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Dynamics of contaminant flow through porous
media containing random adsorbers
Kaj Pettersson ^{1*} , Albin Nordlander ² , Angela Sasic Kalagasidis ² , Oskar Modin ² , Dario Maggiolo ¹
 ^{1*}Department of Mechanics & Maritime Sciences, Chalmers University of Technology, Chalmersplatsen 4, Göteborg, 412 96, Sweden. ²Department of Architecture & Civil Engineering, Chalmers University of Technology, Chalmersplatsen 4, Göteborg, 412 96, Sweden.
*Corresponding author(s). E-mail(s): kajp@chalmers.se;

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Contributing authors: Angela.Sasic@chalmers.se; oskar.modin@chalmers.se; maggiolo@chalmers.se;

Abstract

Many porous media are mixtures of inert and reactive materials, manifesting 13 spatio-chemical heterogeneity. We study the evolution of scalar transport in a 14 chemically heterogeneous material that mimics a green roof soil substrate, frac-15 tionally composed of inert and reactive adsorbing particles. These adsorbing 16 particles are equivalent to biochar within a real soil substrate. The scalar trans-17 port evolution is determined using experiments and simulations calibrated from 18 experimental data. Experiment 1 is used to determine the equilibrium capacity 19 and adsorption rate of two biochar types when immersed in a methylene blue 20 solution. Breakthrough curves of a packed bed of glass beads with randomly 21 interspersed biochar are determined in experiment 2. Simulations are then run 22 to investigate the solute transport and adsorption dynamics at the pore-scale. 23 An analytical model is proposed to capture the behavior of the biochar adsorp-24 tion capacity and the simulation results are compared with experiment 2. A 25 pore-scale analysis showed that uniformly sized beds are superior in contaminant 26 27 breakthrough reduction, which is related to the adsorptive surface area and the 28 rate at which adsorption capacity is reached. Cases using the adsorption capacity model display a tight distribution of particle surface concentration at later simu-29 lation times, indicating maximum possible adsorption. The beds with dissimilar 30

particle sizes create more channeling effects which reduce adsorptive particle efficiency and consequently higher breakthrough concentration profiles. Comparison between experiments and simulations show good agreement. Improved biochar performance can be achieved by maintaining particle size uniformity alongside high adsorption capacity and adsorption rates appropriate to the rainfall intensity.

37 **Keywords:** biochar, lattice Boltzmann, experiment, adsorption, methylene blue

³⁸ Article Highlights

- ³⁹ Experiments on biochar adsorption of methylene blue compared to simulations, with
- 40 good agreement.
- ⁴¹ Simple analytical model proposed to capture biochar adsorption capacity, results
- ⁴² compared to experiments.
- Performance inferior in polydisperse beds due to underutilized particles and
 inhomogeneous concentration front profile.

45 1 Introduction

Porous media containing a proportion of inert and chemically reactive elements can found in biology in the form of biofilters, bioreactors, and organic tissue. They are 47 also found in geological elements such as rocks or soils, which are also highly chemi-48 cally heterogeneous, containing a wide variety of chemically dissimilar minerals and/or 49 organic elements. Research into this type of porous media has formerly been primar-50 ily driven by the petrochemical industry, however this has shifted in recent years to 51 a more environmental focus. Green roofs are one such example, consisting of living 52 greenery ranging from trees to grasses growing in a soil substrate whose composition 53 can vary widely, from crushed brick to manure to peat moss. The benefits of green 54 roofs in urban environments are well documented, such as their contributions to the 55 reduction in urban noise, air, and water pollution; their effects on the urban heat island 56

⁵⁷ as well as building envelopes themselves; and urban rainwater runoff management
⁵⁸ Aguilar Fajardo et al. (2022).

Green roofs require regular maintenance, including the administration of fertilizer 59 upon installation and potentially thereafter as the need arises. The fertilizer consists 60 of a fast-acting agent which gives the plants a boost to survive the initial period of 61 installation and establish themselves; and longer-acting nutrients which aim to keep 62 the additional required maintenance to a minimum. Excess fertilizer which cannot be 63 stored by the soil or used directly by the vegetation can be carried by rainwater from 64 the soil through the drainage system to locations where it acts as a contaminant Wang 65 et al. (2017). The addition of biochar to the soil to adsorb excess fertilizer (contaminant 66 or solute) is currently being put forth as viable solution to this problem. 67

⁶⁸ 1.1 Biochar in green roofs

Biochar is essentially any organic material that has been carbonized under high temperature through a process known as pyrolysis and can vary in material properties and morphology significantly. A thorough overview of the types of biochar in use commercially can be found in Novotný et al. (2023).

Biochar is incorporated into green roof soil via three strategies; random application 73 to the top of the bedding material, thorough mixing into the substrate composition 74 prior to vegetation, and set as a layer at the bottom of the substrate. Thorough 75 mixing and layering at the soil base are considered the most effective, with the latter 76 shown to be effective for reducing the leaching of total nitrogen and total phosphorous 77 Kuoppamäki et al. (2016). The proportion of applied biochar varies, from 5% by 78 weight/volume up to about 40%, however excessive proportions of biochar will have 79 adverse effects on plant growth and contribute to increased contamination Xiang et al. 80 (2021).81

There are few large-scale projects incorporating biochar use on green roofs though a few exist, such as the NWE CASCADE project, in which France intends to implement the use of biochar to enhance stormwater management within the Brittany region. Other similar solutions will likely be implemented within a larger initiative led by Bloomberg Philanthropies as touched upon in Senadheera et al. (2024).

The industrial-scale implementation is dependent upon several factors, such as the 87 ecological impact of large-scale biochar use and the release of additional contaminants present within or on the surface of biochar as a result of its preparation and function. 89 For example, Premarathna et al. (2023) discusses the almost complete removal of 90 ammonia from the soil when excessive biochar is applied, which can be problematic 91 for soil health. Another key factor is availability of biochar itself, given the different 92 feedstock and treatment methods for its creation. A final major consideration is the 93 lifetime of the biochar as an adsorbent and its end-of-life handling. Little work has been 94 done on the effects of aging on biochar performance due to limited long-term projects 95 but what is known is that degrading performance is dependent upon the contaminant and environmental conditions. For example, if the primary goal of the biochar is to 97 trap micro and nano-plastics (MPs and NPs) then both the aging of the biochar and 98 the plastics themselves play an important role in the removal efficiency J_i et al. (2024). 99 Physical degradation of the biochar also occurs during the aging process and leads 100 to biochar dust, which no longer serves its purpose of entrapping pollutants but can 101 act to spread captured plastics or heavy metals due to its increased mobility. It has 102 been shown the granulated biochar in particular is more resistant to this degradation 103 process, thus making it more suitable for applications such as green roofs where the 104 environmental exposure is high Lee and Kwon (2024). 105

Related to the issue of biochar disposal is the issue of regeneration, which heavily depends upon the captured contaminants. It is possible to regenerate biochar that has captured MPs/NPs using organic solvents such as acetone, however this is not

feasible in a green roof environment due to environmental concerns. Water rinsing 109 performs relatively poorly for removing carbon-based adsorbents but can be used Ji 110 et al. (2024). Adsorbed copper can be almost completely recovered Bashir et al. (2023), 111 as can cadmium Cui et al. (2022). Biochar regeneration over five adsorption-desorption 112 cycles was assessed with regard to volatile organic compounds (VOCs) with reported 113 88% to 96% regeneration Rajabi et al. (2021). While these reports are promising, the 114 scalability for some of the processes used for regeneration is in doubt as they are either 115 inappropriate for green roofs due to their open nature, or less efficient if alternatives 116 are used Gupta et al. (2020). 117

