## Modelling and Upscaling of Transport in Carbonates During Dissolution: Validation and Calibration with NMR Experiments

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### Abstract

We present an experimental and numerical study of transport in carbonates during dissolution and its upscaling from the pore (~  $\mu$ m) to core (~ cm) scale. For the experimental part, we use nuclear magnetic resonance (NMR) to probe molecular displacements (propagators) of an aqueous hydrochloric acid (HCl) solution through a Ketton limestone core. A series of propagator profiles are obtained at a large number of spatial points along the core at multiple time-steps during dissolution. For the numerical part, first, the transport model—a particle-tracking method based on Continuous Time Random Walks (CTRW) by [1]—is validated at the pore scale by matching to the NMR-measured propagators in a beadpack, Bentheimer sandstone, and Portland carbonate [2]. It was found that the emerging distribution of particle transit times in these samples can be approximated satisfactorily using the power law function  $\psi(t) \sim t^{-1-\beta}$ , where  $0 < \beta < 2$ . Next, the evolution of the propagators during reaction is modelled: at the pore scale, the experimental data is used to calibrate the CTRW parameters; then the

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shape of the propagators is predicted at later observation times. Finally, a numerical upscaling technique is employed to obtain CTRW parameters for the core. From the NMR-measured propagators, an increasing frequency of displacements in stagnant regions was apparent as the reaction progressed. The present model predicts that non-Fickian behaviour exhibited at the pore scale persists on the centimetre scale.

### Nomenclature

### Acronyms

CTRV	V Continuous time random walk			
PFG	Pulsed field gradient			
PTM	Particle-tracking method			
TPL	Truncated-power law			
Greel	Greek Symbols			
$<\zeta>$	$_0$ Mean particle displacement	m		
β	Power-law coefficient			
Δ	NMR observation time	S		
$\phi$	Porosity			
$\psi(t)$	Transit-time distribution			
τ	Normalized time $t/t_1$			
$\zeta$	Particle displacement	m		
Roman Symbols				
A	Normalization constant			
d	Core diameter	m		
$D_m$	Diffusion coefficient	$\mathrm{m}^2~\mathrm{s}^{-1}$		

Damköhler number		
Core length	m	
Probability density function		
Probability of a particle moving from $i$ to $j$		
Péclet number		
Flux of fluid	$\mathrm{m}^3~\mathrm{s}^{-1}$	
Transit time	S	
Average advection time	S	
Diffusion cut-off time	S	
Experimental time	S	
Interstitial velocity	${\rm m~s^{-1}}$	
Subscripts		
Core scale		
	Damköhler number Core length Probability density function Probability of a particle moving from <i>i</i> to <i>j</i> Péclet number Flux of fluid Transit time Average advection time Diffusion cut-off time Experimental time Interstitial velocity <b>ripts</b> Core scale	

- CP Core-plug scale
- i, j Node indices
- k Link indices
- P Pore scale

### 1 1. Introduction

Transport and reaction of fluids in porous media is important in many hydrogeological problems. Examples include stimulation in petroleum reservoirs by acidization [3], water and contaminant management [4], and geological storage of carbon dioxide [5, 6]. Rock matrix dissolution refers to reactions at fluid/solid boundaries that result in the dissolution of the solid grains, pore growth, and variation of flow characteristics. For practical applications, the main difficulties in building models with predictive capabilities

are twofold: first, reaction changes the microstructure of the rock, and thus 9 the structure heterogeneity starting at the pore scale. Second, there is a large 10 disparity between the scale at which transport can be understood from first 11 principles, and the scale at which practical predictions are needed [7]. Since 12 in many cases, the formal closure problem may be too complex for general 13 solution, we propose to study the effects of reaction on solute transport from 14 micrometre to centimetre scales using a heuristic multiscale modelling ap-15 proach which does not impose a particular form to the governing equations, 16 in conjuction with NMR fluid propagator method to validate and calibrate 17 the model at the pore scale sequentially during dissolution.



Figure 1: A reactive transport experiment where HCl solution is injected through a core of Ketton limestone core. Flow channels are formed due to solid dissolution. The figures depict (a) the dimension of the core, and (b) the difference in porosity between the beginning and the end of the experiment provided using NMR imaging, with blue being the smallest and green the largest; white indicates no change in porosity.

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The heterogeneity of porous media in geological formations is embodied 19 by the pore structure as well as the mineralogical heterogeneity resulting 20 from multiple components. In heterogeneous porous media, the observed 21 reactive transport processes frequently do not behave according to the trans-22 port laws that can be derived for homogeneous media, see [8, 9, 10, 11, 12, 13], 23 such as the classical advection-diffusion-reaction equations. Because reactive 24 transport modeling is typically applied at large scales, it necessarily ignores 25 spatial heterogeneities at scales smaller than the size of model discretization 26 , see [14, 15]. Several techniques have been introduced as a remedy, i.e to 27 compute effective parameters which capture subscale effects, see [16, 17]. 28 Furthermore, while under limited circumstances the homogeneity assumption 29 is reasonable, the pore-scale heterogeneities can result in a significant scaling 30

<sup>31</sup> effect because of the spatial variations of concentrations and reaction rates,

<sup>32</sup> leading to the breakdown of the homogeneity assumption. This scaling effect

may be one of the causes of the order-of-magnitude differences between laboratory measured reaction rates and that obtained from field measurements,
see [18, 15]. Therefore, it is important to understand the effects of pore-scale
heterogeneities on the reactive transport processes.

