On the validity of effective formulations for transport through heterogeneous porous media

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Abstract

Geological heterogeneity enhances spreading of solutes, and causes transport to be anomalous (i.e., non-Fickian), with much less mixing than suggested by dispersion. This implies that modeling transport requires adopting either stochastic approaches that model heterogeneity explicitly or effective transport formulations that acknowledge the effects of heterogeneity. A number of such formulations have been developed and tested as upscaled representations of enhanced spreading. However, their ability to represent mixing has not been formally tested, which is required for proper reproduction of chemical reactions and which motivates our work. We propose that, for an effective transport formulation to be considered a valid representation of transport through Heterogeneous Porous Media (HPM), it should honor mean advection, mixing and spreading. It should also be flexible enough to be applicable to real problems. We test the capacity of the Multi-Rate Mass Transfer (MRMT) to reproduce mixing observed in HPM, as represented by the classical multi-Gaussian log-permeability field with a Gaussian correlation pattern. Non-dispersive mixing comes from heterogeneity structures in the concentration fields that are not captured by macrodispersion. These fine structures limit mixing initially, but eventually enhance it. Numerical results show that, relative to HPM, MRMT models display a much stronger memory of initial conditions on mixing than on dispersion because of the sensitivity of the mixing state to the actual values of concentration. Because MRMT does not restitute the local concentration structures, it induces smaller non-dispersive mixing than HPM. However long-lived trapping in the immobile zones may sustain the deviation from dispersive mixing over much longer times. While spreading can be well captured by MRMT models, in general non-dispersive mixing cannot.
1 Introduction

Transport is anomalous in heterogeneous porous media. Anomalous transport observations include tailing in concentration breakthrough curves and plumes, or the strong increase in the rate of spreading of plumes. Several frameworks have been developed to generalize the Advection Dispersion Equation (ADE) and overcome its limitations [Fripiat and Holeyman, 2008]. All these alternative frameworks share the goal to model complex permeability, velocity and concentration patterns in unified parsimonious effective equations. The limited number of parameters makes them efficient for the limited quantity of data usually available. In fact, they can be parameterized from breakthrough curves. They comply with the broad residence time distributions and non-local transport processes observed in reality [Gjetvaj et al., 2015; Le Borgne and Gouze, 2008; Willmann et al., 2008]. They represent the consequences of complex concentration patterns, of simultaneous concentration trapping and fast progress on residence times while averaging out all the fine concentration structures in the upscaling process. These anomalous transport frameworks have proven to be highly effective for residence times, transport time distribution and effective spreading both phenomenologically and practically [Berkowitz et al., 2006; Neuman and Tartakovsky, 2009]. However, their ability to reproduce mixing, which is required for properly reproducing chemical reactions, has not been tested.

We argue that an effective transport formulation should honor not only the mean advection, and spreading observed in Heterogeneous Porous Media (HPM), but also the evolution of mixing. This should not be understood as limiting anomalous transport frameworks but at extending them to handle broader ranges of physical and chemical processes, and at further promoting the approach of effective equations that upscale out the fine scale structures to retain only their main consequences in terms of transport, reactivity and reactive transport couplings.

Here, we investigate the relevance of Multi-Rate Mass Transfer (MRMT) framework to model not only spreading but also mixing. MRMT is taken as a typical anomalous transport framework. Its advantage lies in providing local concentrations, which can be straightforwardly used to evaluate concentration variance, mixing and mixing induced reactivity [Babey et al., 2014; Carrera et al., 1998; de Dreuzy et al., 2013; Haggerty and Gorelick, 1995], as well as the apparent reduction in the rate of kinetic reactions [Dentz et al., 2011]. The question is whether its validity as a representation of transport through
heterogeneous porous media (HPM) can be extended to reproduce the effects of the evolution of mixing rates resulting from the stretching and folding associated to complex velocity structures [de Anna et al., 2014b; Jimenez-Martinez et al., 2015; Le Borgne et al., 2015].

This comparison is especially appropriate as anomalous transport processes are currently extended to simulate reactive transport processes [Cirpka and Valocchi, 2007; Clement, 2001; de Barros et al., 2012; Donado et al., 2009; Hochstetler et al., 2013; Luo et al., 2008; Luo and Cirpka, 2011; Orgogozo et al., 2013; Schneider et al., 2013]. They deal with chemical reactivity either in a stochastic manner, representing reactivity with molecular analogies, or in classical approaches by means of concentrations [Bolster et al., 2010; Cirpka et al., 2012; Ding et al., 2013; Hayek et al., 2012; Knutson et al., 2007; Zhang et al., 2013]. Extensions are both required for applications purposes and attractive for capturing the consequences of anomalous transport to potential "anomalous" and enhanced reactivity [Battisti et al., 2009; Sadhukhan et al., 2014; Scheibe et al., 2015; Tartakovsky et al., 2009].