Biochar may raise the production costs of green roofs, resulting in higher final prices 118 for these products Khan et al. (2021). To evaluate its profitability for both producers 119 and end users, it is essential to compare its effects against using fertilizers alone or in 120 combination with fertilizers Ye et al. (2020). Additionally, given its environmental ben-121 efits—such as minimizing nutrient leakage and boosting carbon sequestration—efforts 122 are underway to improve cost calculation models to include both private and soci-123 etal costs and benefits Campion et al. (2023). However, these calculations are highly 124 case-specific, influenced by factors like location, feedstock, scale, pyrolysis conditions, 125 biochar pricing, and crop type. To promote greater adoption of biochar, Campion 126 et al. (2023) suggest developing standardized calculation models and emphasizing 127 additional societal benefits of biochar, such as improved water retention. While the 128 proposed model is comprehensive, it remains unclear how potential hazards associated 129 with biochar Xiang et al. (2021) will be addressed with it. Thus, the technoeconomic 130 benefits of biochar will be a focus of future research. 131

132 1.2 Biochar adsorption capabilities

Many variants of biochar have been employed for the purpose of removing contaminants in a soil or packed bed. Afroze et al. (2016) determined the adsorption of

methylene blue (MB) by eucalyptus bark biomass in a packed bed and found that it 135 performs well in the removal of dye-containing effluents. Dawood et al. (2019) ana-136 lyzed the adsorption of MB in pine cone biochar in a packed bed column of Kaolin 137 clay and utilized several analytical models to determine breakthrough curves. These 138 curves were compared under varying experimental conditions and the biochar/clay 139 packed bed was found to perform best under low flow rates, high MB concentration, 140 and larger bed depth. Zanin Lima et al. (2023) analyzed competitive sorption and des-141 orption of zinc, cadmium, and lead between compost, biochar, and peat. They found 142 that biochar had the highest adsorption capacity and lowest desorption rate, with lead 143 removal being the most effective. Beesley et al. (2010) examined the efficacy of biochar 144 and greenwaste compost on the reduction of zinc, cadmium, and copper within soils 145 and found them to be beneficial in the control of pollutants. Pita et al. (2024) provided 146 a review of the work associated with the use of biomass on the removal of pharma-147 ceutical compounds and reported that biochar made from a variety of plants; such as 148 rice husks, corn, sawdust, and sugar cane, was the most effective in many industrial 149 applications, partly due to their high adsorption capacity. While the aforementioned 150 studies are valuable in understanding biochar as an adsorbent for a variety of contam-151 inants, we are unable to identify the mechanisms by which it performs best, such as 152 available adsorbing surface area, effect on the flow field, or their pore storage capac-153 ity. For this level of detail we must utilize numerical models which can describe such 154 systems and provide the level of detail we require. 155

The physical process of contaminant adsorption in a flow field is described mathematically by the advection-diffusion of species, combined with adsorption/desorption; a model which is present across many fields of application. This general model is used to describe processes such as post-combustion CO_2 capture Pröll et al. (2016), determining optimal geometry of granular heat exchangers Mitra et al. (2018), analyzing methane adsorption in subsurface shale deposits Li et al. (2016), and describing the

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behavior of photocatalytic textiles Robin et al. (2016) to name a few. Investigations 162 into the dominant factors affecting adsorption rates and capacity have shown that 163 particle and pore morphology play an important role Liapis et al. (1999); Jareteg 164 et al. (2022). Adsorption rates themselves also strongly determine the process evolu-165 tion Marin et al. (2014), with lower rates determining adsorption quantity by reaction 166 duration, and higher rates controlling quantity through reaction rate Zakirov et al. 167 (2023). Non-isothermal systems can result in high heterogeneity of concentration and 168 temperature, with performance suffering from regions of stagnant flow and increased 169 near-wall breakthrough Verma and Mewes (2009). 170

When one considers a transported reactive species adsorbed locally by reactive 171 elements within a porous matrix; understanding the underlying interactions which 172 determine the local concentration distribution is necessary. The mixing itself within 173 porous media has been shown to be complex at high Reynolds numbers (Re). As such, 174 the flow is characterized by an initial advection-dominated regime as the flow pene-175 trates the pore network and is followed by a diffusion-dominated regime wherein the 176 molecular diffusion redistributes the scalar more effectively than the advection. The 177 advective regime consists of stretching of the scalar front in longer finger-like struc-178 tures, particularly where the flow is strongest, and can be defined by by the plume's 179 concentration mean, variance, and probability distribution Bonazzi et al. (2023). A 180 thorough overview of current state of research and methods used to solve species 181 transport in porous media can be found in Dentz et al. (2023). 182

At low Re, over a short time these structures regress to a more uniform distribution through molecular diffusion. However, even at low Re, the process of homogenization by diffusion may be retarded by the presence of local chemical heterogeneities in the form of solid adsorbing particles which affect the concentration field. The effect of varying the quantity of uniformly distributed adsorbers within a porous medium was undertaken previously, with two regimes in the evolution of the concentration field

¹⁸⁹ identified and the rate at which one regime reaches the other is determined by the
¹⁹⁰ number of adsorbing elements present Maggiolo et al. (2023).

The previous investigations cover primarily changes in adsorptivity, usually a static 191 value, with the values not necessarily reflecting realistic conditions. The geometries 192 generally tend to be regular, not reflecting in any way a realistic soil sample. We 193 will extend this characterization by introducing the presence of polydisperse particles 194 within the packed bed as well as an adsorption rate dependent upon a proposed adsorp-195 tion capacity model for the biochar. In addition, we make use of experimental data to 196 correctly tune the simulations to realistic adsorption rates and determine how these 197 factors impact the previously identified regimes as well as report additional influences 198 on the biochar performance. The results are directly compared to experimental data 199 to assess the predictive accuracy of our model, quantified by breakthrough curves. 200

²⁰¹ 2 Experiments

The experiments outlined in this section have been undertaken for two purposes. Experiment 1 was used calculate the adsorption capacity of the biochar which in turn is an input in the numerical simulations used to compare against experiment 2. The results for experiment 2 were directly compared to the numerical results to determine the accuracy of the model used in this work. All experimental information is given in this section, with the common materials introduced first, followed by each experiment and its respective results.

The adsorption capabilities of two types of biochar were experimentally determined: wood-chip biochar (WCB) produced mainly from European spruce and granulated biochar (GB) produced mainly from agricultural seed waste. The wood chip biochar was produced by Hjelmsäters Fastigheter AB while the granulated biochar was produced by Skånefrö AB; both companies being based in Sweden. Figure 1 displays

the types of biochar used in the experiments. Alongside the types of biochar, acidwashed glass beads were also used and acted both as an inert, transparent packing material for the biochar and as a control case with minimal to no adsorption.

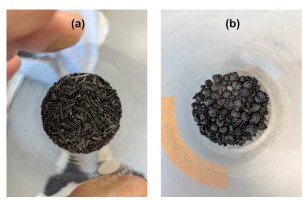


Fig. 1: Types of biochar (a) WCB (Hjelmsäters Fastigheter AB) and (b) GB (Skånefrö AB).