Conventionally transport simulation is performed by solving the advection-37 dispersion equation (ADE) with known, albeit complicated, boundary con-38 dition. In some cases, depending on the investigated conditions and on the 39 quality of the characterization of the heterogeneity of the system, the ADE 40 can still be used effectively, see [19]. Furthermore, it is difficult to deter-41 mine the correct values of the coefficients in the model. As the solution 42 of the ADE at a fine scale over the full extent of the geological hetero-43 geneity is prohibitively difficult, as we have no general way to incorporate 44 uncertainty in the description of the reservoir model for the prediction of 45 transport. Motivated by this problem, Rhodes et al in [1, 20] presented a 46 particle-tracking method based on CTRW (from here-on called PTM-CTRW) 47 for solving single-phase transport across a hierarchy of length-scales. Unlike 48 other upscaling methods which rely on special basis functions, or homogeni-49 sation to capture the subscale effects (see [21, 22, 23, 16]), the method does 50 not pre-suppose the functional form of the upscaled transport equations, and 51 automatically accounts for uncertainty in the field-scale description. PTM-52 CTRW has been tested for simulating transport in sandstones. Here, PTM-53 CTRW forms the basis of our solute transport simulations and its application 54 is extended to reactive transport in carbonates. 55

To rid geological transport simulation of uncertainties due to upscaling 56 it is imperative that a numerical model undergoes rigorous laboratory vali-57 dations. In our study, the model and its validation are built upon pore-scale 58 information. The distribution of molecular displacement (or propagators) in 59 the preasymptotic dispersion regime can provide the basis for validation of 60 transport models that are based on X-ray microtomography images of the 61 pore space—see [24, 25]; and [26]. In recent years, Nuclear Magnetic Res-62 onance (NMR) has been used to probe transport signatures in porous glass 63 beads, see [27]. It has also been used in beadpack, sandstone and carbonate 64 samples in the preasymptotic dispersion regimes e.g. in [2], and [28]. This 65 paper augments previous work and describes how NMR 1D-imaging and fluid 66 propagator measurements are employed to provide experimental insights of 67 hydrochloric acid (HCl) flow through Ketton carbonate cores at multiple 68 time increments during dissolution. First HCl solution is injected into the 69 core as illustrated in figure 1. The change in porosity, and propagators at a 70

<sup>71</sup> large number of spatial points along the core can be monitored throughout <sup>72</sup> the experiment. These propagators are then used to calibrate our model at <sup>73</sup> a pore ( $\sim \mu m$ ), and core-plug scale ( $\sim mm$ ), as well as to derive the local <sup>74</sup> probability density functions (PDFs) of transit times, the combination of <sup>75</sup> which, will be used to derive the PDF at the core scale ( $\sim cm$ ).

The scope of this work can be summarized as follows. First, PTM-CTRW 76 is employed to reproduce the NMR-measured propagators through a bead-77 pack, Bentheimer sandstone, and Portland carbonate cores and thereby val-78 idate the described model. Second, the pulsed field gradient NMR technique 79 is used to find a series of reactive propagators in preasymptotic flow through 80 Ketton carbonate core at multiple times during dissolution. Third, these 81 propagators are reproduced numerically, an array of time-transit distribu-82 tions is obtained, and thereby calibrate the present model at the pore scale. 83 Finally, these propagators are used as the bases of our core-scale simulation 84 and derive the upscaled CTRW parameters at the beginning and the end of 85 the experiments. This model can then be used to predict transport at any 86 scale of interest. 87

### <sup>88</sup> 2. Continuous Time Random Walks

The description of CTRW here is by no means exhaustive; for details on the application of CTRW in a geological context, the reader is referred to an excellent review by Berkowitz et al [29].

Anomalous or non-Fickian transport is prevalent in heterogeneous porous 92 media, and is ubiquitous in the context of tracer migration in geological 93 formations. Anomalous transport can be described elegantly as a continuous 94 time random walk. In a CTRW framework, dispersion, which results in 95 solute spreading at the scale of observation, is accounted for by a transit 96 time distribution function  $\psi(t)$ . For many systems,  $\psi(t)$  exhibits power-law 97 dependencies:  $\psi(t) \sim t^{-1-\beta}$ , where  $\beta < 2$  is an exponent. For such systems, 98 this leads to the scaling of outlet concentration  $C(t) \sim t^{-1-\beta}$ , see [30]. 99

### 100 3. Transport Model Description

Traditionally CTRW has been applied to find the ensemble average behaviour of a plume in a macroscopically homogenous domain, see [29, 31]. CTRW has been applied to heterogeneous media, but for relatively coarsely gridded two-dimensional systems where the solution involves the numerical

inversion of a multi-dimensional Laplace transform, see [32]. Rhodes et al 105 in [1, 20] developed a simpler approach, PTM-CTRW, to describe trans-106 port spanning across microns to kilometre scales. The stochastic framework 107 also allows more complicated boundary conditions and various types of dis-108 tribution function to be used. Here the modelling framework according to 109 PTM-CTRW is explained, in which transport is seen as a series of random 110 hops from one node in a 3D lattice to its neighbouring node. Particles move 111 between a series of discrete nodes or sites with a probability  $\psi(t:i,j)$  that 112 a particle that first arrives at site i will move to site j in a time t + dt. 113