Some assessment of MRMT to model reactivity in HPM has been made in former works [Willmann et al., 2010]. Equivalent reactivity has been evaluated at some well-defined travel distances on MRMT calibrated on residence time distributions. Here we follow a different avenue by analyzing the temporal development of spreading and mixing. We extend the integrated assessment of mixing-induced reactivity at given travel distances to its temporal development.

Our contribution concerns the comparison of different models much more than the HPM and MRMT model themselves. For the sake of completeness, we recall model equations and simulation methods in section 2 (models and methods) and measures of spreading and mixing in section 3. We use these measures to propose the conditions that should be met by effective (upscaled) transport formulations to be considered valid representations of transport through heterogeneous porous media (Section 4). We then test whether MRMT formulations meet the proposed conditions (Section 5). While this last section depends on the specific choice of the MRMT framework as an equivalent transport model, the comparison methodology is independent of it and can be used to assess transport equations respecting both spreading and mixing.
2 Model and Methods

We present sequentially the Multi-Rate Mass Transfer (MRMT) and Heterogeneous Porous Media (HPM) models. As they are both well known, we present only the main equations and highlight the critical assumptions of importance in this study.

2.1 Multi-Rate Mass Transfer model (MRMT)

Multi-Rate Mass Transfer models express anomalous transport by the interaction between transport in a mobile zone and a series of immobile zones \cite{Carrera et al., 1998; Haggerty and Gorelick, 1995}. Transport in the mobile zone is advective and dispersive with a mean solute velocity $v$ (water flux divided by mobile porosity, $\phi$) and a dispersion coefficient $d$. Each immobile zone $i$ is parameterized by a characteristic rate $\alpha_i$ (inverse of a characteristic exchange time) and an immobile porosity $\phi_i$. The concentrations $c$ and $c_i$ ($i=1..N$) in the mobile and immobile zones, respectively, are determined by the following set of equations:

$$\phi \frac{\partial c}{\partial t} + \sum_{i=1}^{N} \phi_i \frac{\partial c_i}{\partial t} = -q \frac{\partial c}{\partial x} + d \frac{\partial^2 c}{\partial x^2}. \tag{1}$$

$$\frac{\partial c_i}{\partial t} = \alpha_i (c - c_i) \text{ for } i = 1,..,N \tag{2}$$

The ratio of immobile to mobile water volumes is rated by the total capacity ratio $\beta = \sum \phi_i / \phi$. The term capacity derives from the fact that MRMT formulations were originally devised to represent trapping by sorption in hard-to-reach sorption sites, which were characterized by capacity (including both dissolved and sorbed solute mass) \cite[see, e.g., Haggerty and Gorelick, 1995]{Haggerty and Gorelick, 1995}. We use here an equivalent MRMT formulation for non-sorbing solutes, so as to facilitate comparison with HPM.

Initial and boundary conditions will be described later for both MRMT and HPM models. MRMT models differ by the distributions of characteristic rates $\alpha_i$ and immobile porosities $\phi_i$. Among the available models \cite[Cvetkovic, 2012; Haggerty et al., 2000]{Cvetkovic, 2012; Haggerty et al., 2000}, we choose a uniform distribution for characteristic times (1/$\alpha_i$) bounded by the two extreme rates $\alpha_1=1/t_1$ and $\alpha_N=1/t_N$ ($t_1 < t_N$) and a power-law distribution for $\phi_i$:

$$\phi_i \sim \alpha_i^{m-3}. \tag{3}$$
The power-law distribution is consistent with observed breakthrough curves in Heterogeneous Porous Media, which often display long tails that appear linear in log(c) versus log(t) [Gouze et al., 2008; Haggerty et al., 2004; Li et al., 2011; Silva et al., 2009; Willmann et al., 2008]. This tailing is well modeled by a power law, such that the breakthrough concentration $c_t$ evolves as $c_t \sim t^{-m}$. Haggerty et al. (2000) showed that the slope $m$ relates to the exponent of the power-law distribution of the MRMT rates (equation (3)). $m$ is generally found to be in the interval [1.5;2.5] but little is known about its relationship to the geological heterogeneity. Willmann et al (2008) found some correlation between the degree of connectivity and the slope. The more connected the field, the smaller the slope. In this context, fracture/matrix exchanges in fractured media represent the lowest bound ($m=1.5$), which is controlled by diffusion into immobile regions [Haggerty and Gorelick, 1995]. On the contrary, a slope $m$ of 2.5 may represent a heterogeneous but poorly connected hydraulic conductivity field, where late time arrival is controlled by slow advection.

We simulate MRMT models with a standard time- and space-adaptative method that preserves mass [de Dreuzy et al., 2013] and always complies with the CFL conditions [Daus et al., 1985]. The advective and the diffusive processes in the mobile zone as well as the exchange with the immobile zones are treated with a sequential non-iterative coupling method. These methods lead to efficient simulations of large spatial domains and extended times with initial refined resolutions. We have successfully compared them with a more classical fixed-time Galerkin finite element methods integrated with the 4th order Runge-Kutta method (ode45 function of Matlab) and found relative differences less than $10^{-3}\%$. Simulations have been performed over the time required for transport to reach its asymptotic regime.