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²¹⁷ MB, chemical formula $C_{16}H_{18}CIN_3S$, was used as the adsorbate; diluted to 6 mg/L ²¹⁸ in a phosphate buffer solution containing 380 mg/L KH₂PO₄ and 300 mg/L K₂HPO₄. ²¹⁹ It is worth noting that due to its properties, methylene blue can be seen as an analog ²²⁰ for some organic pollutants however it is by no means representative of all possibilities. ²²¹ Other compounds or the direct pollutant itself may be required for more accuracy and ²²² the numerical model must be appropriately modified as well.

The MB concentration was measured using spectrophotometry with absorbance measured at 680nm. Figure 2 shows the experimental setup that was used for experiments 1 and 2.

226 2.1 Experiment 1

In experiment 1, a known mass of biochar or glass beads (1mm diameter) was placed
in a 5cm diameter Plexiglas column, as shown in Figure 3. The quantities of biochar

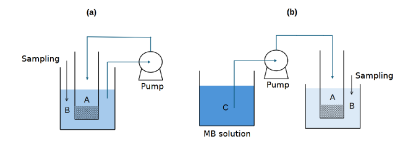


Fig. 2: Experimental setup for (a) experiment 1 and (b) experiment 2. A - Inner annular column containing porous medium, B - Outer annular beaker where MB sampling occurs, C - MB reservoir with original concentration c_0 used in experiment 2

²²⁹ were chosen such that differences in behavior would be evident while allowing for a steady-state to be reached within a reasonable time frame.

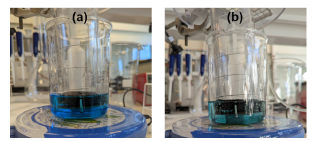


Fig. 3: Experiment 1 at (a) t = 0 min and (b) t = 135 min

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The bottom of the column was covered with a steel mesh which kept the biochar/-231 glass beads within the column. The column was submerged in a glass beaker containing 232 100mL of the MB solution, which was circulated through the biochar-packed column 233 using a peristaltic pump operating at a flow rate of 4mL/min. The column is fully 234 immersed in the MB solution to allow for the adsorption to occur at the maximum 235 possible rate, which is a property of the biochar itself rather than a consequence of 236 the local availability of adsorbent. Liquid samples from the solution being circulated 237 through the Plexiglas column were collected regularly from the glass beaker. The quan-238 tity of biochar used in this experiment is determined such that the measured difference 239 in adsorbance is clear as the quantity of biochar is increased.

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Figure 4 shows the evolution of the normalized outflow concentration over time in experiment 1. Absorbance (A) is converted to concentration and normalized against the initial concentration c_0 . We define absorbance as $A = \log_{10}(I_0/I)$ where I_0 is the intensity of incident light at 680nm wavelength and I is the transmitted intensity. There is no large difference in the performance for the cases using WCB, even when

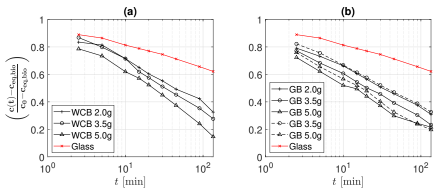


Fig. 4: Normalized outflow concentration for (a) Wood chip biochar (WCB), (b) Granulated biochar (GB)

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the quantity of biochar is increased to 5 grams; with the results for the GB being
similar. To calculate the adsorption capacity of the biochar we employ the conservation
equation

$$V_{bio}c_{eq,bio} = V_{liq}(c_0 - c(t_{end})) - V_{glass}c_{eq,glass},$$
(1)

where V_{bio} , V_{glass} and V_{liq} are the biochar, glass and liquid volumes, c_0 is the initial concentration, c is the measured concentration, and $c_{eq,bio}$, $c_{eq,glass}$ are the measured adsorption capacities (equilibrium concentrations) within the biochar and glass, which is the quantity at which they cease to adsorb addition solute, at t_{end} . Table 1 gives the substrate composition for experiment 1 and the resultant calculated adsorption capacity (equilibrium concentration).

Equation (1) is ideal for calculating the maximum adsorption capacity and biochar equilibrium concentration given that experiment 1 runs until a steady-state is reached,

 $3.3e - 4. [s^{-1}]$ Material WCB Glass GB 5.0Weight [g] 20.0 2.03.55.02.03.53.55.00.420.450.430.470.240.520.420.480.49 c_{eq}

Table 1: Experiment 1, Q = 4mL/min MB solution. WCB - woodchip biochar, GB - granulated biochar. Calculated adsorption rate k =

²⁵⁷ however this equation is unsuitable for calculating the evolution of the concentration
²⁵⁸ at the adsorbing surface over time. To this end we employ the formulation for the
²⁵⁹ evolution of the concentration at the adsorber surface

$$-V_p \frac{dc}{dt} = S_p k(c(t) - c_{eq,bio}), \qquad (2)$$
$$c(0) = c_0,$$
$$(t \to \infty) = c_{eq,bio},$$

 $_{260}$ where the solution takes the form

c

$$c(t) = (c_0 - c_{eq,bio})e^{-S_p kt/V_p} + c_{eq,bio},$$
(3)

where V_p, S_p are the reactive particle volume and surface area and the adsorption rate 261 is calculated by solving (3) for k, as all other quantities are known. This formulation 262 is based upon the assumption that the system is reaction-limited, which is valid since 263 the biochar is initially fully immersed in the solution. Fully immersed biochar allows 264 for the entire surface area of a reactive particle to contribute in the removal of the MB 265 with maximum efficiency, hence reaction-limited. The calculated adsorption rate is 266 used in the calibration of the simulations, as explained in section 3.2. It is important to 267 mention that the above formulations cannot be used for the analysis of experiment 2 as 268 it is mass-transport limited and thus we cannot make any assumption on the evolution 269 of adsorption in time. Thus we are forced to model the rate at which the adsorption 270

²⁷¹ capacity of the biochar is reached, and it's subsequent effect on the adsorption rate²⁷² itself.

If one wishes to apply a simple capacity model to the system, the resulting boundary condition at the reactive particle surface will be as in equation (4). Note that this condition bears striking similarity to (2) but is distinct, supporting our claim equation (3) is not suitable for solving the mass-transport limited problem. In these equations D_m is the molecular diffusivity, Da the Damköhler number, and f(c) the capacity model. This condition will be formalized in section 3.2 and is only presented here to demonstrate the differences in boundary condition between the experiments.

$$-V_{p}\frac{dc}{dt} = S_{p}D_{m}\frac{\partial c}{\partial n},$$

$$\frac{S_{p}D_{m}}{Q}\frac{\partial c}{\partial n} = \frac{kS_{\zeta}}{Q}f(c)c(t),$$
(4)

$$= \operatorname{Da} f(c)c^*|_{S_c}.$$
 (5)

280 2.2 Experiment 2

In experiment 2, the Plexiglass column was filled with glass beads mixed with WCB to 281 a height of 32mm. The biochar made up 15% of the column by volume and was either 282 evenly mixed with the glass beads, placed as a layer at the bottom of the column, or 283 placed as a layer at the top of the glass beads. The chosen fraction of 15% is motivated 284 by the standard composition used in green roofs in industrial production, which varies 285 but provides diminishing benefits beyond this fraction. The column was submerged in 286 a beaker filled with approximately 100mL phosphate solution without MB. The MB 287 solution was then fed to the top of the column at a flow rate of 8mL/min and the 288 MB concentration in the solution surrounding the column was measured in regular 289 time intervals. Table 2 outlines the cases shown in Figure 5, including variable particle 290 distributions and the equivalent weight of the included biochar. 291

Material Type	- %Biochar	Biochar [g]	Particle dist. [mm]
Glass Only	0.0	0.0	$\mu = 1.0$
Glass Only	0.0	0.0	$\mu = 1.0, \sigma = 0.5$
Glass + WCB	15.0	0.93	$\mu = 1.0, \sigma = 0.5$
Glass + WCB	15.0	0.93	$\mu = 1.0, \sigma = 0.5$
Glass + WCB	15.0	0.93	$\mu = 1.0$
Glass + WCB	15.0	0.93	$\mu = 1.0$

Table 2: Experiment 2, Q = 8mL/min with MB solution. WCB - wood-chip biochar.

