $CP$, can be determined by integrating the momentum balance term, $\int \phi d\Pi_c$. The counter pressure has been determined for the simulations performed for different Péctel numbers. The results are presented in Fig. 6 where the counter pressure is plotted as a function of the difference between the maximum of the osmotic pressure in the concentrate and the osmotic pressure in the permeate, $\Pi_{c\text{, max}} - \Pi_{c\text{, perm}}$ (that is classically used for the counter pressure estimation). The values obtained from the integration of the interfacial pressure and from the “bulk” osmotic pressure difference are similar. The discrepancy can be explained from the writing of the counter pressure, $CP$, obtained through the integration of the momentum balance for the particle phase (Eq. (S3-1)) along the exclusion layers and the membrane layers:

$$ CP = \int_{EX+MB} \phi d\Pi_c = \Delta \Pi_c + \int_{EX+MB} n\Pi_{c\text{, osm}} dz $$

(28)

This relationship links the counter pressure to the osmotic pressure difference and to the drag force along the exclusion and the membrane layers. Assuming the absence of an additional contribution of a compressive yield stress $\Pi_c\text{, cry}$, (that could represent the pressure drop in a deposit layer), the difference in particle pressure, $\Delta \Pi_c$, is mainly due to the thermodynamic contribution, $\Delta \Pi_c\text{, th}$. The counter pressure is

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure4.png}
\caption{The transient variation of the volume fraction, $\phi$, along $z$ (the vertical lines represent the membrane thickness-full lines and the exclusion layers-dashed lines). The simulation are continuously describing the polarization layer formation, the exclusion near the membrane inlet, the convection-diffusion inside the membrane, the exclusion near the membrane outlet and the filtrate side.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure5.png}
\caption{Membrane retention calculated as a function of the Pe number. The line of symbols represent the simulation results. These results are compared to the ones obtained with an analytical expression, based on a partition coefficient.}
\end{figure}
therefore linked to the difference in osmotic pressure, ΔΠ_{c, th}, at the interfaces between the exclusion and the bulk that are very close to the maximum osmotic pressure and the permeate osmotic pressure (Fig. 4):

\[ \int_{EX+MB} \phi dΠ_{c} = Π_{c, th \, max} - Π_{c, th \, per} + \int_{EX+MB} n_0 dζ \]  

(29)

The difference between the calculated counter pressure and the osmotic pressure difference is due to the drag force that can be seen as the out-of-equilibrium contribution to the counter pressure. This contribution highlights the gap with the spontaneous reversible process (due to the osmotic pressure difference) that is valid only if considering a sequence of equilibrium states (Peppin et al., 2005). This gap is negative when u_m < u_p (for low Péclet numbers) and becomes positive when u_m > u_p. The Peclet number for which the curve crosses the bisector can be associated to the critical Péclet number, i.e. when u_m = u_p. For lower Péclet number, the discrepancy can be described via a Staverman coefficient, \( σ \) of 0.8 (when the counter osmotic pressure is written, \( σ(Π_{c, th \, max} - Π_{c, th \, per}) \)).

The SBM-EM model can enable having a good description of both the concentration profile and the solute transmission through the membrane (Fig. 5) and the impact of the accumulation on the out-of-equilibrium counter osmotic pressure (Fig. 6). The counter osmotic pressure is therefore explained by the interfacial interaction between the colloids and the interface. These interactions prevent the colloids from leaving the volume, which then lead to a modification of the movement of the particles at the interface that becomes non-isotropic near the interface. The colloids exchange momentum with the interface and a part of this momentum is returned to the dispersed phase (via the energy principle of action and reaction), thus generating additional gradient in colloid pressure. This gradient of particle pressure, induced by the interface, generates a force that is transmitted to the fluid (via the drag force), that result in osmosis. The model can also describe the shear-induced mechanisms and the formation of gel or deposit layer at the membrane surface (Bacchin et al., 2002). Another interesting aspect of this model lies in the fact that the energy map can easily be adapted to describe more complex transfer problems that can be encountered in membrane processes.

