# Good modeling practice for calibration applied to ion exchange breakthrough prediction

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# 12 Abstract

13 Ion exchange (IX) is a key technology in resource recovery processes for demineralization and fit-for-purpose 14 water production due to its inherent ion-selective recovery properties. A major bottleneck in the optimization of the IX process is the accurate prediction of ion breakthrough times, which has the potential to save on 15 16 regeneration chemicals by maximizing resin utilization. However, the models used to predict ion breakthrough 17 times are often unreliable due to poor calibration methods and significant uncertainty in parameter estimates. 18 Consequently, we conducted local and global sensitivity analyses to identify the design and operational 19 parameters that contribute most to the prediction of breakthrough curves. The global sensitivity analysis 20 enabled the selection of a limited subset of parameters for calibration, demonstrating that only two 21 parameters, namely the maximum adsorption capacity isotherm parameter and the resin bead particle size, 22 require thorough calibration, resulting in a 76% improvement in the breakthrough prediction. We also showed 23 that the calibration of additional, less sensitive or correlated parameters results in an insignificant 24 improvement of the predictive power, with a 16% to 60% increased uncertainty in the breakthrough time 25 prediction. The model was validated using three independent data sets, which showed a fairly accurate 26 breakthrough time prediction, with a relative error ranging from 1% to 11%. Herein, we propose a robust 27 calibration procedure, based on good modeling practice, that encompasses both sensitivity and uncertainty 28 analyses and therefore provides a basis for process optimization. The framework is presented in a manner 29 that allows for its application to analogous process settings.

# 30 Keywords

Adsorption, calibration protocol, fixed-bed column, global sensitivity analysis, mechanistic model,
 uncertainty analysis

# 33 Highlights

- 34 1. Local and global sensitivity analyses identified key parameters for calibration.
- 35 2. Only maximum adsorption capacity and resin bead size require thorough calibration.
- 36 3. Calibrating only two parameters resulted in a 76% improved breakthrough prediction.
- 37 4. Prediction uncertainty increased by 16-60% when calibrating correlated parameters.
- 38 5. Calibration protocol is applicable to any physico-chemical or adsorption processes.

# 39 Abbreviations

40	AIC	Akaike Information Criterion
41	BIC	Bayesian Information Criterion
42	CI	Confidence Interval
43	FMI	Fisher Information Matrix
44	GSA	Global Sensitivity Analysis
45	IX	Ion Exchange
46	LSA	Local Sensitivity Analysis
47	ODE	Ordinary Differential Equation
48	PDE	Partial Differential Equation
49	RMSE	Root Mean Square Error
50	WSSE	Weighted Sum of Square Error

# 51 1. Introduction

52 The global issue of water scarcity is placing increasing pressure on the supply and management of water 53 resources. As a result, resource recovery has emerged as a strategy to increase the circularity of resources 54 and water supplies. Ion exchange (IX) is a key technology in this endeavor, as it enables fit-for-purpose water 55 demineralization through its ion-selective recovery properties (Kabdaşlı and Tünay, 2018; Jegatheesan et al., 56 2021; Taghvaie Nakhjiri et al., 2022). IX, like other adsorption processes, is typically operated as a continuous 57 process in fixed-bed packed columns and used as an end-of-pipe removal treatment (Inglezakis and Zorpas, 58 2012). Notable applications include the recovery of nutrients and by-products in the food production industry 59 (Kammerer et al., 2011); the removal of heavy metals such as nickel from urban wastewater (Ma et al., 2019), 60 petrochemical wastewater (Cechinel et al., 2018) and other industrial waters such as chromium-rich textile waste (Wang et al., 2015); the recovery of precious metals from industrial wastewater (Taghvaie Nakhjiri et al., 61 62 2022); the removal of silica for the production of ultrapure water in various industries, including paper mills 63 and electronics (Chen et al., 2022); or the removal of pharmaceuticals (Chu and Hashim, 2023a) and 64 micropollutants such as PFAS in drinking water production (Smith et al., 2023).