At the heart of PTM-CTRW is the *correct* choice of transit-time distri-114 bution  $\psi(t)$ . In their pore-scale simulation, Rhodes et al [1] employed two 115 types of transit-time distribution, one derived from the advection-diffusion 116 equation, as presented in [33], and another a truncated power-law function 117 as an ensemble averaged transit-time distribution, presented in [31]. They 118 conducted numerical studies comparing the two functions: the former was 119 employed in a 3D lattice with a Berea sandstone derived distribution of 120 throat radii, while the latter was implemented in an effective homogenous 121 lattice. They compared the results from both methods with experimental 122 data for Berea sandstone and found that the truncated power-law function 123 gave the observed transport behaviour and reproduced the dispersion co-124 efficients obtained from experiments accurately. The truncated power-law 125 (TPL) transit-time distribution function, as presented in [31], is 126

$$\psi(t) = Ae^{-t/t_2}(1 + t/t_1)^{-1-\beta},\tag{1}$$

where A is a normalization constant such that  $\int_0^t \psi(t')dt' = 1$ , and  $\beta \leq 2$  is a power-law coefficient.

Using network modelling of transport Bijelijc and Blunt in [34] were able 129 to match the transit-time probability density function measured in links be-130 tween neighbouring pores of a Berea sandstone pore network using equation 131 (1) with  $\beta = 1.8$ . Furthermore, Bijeljic et al [35] performed direct simulations 132 of transport in the pore spaces of micro-CT images of Berea sandstone and 133 Portland carbonate and obtained  $\beta = 1.8$  and  $\beta = 0.7$  respectively. Transit 134 times were now measured as the time particles to migrate from one pore 135 voxel to another. 136

At the Darcy scale an explicit relationship between the histogram of permeability and  $\beta$  has been demonstrated [36]. Here a truncated power-law is also used to describe small-scale transport, where the exponent  $\beta$  exponent



Figure 2: The pore-to-core simulation technique. Transport is modelled as a series of hops between nodes via links with a known transit time distribution  $\psi(t)$ . At the smallest scales, advective and diffusive transport is simulated through a lattice representing the porous medium of interest. Transport from one pore to another is described by  $\psi_P$  that is averaged over all possible statistical realizations of the structure. This  $\psi_P(t)$  is then input into a simulation at the core-plug scale to compute  $\psi_{CP}(t)$  for transitions of particles over the mm scale. Finally transport at a core scale can be represented as a single hop governed by the transit-time distribution function  $\psi_C(t)$ . This figure is adapted from [1].

acts as a measure of heterogeneity. In figure 2 the pore-to-core transport
isimulation framework is described.

To clarify the implementation of our method, in figure 3 we show the behaviour of  $\psi$  given the variety of its parameters. We plot equation 1 where  $\psi$ , is a function of the normalized time  $\tau = t/t_1$  for several Péclet numbers  $Pe = 2t_2/t_1$ . The left figure shows  $\psi$  for  $\beta = 0.6$ , and the right figure  $\beta = 1.8$ . At larger  $\beta$ , the long-time distribution diminish faster as illustrated by the power-law trend  $\psi \sim \tau^{-1-\beta}$ .

At the pore scale (~  $\mu$ m), a transit time distribution function  $\psi_P(t)$  is derived from either a semi-analytic description in an idealized network, or



Figure 3:  $\psi$  as a function of  $\tau = t/t_1$ . (left)  $\beta = 0.6$ , and (right)  $\beta = 1.8$ , for several *Pe* numbers using equation 1.

from direct simulation.  $\psi_P(t)$  will form the basis of simulation at the core-150 plug scale ( $\sim$  mm). Numerical upscaling will be implemented such that 151 transport at this scale can be modelled as a single hop governed by a transit-152 time distribution  $\psi_{CP}(t)$ . For the core-scale simulation (~ cm), a lattice is 153 used that is similar in shape to the rock core used in the experiments—a 154 cylinder—in which  $\psi_{CP}(t)$  is applied in each link, this core-scale lattice will 155 be calibrated a priori. Numerical upscaling will then be used again to obtain 156  $\psi_C(t)$  (see Section 6). 157

In the pore-scale simulation, transport is simulated on a homogeneous 3D 158 lattice consisted of nodes and links. Within each link, transport is governed 159 by the transit-time distribution function  $\psi_P(t)$ , equation (1). First a pressure 160 difference is assigned at the inlet and outlet faces. Then the pressure field is 161 solved by enforcing mass balance at each node, assuming slow, single-phase, 162 Newtonian flow. At each node the mass-flux (q) conservation  $\sum_k q_k = 0$  is 163 applied for each node connected to links k by which the velocity field at each 164 link can be known, see Appendix A for details. Assuming complete mixing at 165 each node, the probability p(i, j) that a particle landing at pore i will move 166 to one of its neighbours is calculated 167

$$p(i,j) = \frac{Gq_{ij}}{1 - e^{-\operatorname{Pe}_{ij}}}; \text{ if } q_{ij} > 0$$

$$p(i,j) = \frac{Gq_{ji}}{e^{\operatorname{Pe}_{ij}} - 1}; \text{ if } q_{ij} < 0$$
(2)

where  $q_{ij}$  is the flux in a link connecting node *i* and *j*, and *G* is a normalization

<sup>169</sup> coefficient such that  $\sum_{j} p(i,j) = 1$ , i.e,

$$\frac{1}{G} = \sum_{\forall q_{ij} > 0} \frac{q_{ij}}{1 - e^{-\operatorname{Pe}_{ij}}} + \sum_{\forall q_{ij} > 0} \frac{q_{ji}}{e^{\operatorname{Pe}_{ij}} - 1}.$$
(3)

Then a number of particles are released either at the inlet face, or randomlyin the lattice.