### 2.2 Heterogeneous Porous Media (HPM)

For reference purposes, we restrict the analysis to heterogeneity of hydraulic conductivity ($K$) as represented by the classical 2D Gaussian correlated multi-Gaussian log-$K$ fields. These are characterized by their isotropic correlation function:

$$C(r) = \sigma^2_r \exp \left( -\left( \frac{r}{\lambda} \right)^2 \right) \quad (4)$$
with \( r \) the distance, \( \lambda \) the correlation length, which is used to scale distances, and \( \sigma_Y^2 \) the variance of the logarithm of \( Y = \log K \). We use simulation results performed in previous studies [de Dreuzy et al., 2012] obtained on 2D domains of sizes \( L_L \) and \( L_T \) in the direction parallel and orthogonal, respectively, to the mean flux. \( L_L \) is large enough to avoid any finite-size effects (from \( 10^2 \) to \( 10^3 \) correlation lengths \( \lambda \)). Boundary conditions for flow and transport are periodic in the transverse direction to minimize boundary effects. \( L_T \) is of the order of 100 times \( \lambda \) to ensure initially ergodic transport conditions. Under such uniform extended injection conditions, transport in HPM can be considered ergodic and can be fundamentally compared with a 1D MRMT model. The immobile zones of MRMT can be viewed as representing the low velocity zones of HPM, so that the mobile zone may represent the high velocity channels.

Flow is solved with a finite volume scheme with permeameter-like boundary conditions under a unit head gradient. Transport is simulated using the ADE, with heterogeneous advection and homogeneous diffusion. Therefore, it is characterized by the Peclet number \( Pe \) equal to the mean velocity times the correlation length divided by the diffusion coefficient. Transport is simulated with a random walk Lagrangian method. Numerical methods are exhaustively described in several previous papers [Beaudoin et al., 2006; Beaudoin et al., 2007; Beaudoin et al., 2011].

### 2.3 Injection and boundary conditions

The same type of injection and boundary conditions are used for both models. Flow has a major flow direction imposed in HPM by a head gradient in the longitudinal direction and periodic boundary direction in the transverse direction. For transport, reflecting and absorbing boundary conditions are used respectively upstream and downstream [Beaudoin and de Dreuzy, 2013]. Injection is performed downstream to the inlet boundary to minimize boundary effects.

Extended injection conditions are used for the HPM and MRMT models. Concentrations are homogeneous orthogonally to the main flow direction within a square wave of longitudinal and transverse widths \( \Delta L_0 \) and \( \Delta T_0 \), respectively. In the HPM case, concentration is a sole function of the coordinate \( x_L \) along the flow direction:

\[
c(x, t = 0) = c_0(x_L)
\]

with \( c_0 \) given by:
\[ c_0(x) = \begin{cases} \frac{m_0}{\phi \Delta T_0 \Delta L_0} & \text{if } x_0 < x < x_0 + \Delta L_0 \\ 0 & \text{otherwise} \end{cases} \]  

(6)

\[ \phi_T \] is the total porosity. To ensure that the same mass \( m_0 \) is injected in the HPM and MRMT cases, we adapt the initial state of the MRMT model to:

\[ c(x, t = 0) = c_i(x, t = 0) = c_0(x) \text{ for } i = 1..N. \]  

(7)

Spreading becomes independent of the injection length when the longitudinal plume size becomes significantly larger than \( \Delta L_0 \). Mixing depends more critically than spreading on the injection conditions, as the initial concentration value depends on the injection width \( \Delta L_0 \) (equation (6)).

3 Measures of spreading and mixing

3.1 Spreading

For an extended plume, spreading is generally measured by the square root of the second centered moment of the spatial distribution of concentration \( \sigma_L \):

\[ \sigma_L(t) = \sqrt{m_L^{(2)}(t) - m_L^{(1)}(t)} \]  

(8)

where \( m_L^{(k)}(t) \) is the k-th order moment of the concentration distribution

\[ m_L^{(k)}(t) = \int_{\Omega} x_L^k c(x, t) d\Omega x / \int_{\Omega} c(x, t) d\Omega x \]  

(9)

with \( x_L \) the coordinate of \( x \) in the direction parallel to the main flow direction (longitudinal direction) and \( \Omega \) the flow domain. With this definition, \( \sigma_L \) can be viewed as the longitudinal extent of the plume (i.e., how far it spreads). Dispersion is the rate of spreading (i.e., time derivative of \( \sigma_L^2 \)), usually characterized by the longitudinal dispersivity \( \alpha_L \) :

\[ \alpha_L = \frac{1}{2v} \frac{d\sigma_L^2}{dt}. \]  

(10)

where \( v \) is the plume velocity equal to the time derivative of the mean position plume \( m_L^{(1)}(t) \).