The results of experiment 2 are shown in Figure 5 in the form of normalized concentration outflow, also called breakthrough. Once again the data is normalized against an initial concentration. We see an initial period where little of the MB is adsorbed,

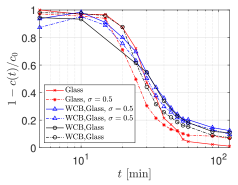


Fig. 5: Evolution of concentration breakthrough $1 - c(t)/c_0$

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whereafter the reaction rapidly reaches saturation within the biochar and little additional MB solution is adsorbed, leading to a "steady state" in the breakthrough curve. These periods; the initial adsorption, the rapid saturation, and the ceasing of additional adsorption can be characterized by the time intervals of i) 0 < t < 20, ii) 20 < t < 70, and iii) t > 70.

We see that the cases using only glass beads perform the worst, as expected. Interestingly these cases also adsorb some measure of MB, which means their behavior should be taken into account when examining the results of the cases which include biochar. Indeed, the inclusion of the quantities of biochar used in these experiments

show that they perform only marginally better than the glass beads themselves. Another important point to note from Figure 5 is that there is little discernible difference between cases in which a variety of particle sizes are present and cases where only a single particle size are used. This will be touched upon later in the numerical simulation results, particularly when the concept of adsorption capacity of reactive particles is present within the system.

310 3 Numerical approach

311 3.1 Lattice Boltzmann method

When one talks about solving the transport of a scalar, we formalize the concept using
the advection-diffusion equation

$$\frac{\partial c}{\partial t} = \nabla \cdot (D_m \nabla c) - \nabla \cdot (\mathbf{u}c) + R, \tag{6}$$

where c is a scalar (in our case concentration), t is time, D_m is the diffusion coefficient, **u** is the advecting fluid velocity, and R is a sink/source term related to reactive processes. At the micro-scale this equation can be represented using the Boltzmann transport equation, which describes motion through particle streaming as well as collisions.

It is an ideal choice for solving flows in porous media due to the complex geometry involved and allows for detailed information of the flow dynamics to be extracted at the pore scale. The system is solved on a lattice structure wherein each lattice element consists of a centroid and nodes placed on a cubic convex hull. The fictive particles travel along the lattice nodes governed by probabilities appropriate for the chosen lattice geometry such that the macroscopic properties of the fluid are preserved Succi (2001). A 3D regular cubic lattice with 19 degrees of freedom for movement (D3Q19) $_{326}$ is used and the solved equation is of the form

$$f_r(\mathbf{x} + \mathbf{c}_r \delta t, t + \delta t) - f_r(\mathbf{x}, t) = -\tau^{-1} (f_r(\mathbf{x}, t) - f_r^{eq}(\mathbf{x}, t)) + F_r$$
(7)

where $f_r(\mathbf{x}, t)$ is the distribution function at position \mathbf{x} and time t along the r-th direction; \mathbf{c}_r is the so-called discrete velocity vector along the r-th direction over time interval δt ; f_r^{eq} is the equilibrium distribution function; and τ is the mean collision time and is related to kinematic viscosity by $\nu = c_s^2(\tau - 0.5\delta t)$. The equilibrium distribution function $f_r^{eq}(\mathbf{x}, t)$ takes the form

$$f_r^{eq} = w_r \rho \left(1 - \frac{\mathbf{u} \cdot \mathbf{u}}{2c_s^2} \right), \ r = 1$$
(8)

$$f_r^{eq} = w_r \rho \left(1 + \frac{\mathbf{c}_r \cdot \mathbf{u}}{c_s^2} + \frac{(\mathbf{c}_r \cdot \mathbf{u})^2}{2c_s^4} - \frac{\mathbf{u} \cdot \mathbf{u}}{2c_s^2} \right), \ r = 2 - 19$$
(9)

where w_r is the appropriate weighting parameter for the D3Q19 lattice; ρ is the density; c_s is the speed of sound; and **u** is the velocity used for defining the equilibrium distribution functions, which can differ from the fluid hydrodynamic velocity, on the basis of the specific forcing scheme used.

The macroscopic flow quantities density and velocity, (ρ, \mathbf{u}) are thus related to the hydrodynamic moments as the following:

$$\rho = \sum_{r} f_r , \qquad (10)$$

$$\rho \mathbf{u} = \sum_{r} \mathbf{c}_{r} f_{r} + \frac{\Delta t}{2} \left(\frac{\Delta P}{L} \right), \tag{11}$$

where $\Delta t = 1$ in our case. The force F_r as formulated by Guo et al. (2002) is applied to the fluid, which mimics the flow rate intensity during a rain event in our case as

³⁴⁰ described later in more detail, and is given by

$$F_r = \left(1 - \frac{1}{2\tau}\right) w_r \left(\frac{\mathbf{c}_r - \mathbf{u}}{c_s^2} + \frac{\mathbf{c}_r \cdot \mathbf{u}}{c_s^4} \mathbf{c}_r\right) \left(\frac{\Delta P}{L}\right).$$
(12)

³⁴¹ 3.2 Solute adsorption implementation

³⁴² A solute adsorption rate is assigned to a percentage of chosen particles ζ such that ³⁴³ their surfaces S follow the first-order kinetics given by:

$$-\frac{\partial c^*}{\partial \lambda_s^*}|_{S_{\zeta}} = \text{Da} \ c^*|_{S_{\zeta}},\tag{13}$$

where Da is the Damköhler number, $c^* = c(\mathbf{x}, t)/c_0$ is the dimensionless concentration at position $\mathbf{x} = (x, y, z)$, and $\lambda_S^* = \lambda_S/d$ is the dimensionless direction pointing inward to the particle surface. The Damköhler number, a ratio of the reaction rate and the advective mass flow rate, is given by

$$\mathrm{Da} = kS_{\zeta}/Q,\tag{14}$$

where k is the adsorption rate on the particle surface, S_{ζ} is the total reactive surface area expressed as a fraction ζ of the total particle surface area, and the flow rate is given by $Q = l_0^2 \varepsilon U$. Here l_0 is the porous domain width, ε is the porosity, and U is the mean streamwise velocity.