At an intersection, a random number a is generated. P(i, j) is then read from memory, and defined as  $P(i, j) = \sum_{m} p(i, m)$ ;  $m \leq j$ . The process is iterated such that

$$P(i, j-1) \le a \le P(i, j). \tag{4}$$

When (4) is satisfied, the particle will move along the link i-j. A random number z is generated and the time t required to move along the link i-jis found by solving, using a root-finding method, F(t) = z i.e,

$$F(t) = \int_0^t \psi_P(t) = z \tag{5}$$

178 where

$$\psi_P(t) = A e^{-t/t_2} (1 + t/t_1)^{-1-\beta} \tag{6}$$

and  $t_1 = l/v$ , l is the link length and v is the fluid velocity within that link.  $t_2 = l^2/D_m$  is the cut-off diffusion time, and  $D_m$  is the self-diffusion coefficient of the working fluid. v and  $D_m$ , thus  $t_1$  and  $t_2$  are known *a priori*; leaving the adjustable parameter  $\beta$  that describes transport heterogeneity.

To obtain the transit-time distribution at the next larger scale, the same 183 technique as in [1] is used, i.e. a number of particles at t = 0 is released 184 at the inlet face of a 3D lattice and the time required for each particle to 185 transit recorded.  $\psi(t)$  can be obtained at the next larger scale by matching 186 the emergent distribution of the transit times of each particle to equation 187 (1). This is illustrated in figure 4 where  $\psi_s(t)$  is the transit time distribution 188 function at a scale larger than where transport is governed by  $\psi_r(t)$ . This 189 methodology is applied to obtain both  $\psi_{CP}(t)$  from  $\psi_P(t)$ , and  $\psi_C(t)$  from 190  $\psi_{CP}(t).$ 191

### <sup>192</sup> 4. Experimental Technique, Apparatus and Results

In this paper, pulsed field gradient nuclear magnetic resonance (PFG-NMR) is used to obtain propagator measurements, i.e. probability distributions,  $P(\zeta)$ , of molecular displacement for a given observation time ( $\Delta$ )



Figure 4: The upscaling methodology. First, flow in each link is solved such that  $t_1$  and  $t_2$  can be determined. Then a number of particles are launched at the inlet face at t = 0. The time required for one particle to reach the outlet face is that particle's transit time.  $\psi_r^k(t)$  is the transit-time distribution function governing transport in link k.  $\psi_s(t)$  is obtained by matching the emergent transit-time distribution with equation (1).

as described in [37]. Here, the experiments are applied such that the water, 196 resident in the rock core, is studied. By observing the displacement,  $\zeta$ , of 197 water molecules over a range of observation times  $\Delta$  and flow velocities v, 198 the fluid behaviour and pore-scale heterogeneity can be characterised. These 199 experiments are time-consuming when the data is fully sampled, requiring 200 experimental durations of the order of hours to complete which is impracti-201 cal for the study of the dynamic, reacting system being investigated here. In 202 this work we have reduced the data acquisition time through undersampling 203 and reconstruction of the smooth acquisition domain signal. Further details 204 of the sampling and reconstruction parameters used here can be found are 205 described in [38]. 206

For the reaction, 10 litres of a 0.01 M HCl solution was flowed at Q =207  $8.3 \times 10^{-7}$  m<sup>3</sup> s<sup>-1</sup>, through a 7.2 cm long by 3.81 cm diameter sample of 208 Ketton limestone. Propagators are recorded in 0.88 cm slices along the core 209 with an observation time ( $\Delta$ ) of 0.25 s. In each slice, the porosity  $\phi$  and 210 propagator are measured throughout the dissolution process with porosity 211 profile measurements being acquired in 0.5 minutes and the propagator mea-212 surements being acquired in 14.5 minutes using the undersampling methods 213 described in [38]. 214

The diffusion coefficient  $D_m$  of water, in water, at 293 K is  $2.1 \times 10^{-9}$ m<sup>2</sup> s<sup>-1</sup>. The initial porosity of the core  $\phi$  is 0.24. The interstitial velocity v is  $\frac{(Q/A)}{\phi} = 3.06 \times 10^{-3}$  m s<sup>-1</sup>. The characteristic length l of the Ketton limestone can be estimated i.e,  $l = \pi/S$ , where  $S[m^{-1}]$  is the specific surface



Figure 5: Experimental results showing the NMR propagator contour at (a)  $t_{\exp} = 0$  s the beginning of the experiment; and at (b)  $t_{\exp} = 11800$  s—the end of the experiment. The average flow velocity v is  $3.06 \times 10^{-3}$  m s<sup>-1</sup>, and the observation time  $\Delta$  is 0.25 s. The propagator  $P(\zeta)$  consists of the normalized probability of displacement  $\zeta$  such that  $\int_{\zeta} P(\zeta) d\zeta = 1$ .