\( \alpha_L \) increases until it converges to an asymptotic value \( \alpha_L A \), thus defining in turn the asymptotic regime [Dagan, 1990; Gelhar, 1993].
In MRMT, spreading comes from the exchanges to the mobile zone. That is, spreading results from trapping. Solutes are slowed down and dispersed by the exchanges with the immobile zones. The resulting dispersivity is a monotonically increasing function of the residence times in immobile zones (both their mean $\langle \tau_{\text{MRMT}} \rangle$ and range $(t_n - t_l)$). The dispersivity induced by the dispersive and diffusive processes in the mobile zone is comparatively negligible and could be disregarded.

In HPM, spreading comes both from diffusive exchanges with low velocity zones and from spatial fluctuations of the velocity field [de Dreuzy et al., 2007; Salandin and Fiorotto, 1998]. The asymptotic dispersivity increases both with the correlation length $\lambda$ and with the log-K variance $\sigma^2_Y$:

$$\alpha_{\lambda\lambda}(\text{HPM}) = \lambda \left[ g(\sigma^2_Y) h(\sigma^2_Y, Pe) \right]$$  \hspace{1cm} (11)

where $g$ is either a linear function for small values of $\sigma^2_Y$ ($\sigma^2_Y < 1$) and a quadratic function at larger values [de Dreuzy et al., 2007]. $h(\sigma^2_Y, Pe)$ is a correction factor accounting for diffusion [Beaudoin et al., 2010]. Local diffusion reduces the effective dispersivity in the high heterogeneity cases by releasing solutes from the low velocity zone and truncating the trapping times induced by slow advection.

Any concentration plume can be approximated by a Gaussian concentration profile $c_D(x,t)$, defined by the two first moments, $m^0(t)$ as mean and $\sigma^2(t)$ as variance. It is the smoothest equivalent profile. Both MRMT and HPM converge asymptotically to this profile. However, it is far away from the full concentration profile $c(x,t)$ at any time as shown by the comparison of Figure 1. At early times (left snapshots on Figure 1), the concentration profile remains heterogeneous especially in the transverse direction with both higher and lower concentrations. Around the advection time, defined as the correlation length $\lambda$ divided by the plume velocity $v$, the deviation reaches its maximum. At this point, the Gaussian concentration profile has become much more diluted than the real concentration field (second from the left snapshot of Figure 1). Concentration inhomogeneities decrease very slowly and remain over very long times even though the range of concentration values decreases (two right-most snapshots of Figure 1) [de Anna et al., 2014a; Jimenez-Martinez et al., 2015; Le Borgne et al., 2011].
In summary, in HPM, dispersivity comes primarily from the velocity structure, which drives the generation of gradients in concentration and thus mixing. Instead, in MRMT, effective dispersivity is controlled by mobile-immobile exchanges and delays the actual mixing between the immobile and mobile solute concentrations.

3.2 Mixing

The Gaussian profile only gives a crude approximation of the concentration field with a strong deviation on the distribution of concentration values, especially at early times when diffusion has not homogenized the concentration field in the transverse direction (Figure 1). Actual concentrations remain much higher and closer to the initial concentration value than in the Gaussian profile prediction. That is the initial concentrations are much less diluted (i.e., mixed) than in the maximum entropy Gaussian distribution. The Gaussian profile $c_D(x,t)$ thus sets a lower bound to the effective concentration variability. Therefore, it is most natural to compare the actual distribution of concentration values to that of the Gaussian profile in order to describe the mixing state. Notice that, contrary to spreading, we are not concerned here with the spatial distribution, but only with the values of concentration and their time evolution, which are most simply characterized by the second moment. We quantified the deviation from the Gaussian mixing regime as the ratio of the actual concentration second moment $M(t)$ to the second moment $M_D(t)$ of the Gaussian profile concentration $c_D(x,t)$ minus 1 [de Dreuzy et al., 2012]:

$$\gamma(t) = \frac{M(t)}{M_D(t)} - 1.$$  \hspace{1cm} (12)

with

$$M(t) = \int_{\Omega} c^2 d^d x$$ \hspace{1cm} (13)

and the second moment of the reference Gaussian concentration:

$$M_D(t) = \frac{m_0^2}{2 \sqrt{\pi} \Delta T_0 \sigma_L}.$$ \hspace{1cm} (14)

$M_D$ is directly the square of the injected mass $m_0^2$ divided by an effective area occupied by the plume $2 \sqrt{\pi} \Delta T_0 \sigma_L$. As $M(t)$ is always larger than $M_D(t)$, $\gamma$ is always positive. $\gamma$ is initially and asymptotically very close to zero. It is however significantly positive while the
concentration distribution is far from the Gaussian profile. \( M(t) \), which we have introduced here as a measure of global concentration variability, is a widely used measure, as its time derivative, dissipation rate, determines the physical constrains of chemical reactivity [de Simoni et al., 2005; Le Borgne et al., 2010]. The dissipation is also closely related to the dilution index, which is another measure of mixing [Kitanidis, 1994; Rolle et al., 2009]. It should be finally noticed that \( \gamma \) and \( M_D \) fully characterize the mixing state given by \( M \):

\[
M = M_D (1 + \gamma).
\]