Numerically, the concentration field $c(\mathbf{x}, t)$ is solved using a second population which is transported by the fluid velocity **u**. The solute is then initialized at the inlet of the porous media as a volume with concentration c_0 . The second lattice population,

denoted g_r , gives the local concentration field by

$$c(\mathbf{x},t) = \sum_{r} g_r(\mathbf{x},t).$$
(15)

We impose a Neumann boundary condition at the adsorbing surfaces for the scalar lattice Boltzmann quantity g_r . The distribution function at a fluid node \mathbf{x} in the proximity of an adsorbing surface placed at $(\mathbf{x} - \mathbf{c}_r)$ is corrected along the wall-normal direction \mathbf{r} as

$$g_r(\mathbf{x}, t+1) = \frac{-A_1 + A_2}{A_1 + A_2} g_r(\mathbf{x}, t) + \frac{2w_r A_3}{A_1 + A_2},$$
(16)

where $A_1 = k$, $A_2 = D_m$, and $A_3 = 0$. This condition was developed by Huang and Yong (2015); Huang et al. (2016).

In order to limit the reactivity as the particle surface reaches high levels of concentration we apply a functional coefficient to the prescribed reaction rate. This limit is designed to act as an analog to the particles reaching a saturated state wherein they can no longer adsorb more of the solute, known as adsorption capacity. Several different functions were tested, including a triangular function as well as a sigmoid function however a simple linear function was chosen, of the form

$$f(c) = 1 - c/c_{eq}, \,\forall c \le c_{eq} \tag{17}$$

which allows for full reactivity at low values of concentration but reaches zero when the particle equilibrium concentration has been reached. This equilibrium concentration is set to $c_{eq} = 0.5$ and is based upon the results of experiment 1. This alteration is reflected in the adsorption boundary condition, becoming

$$-\frac{\partial c^*}{\partial \lambda_s^*}|_{S_{\zeta}} = \text{Da } f(c)c^*|_{S_{\zeta}}, \tag{18}$$

which is the equivalent of equation (4) presented in the previous section.

373 3.3 Domain generation and operating conditions

The pore structure used in the simulations is a packed bed generated by the applica-374 tion of rigid body physics on falling spheres within a cylindrical container. Two such 375 cases are generated; i) a monodisperse case with a fixed particle diameter, and ii) a 376 polydisperse case with the mean diameter equal to that of the monodisperse case. The 377 particles are centered around a diameter of 1.0 with $2\sigma = \pm 0.82$ millimeters. A cubic 378 subdomain is then selected from the cylindrical container and the volume is discretized 379 into a binary matrix of $l_0^3 = 256^3$ computational nodes, giving the mean particle diam-380 eter d = 21.6 nodes. The method by which the domain is generated and characterized 381 can be found in Maggiolo et al. (2023) such that domain integrity is assured and the 382 volumetric porosity is calculated to be $\varepsilon = 0.4, 0.36$ for the mono- and polydisperse 383 cases, respectively. 384

In order to simulate an intense rainfall event the Péclet number Pe > 1 and we 385 assume little inertial influence, thus Re < 1. We set the Damköhler number, Da ≈ 1 , 386 which represents equal magnitudes of molecular diffusion and reaction rates, as calcu-387 lated in the experiment 2. By applying the conditions required above, the simulation 388 has Re = 0.017 and Pe = 6.7, which is representative of a rainfall event with intensity 389 of around 60mm/hr. The flow rate of experiment 2 is Q = 8mL/min, which converts 390 to about 600 mm/hr, about double the world record rainfall intensity, with Pe = 250 391 and Re = 0.17. It is worthwhile to note that intensities higher than the record can be 392 reached when runoff is channeled to a porous surface, which is not naturally occur-393 ring rainfall. While these values are higher than those chosen for the simulation, the 394 governing physics are identically balanced, as the originally defined conditions are all 395 satisfied, including $Da \approx 1$. With the above mentioned flow rates we have been able 396 to measure variations in concentration quantities in a few hours, which we found as a 397

good time lapse to minimize effects from changes in environmental conditions on the
 measurements.

These conditions are used to tune $\Delta P/L$, the molecular diffusivity and viscosity 400 accordingly. It is worth mentioning that common rainfalls are between 3-8mm/hr, 401 however less common extreme events can reach 15-60 mm/hr or more. We can extrap-402 olate the simulation conditions to lower rainfall intensities by accounting for the time 403 scale of the physical processes involved. The soil will not be as thin as that in the 404 simulations and no runoff infiltration front will be pure contaminant. Additionally, 405 one can alter the diffusivity to compensate such that the physically realistic process 406 is retained. 407

The resultant mean flow velocity is averaged over the entire domain, given by $U = (\int_{l_0^3} u_z dl_0^3)/l_0^3 \varepsilon$. The Kozeny-Carman relationship stipulates that if the porosity of a domain is altered the flow rate must be changed accordingly. Since in our cases porosity is approximately unaltered, the resultant mean velocity for the polydisperse case is the same. The biochar percentage is set to 30% in the simulations, double the industry maximum recommendation, to improve statistical analysis on single particle adsorption features without straying significantly from realistic conditions.

The initial concentration $c_0 = 1.0$ is applied to an inlet zone directly adjacent to 415 porous volume. The streamwise boundaries are assigned a periodic condition wherein 416 buffer zones are applied to the inlet and outlet boundaries to prevent any effect on the 417 flow within the porous zone. The lateral boundaries are assigned a free-slip condition. 418 The simulations are run in two steps; first the single phase velocity profile is solved 419 for each geometry and then the solute is added. In this way we solve only the steady-420 state flow in the first stage and only the concentration profile in time in the second 421 stage. Six cases total are run on two different geometries, the details of which can be 422 found in Table 3. 423

Table 3: Case numbers corresponding to operating conditions.Adsorption rate, particle type, andadsorption capacity are alteredbetween cases.

Case	k	Particle	Capacity
1	0.0	Mono	None
2	0.0	Poly	None
3	5.3e-6	Mono	Unlimited
4	5.3e-6	Poly	Unlimited
5	5.3e-6	Mono	Limited
6	5.3e-6	Poly	Limited

424 4 Results

425 4.1 Concentration profiles

We show a comparison of the concentration profile in the domain between the monodisperse and polydisperse geometries for all cases listed in Table 3.

Transverse cuts at $x/l_0 = 140/256$ are shown at the same timestep $t^* = 0.6$, given 428 by $t^* = tU/l_0$, which corresponds to the time at which the majority of the reactive 429 particles begin to reach their full capacity, in the cases with limited capacity. Particles 430 marked with black stars indicate reactive particles. Figures 6(a), (c), and (e) display 431 a more homogeneous front, all with the monodisperse geometry. We do see decreased 432 homogeneity in figures 6(b), (d), and (f), where we have the polydisperse geometry. 433 One can see clearly the difference in front homogeneity and indeed the formation of 434 tendrils of higher concentration within these figures. We also observe a general trend 435 of decreased homogeneity in the polydisperse cases with adsorption, (d) and (f), which 436 we investigate in more detail in subsequent sections. 437

438 4.2 Average surface concentration and flux ratio comparison

The average surface concentration of all reactive particles is given in Figure 7. This quantity is calculated by summing the concentration on all particle surface nodes and dividing by the total amount of such nodes. In Figure 7(a) there is a great similarity

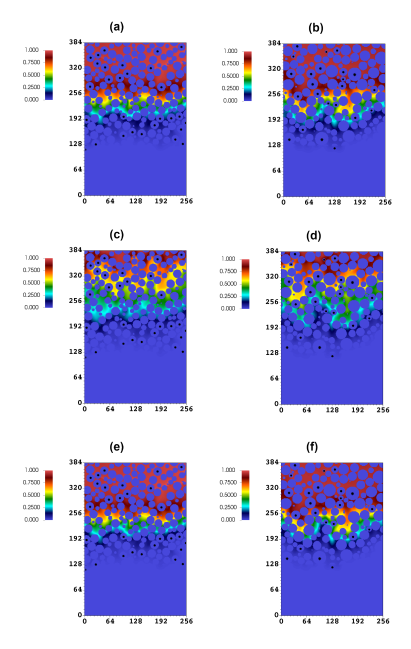


Fig. 6: Concentration profiles cut at $x/l_0 = 140/256$ and $t^* = 0.6 \approx 20$ min. Black stars indicate reactive particles. No capacity: (a) Case 1, (b) Case 2. Unlimited capacity: (c) Case 3, (d) Case 4. Limited capacity: (e) Case 5, (f) Case 6. The polydisperse cases (right panels) appear to generate a less homogeneous concentration front.