area, such that  $l = 4.07 \times 10^{-4}$  m, as presented in [39]. The corresponding Péclet number,  $Pe = lv/D_m$ , is therefore 593. The Damköhler number, the ratio of acid consumed and the acid transported by convection, is defined in [40] as,

$$Da = \frac{\pi r}{vn} \tag{7}$$

where r is the reaction rate constant of pure calcite in 0.01 M HCl solution at 293 K ( $1.5 \times 10^{-3} \text{ mol m}^{-2} \text{ s}^{-1}$ ) measured experimentally in [41]. n is calculated using  $n = \rho_{calcite}[1 - \phi]/M_{calcite}$ .  $\rho_{calcite}$  is the density of pure calcite ( $2.71 \times 10^3 \text{ kg m}^{-3}$ ), and  $M_{calcite}$  is the molecular mass of calcite ( $0.1 \text{ kg mol}^{-1}$ ). In our experiment,  $Da = 7.7 \times 10^{-5}$ .

Figure 5 shows the propagators as a function of axial position along the 228 core-plug, before and after dissolution. Before reaction the propagators are 229 uniform along the length of the core, showing a sharp stagnant region centred 230 on 0 displacement and a broad flowing region extending to a displacement of 231  $\sim 3.5$  mm. After dissolution of the solid matrix has taken place, predomi-232 nantly in the first half of the core, the propagators in this region evolve—fast 233 moving fluid slows as the pore-space is opened up and the overall porosity is 234 increased. 235

### 236 5. Model Validation

# 237 5.1. Comparison with NMR-measured Propagators in a Beadpack, Bentheimer 238 sandstone, and Portland Carbonate

The results of the numerical methods are now compared with the NMRmeasured propagators, without reaction, transport only [2] in a beadpack, Bentheimer sandstone, and Portland carbonate.

The computational domain is a homogenous  $0.008 \times 0.008 \times 0.008$  m<sup>3</sup> 242 lattice consisting of  $80 \times 80 \times 80$  links. In this analysis, the system is homo-243 geneous at the core scale and the measurements of displacement are taken 244 across the whole core. In these computations, the same interstitial velocities 245 are used as in the experiments, namely  $v = 9.1 \times 10^{-4}, 1.03 \times 10^{-3}, 1.26 \times 10^{-3}$ 246 m  $s^{-1}$  for beadpack, Bentheimer sandstone, and Portland carbonate respec-247 tively. Particles are launched at random locations in the lattice at t = 0 s 248 and their movement is tracked. If a particle exits the inlet or outlet, it is 249 randomly reassigned to the opposite face using a flux-weighted assignment. 250

The propagators were measured at different  $\Delta$ . The propagators were 251 computed using the transit-time distribution, equation (1). Average ad-252 vection times  $t_1 = 0.11, 0.097$ , and 0.079 s are known from the interstitial 253 velocities v and the cut-off diffusion time is  $t_2 = l^2/D_m = 15$  s. By fitting 254 the power-law exponent  $\beta$ s it was possible to match the experimental data: 255  $\beta = 1.96, 1.76, \text{ and } 0.63 \text{ yield propagator profiles with those of a beadpack}$ . 256 Bentheimer sandstone, and Portland carbonate respectively, at all studied 257 observation times, as shown in figure 6. The experiments therefore calibrate 258 our model at the pore scale. As expected, transport is the most heteroge-259 neous in the Portland sample, and the least in the beadpack. Our model 260 matches the persistently dominant stagnant region in the Portland sample, 261 which is a manifestation of transport heterogeneity. This is discussed in more 262 detail in [25]. 263

# 5.2. Comparison with NMR Experiments of Transport Involving Mineral Dis solution

The propagators obtained from NMR measurement of reactive transport experiments are now reproduced. The model is calibrated with experimental data of porosities  $\phi$  and propagators in a number of slices along the sample measured at the beginning and the end of the dissolution process. Transport in each slice of the core is computed in a lattice consisting of  $100 \times 100 \times$ 100 links representing a cube of side length  $8 \times 10^{-3}$  m. Each link in the



Figure 6: Computed probability of particle displacement  $P(\zeta) \times \langle \zeta \rangle_0$  as a function of displacement  $\zeta / \langle \zeta \rangle_0$  (solid lines), compared with the propagators obtained with NMR experiments by [2] (dashed lines) for  $\Delta = 0.2, 0.45, 1$  seconds.

lattice has a length of  $8 \times 10^{-5}$  m, which is the resolution of our pore-scale 272 simulation. 100,000 particles are launched at random locations in the lattice 273 at t = 0 s. Periodic boundary conditions are applied at the inlet and outlet 274 faces. If a particle exits the inlet or outlet, it is randomly reassigned to the 275 opposite face using a flux-weighted rule. At this scale, transport is governed 276 by  $\psi_P$  according to equation (1). The CTRW parameters  $t_1, t_2$  are computed 277 using the knowledge of flow rate  $Q = 8.3 \times 10^{-7} \text{ m}^3 \text{ s}^{-1}$ , molecular diffusion 278 coefficient, and porosity  $\phi$  within that slice, from which interstitial velocity 279 v can be computed. For example, for the slice with initial porosity  $\phi = 0.24$ , 280 the initial interstitial velocity is  $v = 3.06 \times 10^{-3} \text{ m s}^{-1}$ . Hence  $t_1 = l/v$  is 281 0.0261 s, whereas  $t_2 = l^2/D_m = 3.03$  s. This leaves  $\beta$  as the only tunable 282 parameter. We match our computations in each lattice with the propagators 283 measured within each slice. 284