In HPM models, resistance to dispersive mixing, as we can also call \( \gamma \), is enhanced by heterogeneity and reduced by large diffusion rates (smaller Peclet number) [de Dreuzy et al., 2012]. \( \gamma \) sharply increases at initial times to a maximum value \( \gamma_{\text{max}} \), at a time \( t_{\gamma\text{max}} \) close to the advection time, and slowly decreases back to 0 (Figure 2). The time range over which \( \gamma \) is significantly non-zero can be characterized by \( r_{\gamma} \), which is the ratio of the upper and lower times at which \( \gamma \) is equal to a quarter of its maximal value \( \gamma_{\text{max}} \). While the amplitude of \( \gamma \) depends on the variability of the velocities and on the rate of advection to diffusion, the shape of the function \( \gamma \) remains unchanged by the \( K \) field heterogeneity \( (\sigma_Y^2) \), the ratio of advection to diffusion \( (Pe) \), and the width of the initial conditions \( (\Delta L_0) \). The time range \( r_{\gamma} \) over which \( \gamma \) is non-negligible also remains constant (Figure 2). Therefore, \( t_{\gamma\text{max}} \) can be used for scaling time, so that \( \gamma \) can be written as:

\[
\gamma(t) = \gamma_{\text{max}} f \left( \frac{t}{t_{\gamma\text{max}}} \right)
\]

where \( f \) is the characteristic scaling function (Figure 2, insert). A similar constant shape behavior has been noted for viscous fingering in heterogeneous velocity fields [Jha et al., 2011a; b].

4 Conditions for effective formalisms of transport through HPM

We propose four conditions for any effective transport formulation to be considered as a valid representation of transport through heterogeneous media. In essence, an effective transport equation should yield the same mean advection, spreading and mixing as the HPM and be sufficiently flexible to represent real problems. Evaluation of these conditions can be done as follows:
(1) **Mean advection** simply requires mean water velocity (i.e., mean plume velocity for non-reactive solutes) to equal \( v = q / \phi_t \). This condition can be met by all published upscaled transport equations, by imposing some simple constrains on their parameters. In MRMT, it is sufficient to impose \( \phi_t = \phi + \sum \phi_i = \phi (1 + \beta) \).

(2) **Spreading** is characterized by dispersivity, which measures the rate of growth of plume size (equation (10)). In cases where asymptotic dispersion is reached, this condition implies that dispersivity of the effective equation should tend to the asymptotic dispersivity of the HPM. Otherwise, dispersion (or directly, spread, as measured by \( \sigma_L \)) can be compared for a spatial scale comparable to the problem dimension (e.g., size of the aquifer, or distance covered by the plume).

In addition, the time required to reach the above dispersion value should also be honored by the effective formulation to ensure that the rate of growth of the plume is reproduced. In our case, where asymptotic dispersion is reached, we propose to define this criterion in terms of \( r_\alpha \), mean distance covered by the plume at the time \( t_{\alpha_{L_A}/2} \) where dispersivity reaches half of its asymptotic value normalized by the asymptotic dispersivity \( \alpha_{L_A} \):

\[
r_\alpha = \frac{v t_{\alpha_{L_A}/2}}{\alpha_{L_A}}
\]  

where \( t_{\alpha_{L_A}/2} \) is implicitly defined by

\[
\alpha(t_{\alpha_{L_A}/2}) = \frac{\alpha_{L_A}}{2}.
\]  

\( r_\alpha \) can also be interpreted as the ratio of advective and dispersive scales like in the definition of the Peclet number.

(3) **Mixing** is required for properly reproducing fast reactions (slow reactions should be properly reproduced if the resident time distribution is honored, which is assured if mean advection and dispersion are reproduced). As discussed above, mixing is essentially dispersive and well characterized by \( M_D \) (equation (14)) for late times. Therefore, assuming dispersion to be well reproduced, an effective transport formulation only needs to reproduce the deviation from dispersive mixing, characterized by \( \gamma \) (equation (12)). In a first stage, the comparison can be restricted to the amplitude of the deviation \( \gamma_{max} \) and the time range over...
which it extends $r_{xy}$. In a more advanced stage, the characteristic shape of the $\gamma$ function, $f$, can be used for comparison.

To compare the timings of spreading and mixing, we define the additional criterion $r_{MT}$ as the ratio of the characteristic spreading time $t_{\alpha_{1/2}}$ to the characteristic mixing time $t_{\gamma_{\text{max}}}$

$$r_{MT} = \frac{t_{\gamma_{\text{max}}}}{t_{\alpha_{1/2}}}.$$  (19)

$r_{MT}$ compares the timing of the development of the resistance to mixing and of spreading and rates the lag between the timing of mixing and spreading.