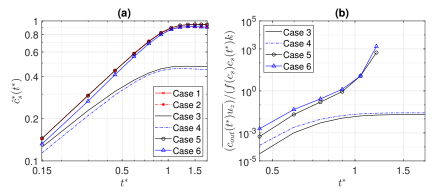


Fig. 7: (a) Average surface concentration and (b) breakthrough flux/adsorption flux ratio as a function of dimensionless time. No capacity: Case 1,2. Unlimited capacity: Case 3,4. Limited capacity: Case 5,6

of behavior between the cases with no capacity and those with a limited adsorption 442 capacity. We see the surface concentrations increase to near unity as the reactive 443 particles are no longer capable of removing more solute from the liquid. On the other 444 hand, if one examines the cases with unlimited capacity we see a significantly reduced 445 surface concentration, around 0.5. This is their effective equilibrium state and further 446 reduction or buildup will not occur without external stimulus. We also note that there 447 is little difference between the monodisperse and polydisperse geometries, implying a 448 similar total reactive surface area despite the difference in individual particle surface 449 areas, which is the case. 450

In Figure 7(b) the ratio of the breakthrough flux and the adsorption flux is shown. 451 This quantity can be seen as a measure of the medium leakage reduction as a con-452 sequence of biochar adsorption. Note that the cases with the limited capacity model 453 end prematurely due to the particles ceasing to adsorb after reaching their adsorption 454 capacity. The ratio in the beginning of the simulations is extremely small, indicating 455 adsorption is the dominant flux in the system, however this reduces significantly as 456 the simulations progress. This is expected since the concentration flux at the outlet 457 will increase over time as the solute travels through the medium. At longer times the 458

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ratio for the unlimited capacity cases stabilizes at around 10^{-2} indicating an equilibrium between the adsorption and the breakthrough flux throughout the simulation lifetime. This is not true in the cases with limited adsorption capacity, wherein we see an inflection point at which the particles cease to react and the breakthrough flux quickly becomes the only contributing factor. This period of approaching particle inertness is quite similar for both geometries, indicating a negligible effect of the front homogeneity on this macroscopic quantity.

466 4.3 Breakthrough curves (vs. experiments)

⁴⁶⁷ Breakthrough curves are calculated for all cases and displayed in Figure 8 where (a)
⁴⁶⁸ is the average concentration present in a small outlet volume the thickness of one
⁴⁶⁹ particle diameter calculated as

$$\overline{c_{out}(t^*)u_z} = \left(\int_{V_{out}} c_{out}(t^*)u_z \mathrm{d}V_{out}\right) / V_{out}$$
(19)

where $V_{out} = l_0^2 \varepsilon d$ and (b) illustrates the comparison of like cases between the experimental results and the simulations. In these graphs, a value of 1 represents a full

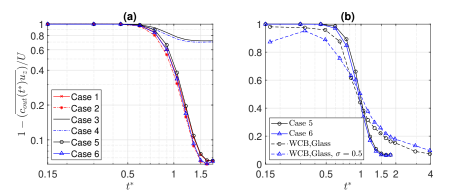


Fig. 8: Breakthrough curves for (a) simulation and (b) between like experimental cases and simulations. 1 - no solute reaches outlet, 0 - no adsorption has occurred, all solute reaches outlet. No capacity: Case 1,2. Unlimited capacity: Case 3,4. Limited capacity: Case 5,6

reaction wherein nothing reaches the outlet and 0 represents no change in concentra-471 tion between inlet and outlet, i.e. nothing has been removed. It is immediately clear 472 in Figure 8(a) that unlimited capacity cases perform significantly better in terms of 473 leakage than those which have a limit or have no reactivity. Interestingly, there is lit-474 the difference in performance between the cases with no adsorption capacity and those 475 with limited capacity. This is due to the fact that the limited capacity particles reach 476 capacity very swiftly and thus contribute very little to removing the solute after the 477 earlier stages of the simulation. 478

This hypothesis is given some weight by Figure 8(b) where two experimental cases 479 are directly compared to the simulation cases with limited adsorption capacity. We 480 find a very similar behavior in that both experiments and simulations swiftly reach 481 capacity and subsequently remove little of the solute. Note that there is an initial 482 period of time for the MB solution to reach the porous media through the piping and 483 we do not take wall effects into account in the simulation; both of which are present in 484 the experiment. Consequently, the experimental curves have a lower gradient and take 485 additional time to reach full capacity, however the steady-state behavior is correctly 486 reflected by the simulations. 487

At lower rainfall intensities we expect to see an increased residence time of the adsorbent near the biochar surfaces, however this will not matter if the biochar is already at capacity. In addition to the uncertain reaction kinetics, lower rainfall intensities may also result in unsaturated flow conditions, radically altering the system and introducing capillary forces, making any prediction based upon the current work void. Finally, a lower flow velocity may transition from more complex mixing caused by the microstructure to a more simplified laminar flow in larger pore spaces.

⁴⁹⁵ 4.4 Flow homogeneity and outlet flow profile

We examine the probability distribution for the streamwise and transverse velocity
magnitude as a quantification of the flow homogeneity within each geometry as well as compare between the two.

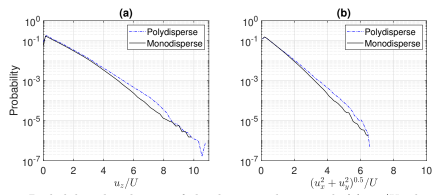


Fig. 9: Probability distribution of the domain velocity with (a) u_z/U where u_z is the streamwise velocity component and (b) $(u_x^2 + u_y^2)^{0.5}/U$ the transverse velocity magnitude

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Figure 9(a) shows the normalized streamwise velocity and it is clear that the 499 streamwise velocity is largely similar to that of the mean velocity. However, varia-500 tions up to 10 times that value exist within the flow field which is also reflected in the 501 distribution of the transverse velocity. There is no real discernible difference in the 502 distributions between the two geometries, thus we can be reasonably confident that 503 the flow alone is not a primary factor in notable differences observed between cases. 504 Indeed, we do see increased inhomogeneity induced by increased adsorption in the 505 front profiles shown in Figure 6. 506

Figure 10 displays the probability distributions for the breakthrough flux, measured $c(t^*)u_z/U$, as a quantification of the flow homogeneity at the outlet. This gives an indication of how the flow interacts with the reactive particles, producing tendrils

⁵¹⁰ of increased concentration as opposed to a more uniformly distributed concentration front.