In figure 7 the porosity profiles at the beginning and the end of the dissolution process are given. Then five propagator profiles at various locations



Figure 7: (a) Evolution of porosities followed by the initial and final (after reaction) propagators at location: (b)  $7.1 \times 10^{-3}$ ; (c)  $2.57 \times 10^{-2}$ ; (d)  $5.22 \times 10^{-2}$ ; (e)  $1.77 \times 10^{-2}$ ; and (f)  $4.16 \times 10^{-2}$  m from the inlet. Propagators are reproduced numerically using  $\psi$  according to equation (1). The corresponding  $\beta$ s are shown in the figures. The propagators are matched with experimental data i.e, dotted lines are the computed ones whereas solid lines are measurements.

along the core—computed at initial and final times respectively—are plotted and matched with the propagators computed according to unique and different  $\beta$  values. First it is worthy of note that the numerical results satisfactorily match the experimental data. Second, as dissolution takes place along the core,  $\beta$  values at the front of the experiments become smaller.

As in section 5.1, the propagators can be computed beyond  $\Delta = 0.25$  s.



Figure 8: Prediction of propagators at larger observation times  $\Delta s$ . We predict numerically that at around 1.6 s mark, a mobile region at around the main displacement will start to occur.

In figure 8 the propagators can be observed, computed initially with  $\beta = 0.8$ and 0.62, with  $t_1$  and  $t_2$  0.0275 and 9.6 s respectively, plotted at observation times  $\Delta$  up to 1.6 s. Our model shows that the stagnant, diffusion dominated regimes persist even at later observation times. Asymptotically, according to [29], Fickian behaviour should be expected at t much larger than  $t_2$ .

#### <sup>298</sup> 5.3. Core-plug Scale CTRW Parameters

Transport at the core-plug scale is modelled as a hop governed by  $\psi_{CP}(t)$ 299 which is obtained using the methodology illustrated in figure 4. At this scale, 300 transport is modelled in each  $8.8 \times 10^{-4}$  m-thick slice of the core using a cubic 301 lattice, with side length  $8.8 \times 10^{-4}$  m, consisting of  $100 \times 100 \times 100$  links; 302 see figure 9. Particles are injected into the inlet face at t = 0 and record the 303 transit-time required by each particle to reach the outlet face. Transport in 304 each link is governed by  $\psi_P = \psi(\beta_P, t_1, t_2)$  where  $\beta_P$  is calibrated for every 305 slice along the core i.e, by matching the NMR-measured propagators during 306 dissolution. An example of how  $\beta_P$  is obtained has been reported in section 307 5.2, i.e. by computing the propagators in each section along the core and 308 matching them with those measured in the experiments.  $t_1 = l/v$  can be 309 determined by knowing the constant flow rate  $Q = 8.3 \times 10^{-7} \text{ m}^3 \text{ s}^{-1}$ , and 310 porosities  $\phi$  of each slice. For example, for the slice where porosity  $\phi = 0.32$ , 311 the interstitial velocity is  $v = 2.74 \times 10^{-3}$  m s<sup>-1</sup>. Hence  $t_1 = l/v$  is  $3.2 \times 10^{-3}$ 312 s. The diffusion cut-off time  $t_2 = l^2/D_m$  is 0.036 s in a single micron-scale 313 link.  $\psi_{CP} = \psi(\beta_{CP})$  for every  $8.8 \times 10^{-4}$  m-thick slices are obtained by 314 matching the emergent transit-time distribution with equation (1). 315



Figure 9: Transport in each slice along the core is represented as a single hop governed by  $\psi_{CP}(t)$ . Each slice is represented as a 3D lattice consisting of  $100 \times 100 \times 100$  links. In each link, transport is governed by  $\psi_P(t) = \psi(t_1, t_2, \beta_P)$ .  $t_1, t_2$  are computed using the knowledge of flow rate,  $Q = 8.3 \times 10^{-7}$  m<sup>3</sup> s<sup>-1</sup>, and porosity  $\phi$ . For each slice,  $\beta_P$  has been calibrated by matching the NMR-measured propagators. Next, we use the upscaling methodology [1] to obtain  $\psi_{CP}(t)$ .  $\psi_{CP}(t)$  for every slice is tabulated in Appendix B.

 $\psi_{CP}(t)$  from  $\psi_{P}(t)$  are obtained using the upscaling methodology pre-316 sented in [1], which is illustrated in figure 9. First, we run a particle tracking 317 simulation in each core-plug lattice described above. Then, the emergent 318 transit-time distribution  $\psi_{CP}$  is matched with equation 1 by selecting the 319 correct  $\beta_{CP}$  value whereas  $t_1$ , and  $t_2$  are assigned according to the lattice di-320 mension, flow rate, and porosities at before and after dissolution. The list of 321  $\beta_{CP}$  coefficients, and measured porosities, obtained before and after reaction, 322 at a number of points along the core, can be found in Appendix B. After re-323 action, we found that the markedly increased porosities especially in the first 324 half of the core near the inlet, do not lead to a more homogeneous spread of 325 particle displacements. Rather, the overall transport process becomes more 326 heterogeneous as shown by the change of propagator profiles before and after 327 reaction. Quantitatively, this is shown by the decrease of  $\beta_{CP}$ . This shows 328 that emergent channels in the core result in some particles experiencing an 329 increase in velocity, whereas other particles who remain in the slower regions 330 now become even more stagnant in comparison. The non-Fickian features 331 are more pronounced after dissolution. This is characterised and quantified 332 by a smaller beta values after dissolution. A smaller beta value means a 333 higher probability of long transit times. This is apparent when seeing the 334 propagator profiles where the most common displacement after dissolution 335