(4) **Flexibility.** Most of the work on effective transport is of a theoretical nature, but the ultimate goal should be application to real problems. This implies that a valid transport formulation should be able to accommodate different types of boundary conditions and flow regimes (i.e., transient flow) and dimensions. Most importantly, it should accommodate characterization. Dispersion usually includes the effects of heterogeneity and uncertainty. Whereas the latter is reduced by aquifer characterization, the former is not. Specifically, hydrologists use geology, hydraulics, geophysics, hydrochemistry and isotopes to figure out, among other things, the patterns of spatial variability of hydraulic conductivity. The resulting models display variability not only in the mean log-$K$ but also on their correlation distance and variance. An effective transport formulation should be able to honor this variability.

5 **Results and discussion**

We consider well established that MRMT, and other non-local in time formulations, can reproduce mean advection and spreading, as discussed in the introduction. Mean advection in the MRMT approach is equivalent to that of the HPM provided that flux and total porosity are equivalent. And the distribution of residence times in immobile zones can be adapted so that the asymptotic dispersivity of the MRMT model be equal to that of the HPM model in equation (11). It is always possible as dispersivity is an increasing function of the residence times. This imposes a condition on the temporal range of $t_1,...,t_N$ or equivalently on their mean residence time $\langle \tau_{\text{MRMT}} \rangle$. As trapping in the immobile zones is the main dispersive mechanism, the mean residence time is logically adapted to calibrate the asymptotic dispersivity. With the total flow imposed to be set by the HPM, the characteristic spatial scale is the typical plume position at $\langle \tau_{\text{MRMT}} \rangle$. As the characteristic spatial and temporal scales are interrelated to ensure
consistent asymptotic behaviors, comparison of results can be performed on dimensionless terms and should ensure consistent preasymptotic regimes. In fact, MRMT are calibrated on tracer tests and breakthrough information, but this does not ensure a good reproduction of mixing [Luo and Cirpka, 2011]. Therefore, we restrict our comparison to mixing criteria and sensitivity to initial conditions.

5.1 Comparison of mixing in HPM and MRMT

In Heterogeneous Porous Media (HPM), the temporal extension of the deviation from the dispersive mixing regime \( r_T \) does not depend significantly on the permeability heterogeneity, as also expressed by the constancy of the shape of \( \gamma \) (Figure 2). We thus compare the shape of \( \gamma \) obtained for the HPM with \( \sigma_Y^2=9 \) (function of equation (16)) to shapes of \( \gamma \) obtained for various MRMT models obtained under consistent injection conditions (equations (6) and (7)).

For MRMT, extreme values have been investigated to get the possible range of behaviors. For slopes \( m \), we adopted the range observed in nature as discussed in section 2.1 with \( m \) varying between \( m=1.5 \) (typical fracture/matrix case) and \( m=2.5 \). The Single Rate Mass Transfer (SRMT) is also shown for comparison. The porosity ratio \( \beta \) does not have an upper bound. In fact, ideally, the mobile porosity could be zero. We adopted \( \beta=150 \) as a large upper value. Larger upper values would not affect results and might cause numerical difficulties. The same can be said for \( t_N/t_1 \) for which we took \( t_N/t_1=10^3 \) as the upper bound. The analysis presented hereafter has been made for different combination of parameters within these bounds. As all models lead to consistent conclusions, we only present the most characteristic results.

All MRMT models capture the sharp rise of \( \gamma \) at times smaller than \( t_{\gamma_{\text{max}}} \) (Figure 3). The sharp rise comes from a strong initial divergence from the equivalent Gaussian concentration profile. Initial behavior is dominated by the contrast of the quickly progressing concentrations in the mobile zones and trailing concentrations in the low flow or immobile zones. On one hand, dispersion induces a sharp decrease of \( M_D \), which is inversely proportional to \( \sigma_L \) (equation (14)). On the other hand, trapping maintains high concentrations in the immobile zone and high values for the second moment of the concentration distribution \( M \). Divergence of \( M \) from \( M_D \) increases until it reaches its maximum at \( t_{\gamma_{\text{max}}} \).

At larger times, progressive release of solute mass from the immobile zone and equilibration with the concentration values in the mobile zone let \( f \) decrease. The insert of Figure 3 highlights in a lin-lin graph the differences of the decrease stage. The MRMT model that best
matches the scaling function $f$ is obtained for $m=2.25$. For power law slopes $m$ larger than 2, the decrease of $f$ is qualitatively similar to that of the single-rate mass transfer model. For $m$ values closer to 2, the deviation is more sustained but displays the same decreasing trend. The behavior changes significantly when $m$ becomes smaller than 2, with a very slow decrease of $f$ coming directly from the effect of the slow drainage of the immobile zones having the smallest rates (long exchange times). At least in MRMT models with $m$ lower than 2, trapping displays a much longer memory effect in MRMT than in HPM.