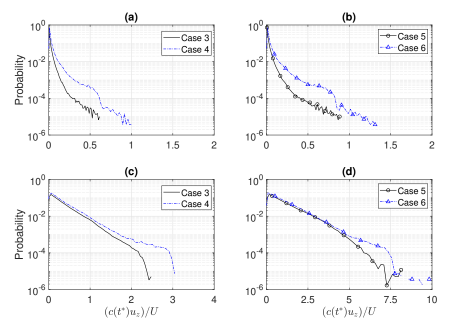


Fig. 10: Breakthrough flux distributions for (a) Unlimited capacity, $t^* = 0.6$ (b) limited capacity, $t^* = 0.6$ (c) Unlimited capacity, $t^* = 1.5$ (d) limited capacity, $t^* = 1.5$

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We compare the distributions during the period where the solute first reaches the 512 outlet $(t^* = 0.6)$, figures 10(a) (unlimited capacity) and 10(b) (limited capacity) and 513 when the steady state at the outlet is reached $(t^* = 1.5)$, figures 10(c) (unlimited) 514 and 10(d) (limited). A value near 1 indicates tendrils of high concentration trans-515 ported at a velocity around that of the streamwise mean velocity. Deviations from 516 this imply a change to the local concentration or streamwise velocity. In other words, 517 fluctuations around the mean of the concentration flux. Note a strong similarity in dis-518 tribution between 10(a) and 10(b), though the measured concentration flux is slightly 519 higher in the limited capacity case. In all cases the monodisperse geometry exhibits 520 a lower and tighter distribution of outflow concentration, indicating higher and more 521

efficient adsorption of the solute. The largest deviation is found in Figure 10(d), which compared to the unlimited capacity case in 10(c), has a higher concentration flux magnitude. We know from our analysis of the flow itself there is little influence on the flow induced by each geometry, thus the adsorption rate spatial distribution is the likely cause for this phenomenon.

527 4.5 Particle adsorption capacity

Figure 11 displays the PDFs of average particle surface concentration c_s^* during the transition stage $t^* = 0.6$, figures 11(a),(b) wherein particles are reaching their capacity and begin to become non-reactive. The lower figures, (c),(d) show the same cases, however the timestep is altered to reflect the steady-state distribution found at $t^* =$ 1.5. Figures 11(a),(c) are unlimited capacity and (b),(d) are limited capacity. The order of cases and times is replicated for Figure 12.

Figure 12 displays PDFs for the surface concentration flux $c_s^* S_p / S_{\mu}$, using the prefactor S_p / S_{μ} , where S_p is individual particle surface area and S_{μ} is mean particle surface area. This ratio is the measure of the available adsorptive surface area compared to that of the ideal surface area of a representative particle. When combined with a particle's surface concentration we can get an idea of which particles are more effective individually in terms of the adsorptive flux.

In Figure 11 we show the distribution of each particles' average surface concentration, taken at an intermediate and steady-state time. We observe no notable differences between the monodisperse and polydisperse geometries. At the intermediate time the distribution for the limited capacity cases exhibits the U-shaped distribution typical of diffusion-dominated systems, with two peaks at the extrema. In the cases with unlimited reactivity the peak at higher concentration disappears due to the continuous adsorption, figure 11(a). At the steady-state, the cases with limited adsorption

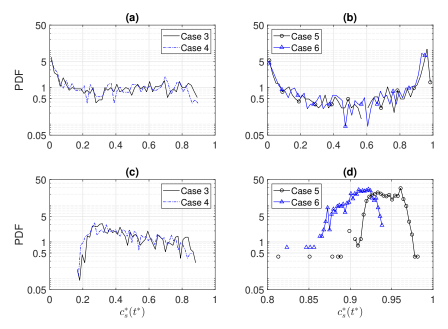


Fig. 11: Particle average surface concentration PDFs for (a),(b) $t^* = 0.6$ and (c),(d) $t^* = 1.5$. Unlimited capacity: Case 3,4. Limited capacity: Case 5,6

tend towards concentration of 1 with a very narrow distribution of values whereas the unlimited are more widely distributed, figures 11(c),(d).

⁵⁴⁹ When one examines the surface concentration flux $c_s^* S_p / S_\mu$ in the cases with unlim-⁵⁵⁰ ited capacity, figures 12(a),(c), we observe a similar behavior to that of the average ⁵⁵¹ particle surface concentration. As expected, a similar trend is also evident in the ⁵⁵² monodisperse case with limited capacity, figures 12(b),(d), when compared to the dis-⁵⁵³ tribution evolution in time of figures 11(b),(d). Indeed $S_p / S_\mu \approx 1$ in the monodisperse ⁵⁵⁴ case.

This is not reflected in the polydisperse case with limited capacity. At longer times where the concentrations are uniformly $c_s^* \approx 1$, the fluxes will tend to the distribution of S_p/S_μ , figure 12(d). Due to the fact that the particle radii are normally distributed, the probability of S_p/S_μ will be χ^2 distributed, exhibiting an exponential decay away from the mean.

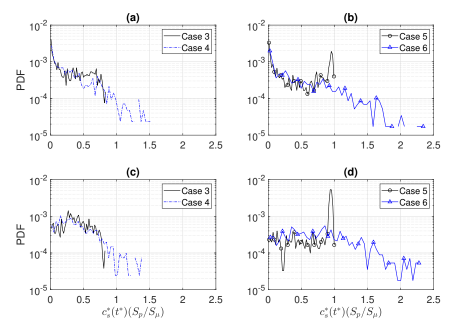


Fig. 12: Surface concentration flux PDFs for (a),(b) $t^* = 0.6$ and (c),(d) $t^* = 1.5$. Unlimited capacity: Case 3,4. Limited capacity: Case 5,6

Table 4: Ratio of mean particle size with con-
centration higher than concentration threshold
vs. mean particle size of the rest at $t^* = 0.6$ Concentration0.30000.50000.70000.9000Ratio of means0.99240.99080.95150.8853

Interestingly, at the intermediate time, the distribution of the fluxes preserves the 560 peak at low concentration whereas the peak at higher concentration visible in the 561 monodisperse case here is absent for the polydisperse case. We calculate the ratio of 562 mean particle size with concentration higher than a certain threshold against the mean 563 size of the remainder particles, shown in Table 4. It is noticeable that high surface con-564 centrations, that is above a specified threshold, are found on smaller particles. In turn 565 the distribution of fluxes $(c_s^* S_p / S_\mu)$ decreases uniformly as the flux itself increases, 566 figure 12(b). This may indicate that some particles in the polydisperse geometry are 567

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⁵⁶⁸ being underutilized. If one examines the cuts shown in Figure 6 where the polydis-⁵⁶⁹ perse cases display preferential pathing we see some occurrences of preferential paths ⁵⁷⁰ probably following high concentrations around smaller particles which have reached ⁵⁷¹ capacity.