### is much smaller than the average.

### 337 6. Transport at the Core Scale: Obtaining $\psi_C(t)$



Figure 10: 3D cylindrical lattice for core-scale simulation with diameter  $3.8 \times 10^{-2}$  m, and length  $7.16 \times 10^{-2}$  m. The computational domain is a 3D lattice consisting of  $40 \times 40 \times 82$ links. v(x, y) = 0 for  $x^2 + y^2 > (\frac{3.81 \times 10^{-2}}{2})^2$  m<sup>2</sup>. Transport in each link is governed by  $\psi_{CP}(t)$ . Then the upscaling method [1] is used to obtain  $\psi_C(t)$ .

At the core scale, transport can be interpreted as a single hop with corresponding  $\psi_C$ . For transport at this scale a cylindrical lattice is used, see figure 10, with length and diameter similar to the core plug used in the experiments. Within each link in the lattice, transport is governed by a transit-time distribution  $\psi_{CP}$ , which has previously been computed. The diameter of the lattice is 3.8 cm, and the length 7.16 cm—identical to the core used in the experiments. The lattice comprises  $40 \times 40 \times 82$  links.

The relation  $\psi_{CP} = Ae^{-t/t_2}(1+t/t_1)^{-1+\beta_{CP}}$  is applied in each link with  $t_1$ and  $t_2$  equal to 0.3 s and 1161 s respectively. 100,000 particles are launched at the inlet face at t = 0 s.

The flux Q is determined from experiment. Having measured the porosity 348 of every slice along the core, the interstitial velocity v is computed for every 349 link along the flow direction. The times required for each particle to break 350 through and reach the outlet face are recorded. The emergent transit-time 351 distribution function is plotted for three Pe numbers—59.3, 593 (the Pe352 number of the experiments), and 5930—before and after reaction, see figure 353 11, and have them matched to a functional form i.e, equation (1). Thus the 354 corresponding  $\beta_C$  at initial and final experimental times can be obtained i.e. 355



(b) After reaction

Figure 11: Core-scale ensemble averaged transit-time distribution function  $\psi_C(\tau)$ , at initial  $t_{\exp} = 0$  and final time  $t_{\exp} = 11800$  seconds, where  $\tau = t/t_1$  are the normalized transit times. Three *Pe* numbers are used where 593 is the *Pe* number of the experiments.

<sup>356</sup> 0.75 and 0.65 respectively. Note that the power-law behaviour continues to <sup>357</sup> exist at the core, cm, scale.

Prior to reaction, the emergent transit-time distribution function showed a power law behaviour with  $\beta_C = 0.75$ . According to [31], in this region i.e,  $1/2 < \beta < 1$ , the longitudinal dispersion is *super* diffusive. This can be seen from the propagators (figure 7). Initially, the bulk of displacement occurs below the mean displacement. With reaction, transport becomes more heterogeneous and even more super-diffusive, as seen in other dissolution experiments [40].

### <sup>365</sup> 7. Conclusions

A robust multiscale modelling of transport based on CTRW is validated with a combination of NMR imaging and transport experimentation to study

reactive transport signatures at a pore, and core-plug scale. For the specific 368 conditions investigated in our work, the analysis of the propagators before 369 and after reaction show that transport becomes more heterogeneous after 370 reaction. The present model reproduces these results well and quantifies the 371 increase in heterogeneity by the decrease of  $\beta$  values. For a beadpack, Ben-372 theimer sandstone, and Portland carbonate systems, the numerical results 373 agree with the experimental data, validating the pore-scale CTRW model 374 for different porous-media heterogeneities. 375

By predicting the propagators at longer observation times, it is shown that non-Fickian behaviour persists at the cm scale. Truncated power law behaviour is demonstrated for transport at the core-plug scale, emerging from the pore-scale representation of heterogeneity.

In future work the modelling could be extended to include reactive transport at the pore-scale and hence predict the change in transport properties with time.

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### <sup>604</sup> Appendix A. Pressure solver

Flow in each link is computed by solving mass conservation equation in each node *i* such that for each link *k* adjacent to node *i*,  $\sum_{k} q_k = 0$  applies, or in a matrix form,

$$[B] \mathbf{q} = \mathbf{0} \tag{A.1}$$

where [B] is an incidence matrix -  $N_k \times N_i$  where  $N_k$  is the number of links and  $N_i$  is the number of nodes - describing the topology of network.  $\mathbf{q} = q_k$ is the mass flux vector. Applying Darcy's law, flux can be expressed in terms of the pressure drop such that

$$\mathbf{q} = -\left[C\right]\left[B\right]^T \mathbf{p},\tag{A.2}$$

where [C] is a conductivity matrix; an  $N_k \times N_i$  diagonal matrix with entries  $C_k = \frac{K_k A_k}{l_k}$  where K is the permeability of the link, A the cross-sectional area,  $\mu$  the viscosity and l the length of the link. Substituting equation A.2 into A.1 we derive

$$[B] [C] [B]^T \mathbf{p} = \mathbf{0}.$$
(A.3)

The nodal pressure p in the network is obtained by solving the linear equation
A.3 using MUMPS: a MUltifrontal Massively Parallel sparse direct Solver—
see [42].