6. Conclusions

The concept of the energy map is implemented in the Suspension Balance Model (SBM) in order to account for transport phenomena due to particle/wall interactions. This Eulerian model can then simultaneously describe the effect of multibody particle-particle interactions (through the osmotic pressure or the generalized concept of particle pressure) and the particle-wall interactions (through an interfacial pressure relative to the free energy due to the colloidal-wall interaction). The interfacial pressure is a key parameter that characterizes the particle/wall interaction: the energy map represents the value of the interfacial pressure spatially. The gradient in the energy map leads to additional terms of transport phenomena (implemented in the mass balance) and of momentum exchange (implemented in the momentum balance). A new set of equation has been established for this full model (SBM-EM). The coupling of the additional new terms enables accounting for the Boltzmann exclusion of colloids near the interfaces, the presence of heterogeneous critical flux (when the advection leads to overcome the map energy barrier) and the osmotic flow (through the momentum exchange term). The model is applied for a 1D modeling of the transfer through a membrane. It enables the description of the retention of a solute through a membrane (i.e. through the polarization layer, the exclusion layers and inside the membrane) and the determination of the out-of-equilibrium counter pressure (that can be seen as the direct consequence of the colloid’s interaction with the semi-permeable membrane wall). The model is compared to the analytical relationships that exist for 1D problem. At the end, the SBM-EM model enables the representation of the interaction of the colloids with their environment (for example when the dispersion flows in a confined media). The energy map can allow flexibility in incorporating the physical and chemical heterogeneities of the interacting surfaces. It might be of interest to explore the transfer of colloids along or inside real surfaces (being a mosaic of nano or microscale domain with specific interactions).

Acknowledgements

The author thanks Yannick Hallez, Martine Meireles and Pierre Aimar for the fruitful discussions on this modeling.

Appendix A. Supplementary information

Supplementary information associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.ces.2016.10.024.

References


Fig. 6. The counter osmotic pressure deduced from the simulation, \( \int \phi dΠ_{c} \), as a function of the counter osmotic pressure determined through the difference in osmotic pressure between the maximum of the osmotic pressure in the polarization layer and the osmotic pressure in the permeate, \( Π_{c, max} - Π_{c, per} \). The arrow on the dashed line indicates the direction of the increasing Péclet.

The counter osmotic pressure is therefore explained by the drag force induced by the interface (Fig. 4). This contribution highlights the gap with the spontaneous reversible process (due to the osmotic pressure difference) that is valid only if considering a sequence of equilibrium states (Peppin et al., 2005). This gap is negative when \( u_m < u_p \) (for low Péclet numbers) and becomes positive when \( u_m > u_p \). The Péclet number for which the curve crosses the bisector can be associated to the critical Péclet number, i.e. when \( u_m = u_p \). For lower Péclet number, the discrepancy can be described via a Staverman coefficient, \( σ \) of 0.8 (when the counter osmotic pressure is written, \( σ(Π_{c, max} - Π_{c, per}) \)).

The SBM-EM model can enable having a good description of both the concentration profile and the solute transmission through the membrane (Fig. 5) and the impact of the accumulation on the out-of-equilibrium counter osmotic pressure (Fig. 6). The counter osmotic pressure is therefore explained by the interfacial interaction between the colloids and the interface. These interactions prevent the colloids from leaving the volume, which then lead to a modification of the movement of the particles at the interface that becomes non-isotropic near the interface. The colloids exchange momentum with the interface and a part of this momentum is returned to the dispersed phase (via the energy principle of action and reaction), thus generating additional gradient in colloid pressure. This gradient of particle pressure, induced by the interface, generates a force that is transmitted to the fluid (via the drag force), that result in osmosis. The model can also describe the shear-induced mechanisms and the formation of gel or deposit layer at the membrane surface (Bacchin et al., 2002). Another interesting aspect of this model lies in the fact that the energy map can easily be adapted to describe more complex transfer problems that can be encountered in membrane processes.