The behavior discussed above regarding the monodisperse geometry is indicative 572 of a more homogeneous front, where the particles are either being approached by the 573 front or are surrounded by it, whereas the front reaches the particles in the polydisperse 574 geometry at different times, leading to a higher distribution of different magnitudes. 575 We note that the monodisperse case exhibits an overall higher surface concentration 576 at the steady-state than that of the polydisperse cases. This is coupled with a simi-577 larly lower breakthrough flux. One explanation for this is the increased inhomogeneity 578 observed in the concentration profiles for the polydisperse geometry. This preferential 579 pathing by the formation of tendrils of higher concentration removes solute from reach-580 ing particles that would otherwise aid in the removal of concentration from the flow. 581 This channeling effect reduces the efficiency of the reactive particles as some avail-582 able surface area is left unused, effectively underutilizing them. A more homogeneous 583 front allows for the maximum surface area to be reached at any given time. 584

Figure 13 shows the particle diameter and surface area distributions to aid in the 585 interpretation of the results as discussed above. Figure 13(a) shows the distribution 586 of reactive particles only and all particles for the polydisperse case as well as the 587 resultant normal distribution curve. The distribution of dimensionless surface area is 588 shown in Figure 13(b) for the polydisperse case only. It is compared to the ideal PDF 589 of diameter squared d^2 which follows a χ^2 distribution. The difference observed are 590 due to surface contact or cutting along the domain edges. Despite this fact, we are 591 satisfied that the distributions shown here justify the discussion we have entertained 592 regarding geometry effects and the potential for preferential pathing induced by the 593 polydisperse geometry in particular. 594

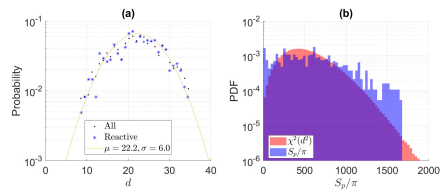


Fig. 13: (a) Particle diameter d distribution in l.u.. Dots indicate all domain particles, stars are just reactive particles. Line is the normal distribution function with $\mu = 22.2, \sigma = 6.0$. (b) PDF of reactive particle surface area S_p/π (blue), $\chi^2(d^2)$ distribution of particle diameter d (red)

We can now state with reasonable certainty that reactive particles do influence the concentration profiles, particularly the available reactive surface area. An advantage may be achieved by using more mondisperse particles, as they promote a more homogeneous front for the advancing solute and reduce the solute leakage, particularly in the case of particles with limited adsorbing capacity.

4.6 Modifying the reaction kinetics

We close the paper with a short discussion on the practicalities of accurately modelling 601 the adsorption dynamics present in reality on the surface of biochar. There are several 602 different mechanisms by which adsorption occurs; namely pore-filling, hydrophobic 603 interaction, surface complexation, electrostatic interaction, hydrogen bonding inter-604 action, $\pi - \pi$ electron-donor-acceptor (EDA) interaction, and intra-particle diffusion. 605 In addition it has been shown that thermal conditions as well as the pH of the envi-606 ronment play a non-negligible role in the reaction speed Ji et al. (2024); Gupta et al. 607 (2020), as does aging or the presence of dissolved organic matter. The sum total of 608 considering individual biochar characteristics, their intrinsic modes of adsorption, as 609 well as environmental factors renders the modelling a significant challenge. In general 610

current works have made use of first and second order pseudo-kinetic models along-611 side Langmuir and Freundlich isotherms. Some forays have been undertaken in the 612 realm of applied artificial neural networks and statistical chemical modelling however 613 this is still in its infancy for such an application. If one wishes to enhance the models 614 governing the reaction kinetics we need more experimental data to determine which of 615 the above modes of adsorption should be considered and how the environmental and 616 aging aspects can be included in the models. While some kind of pre-factors can be 617 easily applied they are not representative of any physical phenomenon and some possi-618 ble empirical model must be constructed which accurately captures these effects may 619 be a simple and efficient solution. Alternatively the full chemical species and reactions 620 can be included in the modeling but this undertaking may not be compatible with 621 how the lattice Boltzmann method operates but would ensure a more chemically con-622 sistent interaction. Pore filling and intra-particle diffusion might be handled via some 623 volumetric functions that account for internal pore volume and pore network but this 624 is also impractical to implement. Whichever strategy is used, more experimental data 625 is likely to be required and understanding the fundamental physical interactions in 626 such a system are a good place to aid in determining where to next add the increased 627 realism required for more accurate and applicable predictions. 628

⁶²⁹ 5 Conclusion

Experiments have been undertaken to determine the adsorption rate and capacity of two different kinds of biochar alongside breakthrough curves for same. The breakthrough curves are calculated using the absorbance of an MB solution over a period of time. Simulations are run using the experimental data and a comparison is made between the predicted breakthrough curves and the experimental results, with good agreement found.

A deeper investigation of the underlying factors determining the concentration field 636 evolution is described. This analysis consists of flow velocity and breakthrough con-637 centration profiles, average and individual surface concentration distributions, particle 638 capacity distributions, and a flux ratio comparison. The results show that the flow 639 velocity distribution remains the same for both simulated geometries, implying that 640 the differences in concentration distribution may not be a result of the flow velocity 641 profile. Examination of the breakthrough curves and individual particle concentra-642 tion distributions show that the polydisperse geometry promotes preferential pathing 643 throughout the medium and leads to a more varied distribution of particle states 644 than those found in the monodisperse cases. This preferential pathing is undesirable 645 in the context of removal of contaminants from the flow, as a more homogeneous 646 front ensures the randomly distributed reactive particles will interact with the solute, 647 whereas tendrils may bypass them entirely and flow directly for the outlet. 648

We have chosen to use spherical packed beds over a realistic soil substrate geometry due to the intrinsic inhomogeneity found within soil scans taken using x-ray microtomography (XMT) or any similar technique. A chosen domain may contain few very large particles and multitudes of tiny particles or anywhere in between, rendering the porosity of any such domains fluctuating wildly, making comparisons difficult. In future it is worthwhile to determine if the trends observed in this work hold true for the realistic soils as well.

If one extends these results to application recommendations, we can state that the observed differences between the mono- and polydisperse cases are larger than those observed when biochar is present or not. This would correspond to cases 1 and 2 (no biochar) and 5 and 6 (biochar with a limited capacity). The concept of limited adsorption capacity is analogous to the realistic adsorbance occurring in the experimental biochar, which cannot remove the solute indefinitely. These cases only slightly outperform the cases in which there are no reactive particles present. This

is due to the rate at which the limited capacity particles reach capacity and become inert. Cases 3 and 4 have unlimited capacity, which one could say is analogous with roots or some other living biological matter that does not have a capacity in the same manner as biochar.

The practical implication here is that monodisperse beds are preferred as they promote more even distribution of the solute within the domain and thus allow the random adsorbers to act with maximum effect, i.e. reaction-limited. Additionally, the presence of adsorbers with an unlimited adsorption capacity must be taken into account as this can dramatically effect the results, particularly in proximity to each other. This effect will obviously increase if the adsorption rate is increased.

If one wishes to increase the efficacy of biochar, a high absorptive rate with a 673 similarly high capacity will greatly reduce breakthrough. This can be accomplished 674 by modifying the material itself and/or increasing the total quantity of biochar within 675 the soil, which will consequently increase its capacity. Current practice suggests no 676 more than 15% biochar by volume is beneficial given common rainfall quantities; 677 however our results show that performance can be increased if one takes into account 678 the factors mentioned above, with the additional mention of total reactive surface 679 area and the idea of induced preferential pathing. Lower rainfall intensities will cause 680 higher residence time of the concentration front near the biochar; however once they 681 saturate, which we show can take place quite quickly, breakthrough will still occur. 682 Thus a more accurate method for modeling the capacity of different adsorbers should 683 be investigated to improve the predictive ability of the models. Finally, given that 684 monodisperse particles promote a more homogeneous concentration front and that 685 total reactive area matters more than individual particle size we suggest that more 686 uniform sized substrates are used. 687

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