While similar to HPM for the extension of the non-dispersive mixing regime, MRMT models with slopes $m$ larger than 2 converge to less anomalous Single Rate Mass Transfer. In terms of mixing, this translates in small amplitudes in $\gamma$ (small $\gamma_{\text{max}}$ values). For $m=2.25$ and $\Delta L_0/\alpha_L = 0.075$, we have computed $\gamma_{\text{max}}$ for a large variety of $t_N/t_1$ and $\beta$ values, the only two remaining parameters of the MRMT model. $\gamma_{\text{max}}$ first increases with $t_N/t_1$ and $\beta$ and quickly saturates to a maximal value of 2.85 (Table 1). $\gamma_{\text{max}}$ varies between 0.57 and 2.85 by a maximum factor of 5. In the HPM case, however, $\gamma_{\text{max}}$ is always larger than 3, reaches values of 15 for a Peclet number $Pe$ of 100, scales like the square root of $Pe$ and is thus not limited in amplitude.

MRMT models cannot match both the amplitude and the timing of $\gamma$. For MRMT models with $m$-slopes larger than 2, mixing is far more dispersive in MRMT than in HPM (smaller deviation values $\gamma$ in the MRMT models). MRMT models with $m$-slopes smaller than 2 induce larger but much too sustainable deviations. As a result, MRMT models have a stronger memory of trapping or display less non dispersive-mixing, without excluding to display both differences simultaneously. The difficulties of MRMT models to capture mixing might be linked to the existence of the structure of concentrations in lamellas where stretching and folding extends the concentration front and enhance the eventual mixing by diffusion (Figure 1) [de Anna et al., 2014b; Le Borgne et al., 2015].

Concerning the non-dispersive mixing shapes of the scaled function $f$, MRMT models display a much broader range of shapes than HPM. The invariant shape property of HPM is not recovered in MRMT. On the contrary, MRMT shapes depend strongly on the distribution of transfer rates. This is an advantage to match a wider range of cases issuing possibly different $\gamma$ functions. But it is a drawback to fit just one case as it restricts the MRMT models that can match HPM simulations with broad ranges of $Pe$ and $\sigma Y^2$ values.
### 5.2 Influence of initial injection size

To qualify the memory effect in MRMT and HPM, we analyze their sensitivity to the initial injection width $\Delta L_0$. Spreading as defined by the characteristic longitudinal plume extension (equations (8), (9) and (10)) does not depend on the initial concentration $c_0$ (equation (6)). $\sigma_L$ is initially of the order of $\Delta L_0$ and quickly becomes larger. Spreading loses quickly any memory of the initial conditions. We note that this is the case because sampling effects do not intervene as the transverse injection scale is assumed large enough to ensure by itself ergodic sampling. Like spreading, the second moment of the reference Gaussian concentration $M_D(t)$ does not depend either on the initial concentration but only on the injected mass divided by the characteristic area occupied by the plume ($\Delta T_0 \sigma_L$) (equation (14)).

The concentration second moment $M(t)$ (equation (13)) however depends critically on the injection width through the relation between injected mass and concentration (equation (6)). At initial times, the concentration second moment is proportional to the injected concentration value. At late times, the concentration second moment has lost the memory of the initial concentration and is only function of the injected mass $m_0$. As a result, we expect that the concentration second moment and the deviation towards the dispersive mixing regime $\gamma$ depends on the injection conditions, here represented by the injection width $\Delta L_0$. On the basis of numerical simulations, we compare the evolution of $\gamma$ with $\Delta L_0$ for HPM and MRMT models.

Results of the $\gamma$ function for both MRMT and HPM models are displayed on Figure 4a and Figure 4b, respectively, for different injection sizes. We have performed simulations for comparable ranges of $\Delta L_0/\alpha_{LA}$ values (0.005-0.1 for HPM and 0.02-0.4 for MRMT). We have checked numerically that the results displayed for the two specific displayed MRMT and HPM cases display generic tendencies. In both models, injection width has a critical influence on $\gamma$ (Figure 4). Smaller injection windows let the initial concentration increase and enhance the deviation towards the dispersive mixing regime. We use the maximum deviation $\gamma_{\text{max}}$ to characterize the overall influence of $\Delta L_0$. In the MRMT and HPM models, maximum deviations $\gamma_{\text{max}}$ have different scaling (Figure 4a and b, inserts):

$$\gamma_{\text{max}} \sim (\Delta L_0)^{-1} \text{ for MRMT}$$

$$\gamma_{\text{max}} \sim (\Delta L_0)^{0.5} \text{ for HPM}$$

(20)
For MRMT, $\gamma_{\text{max}}$ evolves like the initial concentration level (equations (6) and (7)). For HPM, $\gamma_{\text{max}}$ has the same scaling for the initial conditions $\gamma_{\text{max}} \sim 1/\sqrt{\Delta L_0}$ and for the diffusion coefficient as $\gamma_{\text{max}} \sim \sqrt{Pe}$ [de Direuzy et al., 2012]. Doubling the injection width has a comparable effect to doubling the diffusion coefficient. Initial dilution over $\Delta L_0$ and dilution induced by diffusive/dispersive processes reduce the overall deviation to the dispersive mixing regime $\gamma$ in the same proportion. The reduction of concentration in HPM comes from the diffusive/dispersive processes while, in MRMT, it comes from the progressive release of solutes with high concentrations close to $c_0$ trapped in the immobile zone. Because of their differing signatures, both processes cannot be compared and the dispersive/diffusive processes of HPM cannot be modeled as trapping/release mechanisms.