### <sup>619</sup> Appendix B. Porosities and the Corresponding $\beta_{CP}$ Coefficients

The table below shows the porosities and the corresponding  $\beta_{CP}$  coefficients along the core, before and after the experiment. The porosities are measured, whereas the  $\beta_{CP}$  values are computed (see section 5.3).

Distance from inlet	Porosity $\phi$		$\beta_{CP}$	
$(m \times 10^{-2})$	Before	After	Before	After
0.00	0.32	0.59	0.65	0.46
0.09	0.27	0.48	0.77	0.51
0.18	0.26	0.42	0.79	0.51
0.27	0.25	0.38	0.81	0.59
0.35	0.25	0.35	0.80	0.59
0.44	0.25	0.33	0.81	0.61
0.53	0.25	0.32	0.79	0.60
0.62	0.25	0.31	0.83	0.61
0.71	0.25	0.30	0.84	0.62
0.80	0.25	0.29	0.83	0.71
0.88	0.25	0.28	0.82	0.70
0.97	0.25	0.28	0.82	0.71
1.06	0.25	0.28	0.83	0.71
1.15	0.25	0.27	0.82	0.70
1.24	0.25	0.27	0.82	0.75
1.33	0.25	0.26	0.84	0.76
1.42	0.24	0.26	0.84	0.77
1.50	0.25	0.26	0.82	0.76
1.59	0.25	0.27	0.81	0.69
1.68	0.24	0.26	0.86	0.79
1.77	0.24	0.26	0.85	0.73
1.86	0.24	0.26	0.84	0.75
1.95	0.24	0.26	0.84	0.75
2.03	0.24	0.26	0.83	0.74
2.12	0.24	0.26	0.84	0.77
2.21	0.25	0.26	0.85	0.78
2.30	0.25	0.27	0.85	0.78
2.39	0.25	0.26	0.83	0.79
2.48	0.25	0.26	0.87	0.77
2.57	0.24	0.26	0.88	0.80

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Distance from inlet	Porosity $\phi$		$\beta_{CP}$	
$(m \times 10^{-2})$	Before	After	Before	After
2.65	0.25	0.26	0.85	0.81
2.74	0.25	0.26	0.85	0.81
2.83	0.25	0.26	0.86	0.79
2.92	0.24	0.26	0.82	0.78
3.01	0.25	0.26	0.84	0.79
3.10	0.25	0.26	0.82	0.80
3.18	0.25	0.25	0.80	0.79
3.27	0.25	0.25	0.79	0.79
3.36	0.24	0.25	0.80	0.78
3.45	0.24	0.25	0.81	0.80
3.54	0.24	0.24	0.80	0.79
3.63	0.24	0.24	0.81	0.81
3.72	0.24	0.24	0.80	0.80
3.80	0.24	0.24	0.81	0.81
3.89	0.24	0.24	0.79	0.79
3.98	0.24	0.24	0.80	0.80
4.07	0.24	0.24	0.81	0.81
4.16	0.24	0.24	0.80	0.80
4.25	0.24	0.24	0.80	0.80
4.33	0.24	0.24	0.80	0.80
4.42	0.24	0.24	0.79	0.79
4.51	0.24	0.24	0.80	0.80
4.60	0.24	0.24	0.83	0.83
4.69	0.23	0.23	0.83	0.83
4.78	0.23	0.23	0.82	0.82
4.87	0.24	0.23	0.85	0.82
4.95	0.24	0.23	0.83	0.82
5.04	0.23	0.23	0.82	0.82
5.13	0.24	0.24	0.80	0.80
5.22	0.24	0.23	0.80	0.80
5.31	0.24	0.24	0.82	0.82
5.40	0.24	0.24	0.80	0.80
5.48	0.24	0.24	0.80	0.80
5.57	0.24	0.24	0.80	0.80

Distance from inlet	Porosity $\phi$		$\beta_{CP}$	
$(m \times 10^{-2})$	Before	After	Before	After
5.66	0.24	0.23	0.80	0.80
5.75	0.23	0.23	0.80	0.80
5.84	0.23	0.23	0.80	0.80
5.93	0.23	0.23	0.81	0.81
6.02	0.23	0.23	0.82	0.82
6.10	0.23	0.23	0.81	0.81
6.19	0.23	0.23	0.83	0.83
6.28	0.23	0.23	0.83	0.83
6.37	0.22	0.22	0.88	0.88
6.46	0.22	0.22	0.87	0.87
6.55	0.22	0.22	0.88	0.88
6.63	0.22	0.22	0.89	0.89
6.72	0.22	0.21	0.88	0.88
6.81	0.22	0.22	0.87	0.87
6.90	0.22	0.22	0.7	0.7
6.99	0.22	0.22	0.71	0.71
7.08	0.22	0.22	0.73	0.73
7.16	0.22	0.22	0.74	0.74