6 Conclusion

We propose conditions to test anomalous transport frameworks not only on spreading, but also on mixing. We define a minimum set of 6 essential constrains that they should respect in order to retain the main transport, reactivity and reactive transport couplings. These constrains involve the conservation of (1) the mean advection, (2) dispersivity amplitude and (3) timing generally imposed. Beyond these flow and spreading metrics, (4) amplitude and (5) timing of the deviation towards the dispersive mixing regime should be respected. The last condition concerns (6) the respective timings for mixing and spreading. Under ergodic injection conditions, spreading is characterized by the standard dispersivity describing the evolution of the plume size along the main flow direction. Mixing is characterized by the deviation from the dispersive mixing regime $\gamma$ defined as the second moment of the concentration distribution of a conservative tracer divided by the one of a Gaussian concentration pattern with the same spread minus 1 (equation (12)). Zero initially and asymptotically, $\gamma$ traduces the macroscopic effect on mixing of the concentration structures within the solute plume.

We use these criteria to evaluate Multi-Rate Mass Transfer models by comparison to advective-diffusive transport simulations through Heterogeneous Porous Media (HPM) represented by the classical isotropic 2D Gaussian correlated multi-Gaussian log-permeability fields characterized by variances between 1 and 9. A broad range of Multi-Rate Mass Transfer models (MRMT) are considered. We conclude that MRMT models cannot match both the amplitude and the timing of $\gamma$. MRMT models can reproduce observed spreading rates and some non-dispersive mixing. But they tend to induce larger and too sustained deviations from...
dispersive mixing. As a result, MRMT models display a longer memory but less non
dispersive-mixing than HPM. We attribute this divergence to the fact that MRMT represent
non-dispersive mixing through trapping mechanisms, whereas it is controlled by stretching
and folding in HPM. Divergent sensitivities to initial conditions confirm that dispersive-
diffusive induced mixing in HPM cannot be modeled by mobile/immobile models.

Our study does not preclude however the existence of effective transport equations consistent
with spreading and mixing of HPM. But we argue that the proposed criteria and existing
results of HPM should be used as guidelines to set up effective transport equations that
respect spreading, mixing, and eventually reactive transport.

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Table 1: Values of the maximum deviation to the dispersive mixing regime $\gamma_{\text{max}}$ for MRMT models with a power-law exponent of the rate distribution $m=2.25$ and $\Delta L_0 / \alpha_{LA} = 0.075$.

<table>
<thead>
<tr>
<th>$t_N/t_1$=100</th>
<th>$\beta$=100</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta$</td>
<td>$\gamma_{\text{max}}$</td>
</tr>
<tr>
<td>1</td>
<td>0.57</td>
</tr>
<tr>
<td>10</td>
<td>1.96</td>
</tr>
<tr>
<td>100</td>
<td>2.84</td>
</tr>
<tr>
<td>300</td>
<td>2.85</td>
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Figure 1: Concentration fields normalized by their maximal value $c(x,t)/\max(c(x,t))$ and their related Gaussian profile concentrations $c_D(x,t)/\max(c(x,t))$ in the bar over them at the four evolving times indicated on Figure 2. In this case, the time at which the non-dispersive mixing reaches its maximum $t_{\gamma \text{max}}$ is of the same order of the advection time.
Figure 2: Time evolution of the deviation from dispersive mixing $\gamma(t)$ defined by equation (12) in HPM for evolving log-K variances, $\sigma_Y^2$, under a small width injection window ($\Delta L_v/\alpha_{LA} = 0.075$), flux weighted injection conditions and $Pe=100$ (adapted from de Dreuzy et al. [2012]). The similarity of function shapes is highlighted in the insert by the scaling function $f$ of equation (16) where the thick black line is the average of the displayed functions. Note that the time of maximum deviation, $t_{\gamma_{max}}$, is hardly affected by $\sigma_Y^2$ and falls around the characteristic advection time $\lambda/v$. The four dashed lines indicate the times displayed in Figure 1.
Figure 3: Comparison of $\gamma$ shapes ($f$ scaling functions defined in (16)) for HPM and MRMT simulations with slopes of the power-law distributions of MRMT transfer rates, $m$, between 1.575 and 2.5. HPM is represented by the broad black curve obtained from the insert of Figure 2. The insert displays the same curves in arithmetic scale.
Figure 4: Dependence of the deviation from dispersive function $\gamma$ on the injection width $\Delta L_0/\alpha_{L_A}$ for (a) MRMT and (b) HPM models. Insert shows the dependence of $\gamma_{\text{max}}$ on $\Delta L_0/\alpha_{L_A}$. 