65 The challenges of the IX technology include the optimization of the cost of chemicals associated with resin 66 regeneration and the disposal of generated waste streams (Crittenden et al., 2012), as well as the accurate 67 prediction of target ion breakthrough times for downstream compliance during operation and control (Inglezakis and Zorpas, 2012). Consequently, the optimization of IX processes would allow for a reduction in
 the energetic and material impacts of water treatments, thereby enhancing their environmental and economic
 sustainability and further increasing their potential for resource recovery.

71 A number of mechanistic and empirical models have been developed with the objective of improving the 72 understanding of the IX process and addressing the aforementioned challenges by identifying the 73 breakthrough time. A myriad of these models can be found in the literature, including transport and dispersion 74 mechanisms derived from conservation laws, such as surface diffusion (Ma et al., 2019), pore diffusion (Zhang 75 et al., 2015), and other intraparticle diffusion kinetic models (Wang and Guo, 2022). Similarly, numerous 76 equilibrium isotherm models have been developed beyond the classical Langmuir and Freundlich models. A 77 comprehensive review of the most commonly used models can be found in LeVan and Carta (2008), and more 78 recently in Wang and Guo (2023) and Wang et al. (2024).

79 Therefore, in view of the considerable number of existing models, no unified modeling procedure exists, and 80 as a result, a standardized calibration protocol for adsorption models is still missing. This is evidenced by the 81 numerous existing reports on inconsistencies, mistakes, and misconceptions in the modeling of the IX 82 process (Chu, 2023; Haupert et al., 2021; Hu et al., 2021; Lima et al., 2021; Mudhoo and Pittman, 2023; Tran 83 et al., 2017; Xiao et al., 2018), which result in models with limited reliability and prediction power. Recent 84 studies also emphasize the need for repeated experiments and the reporting of data of complete 85 breakthrough curves as essential practices in IX modeling (Hu et al., 2024). Furthermore, the following five 86 common misconceptions in modeling tasks were identified: (i) overfitting, (ii) use of linearized models, (iii) 87 false sensitivity analyses, performed locally and for single parameters, (iv) absence of uncertainty 88 quantification, and (v) confusion between the goal of sensitivity analyses and uncertainty analyses, as 89 highlighted by Saltelli et al. (2019).

90 A review of the above articles from the literature essentially shows that in IX, models can be effectively used 91 to predict ion breakthrough in a multitude of applications. However, the calibration methodologies employed 92 by various authors exhibit inconsistencies in rigor resulting in overfitting, limited reproducibility and high 93 uncertainty regarding the predictions of these models. Consequently, these models frequently fail when 94 subjected to validation or extrapolation to future time series, due to the considerable uncertainty associated 95 with the calibrated parameters. The limited extrapolation capability, coupled with the complexity and non-96 linear nature of the operation, restricts the utility of these models as a monitoring and control tool for water 97 treatment applications. A standardized model calibration procedure is currently missing.

98 In order to achieve reliable and powerful prediction capabilities and establish good modeling practice for the 99 calibration and reproducibility of IX models, we followed general recommendations by Saltelli et al. (2019) 100 regarding sensitivity and uncertainty analysis, which are valid across disciplines. Similar guidelines have been 101 successfully established for the calibration of diverse processes in the wastewater field (Rieger et al., 2012; 102 Vanrolleghem et al., 2003; Verhaeghe et al., 2024). In a recent study, Chu and Hashim (2023b) employed 103 rigorous model selection techniques to evaluate the performance of competing models with varying numbers 104 of fitting parameters. Nevertheless, essential considerations in the calibration process, such as verification of 105 the model structure, a detailed description and uncertainty quantification of the parameter estimation 106 procedure, and the analysis of the data quality and resulting performance, are still absent in numerous studies. 107 Our work is based on the model proposed by Zhang et al. (2015), which considered an advection-diffusion-108 reaction model for fixed-bed ion exchange columns. This allowed us to address a few of the aforementioned 109 limitations and resulted in the formulation of the proposed framework for model calibration. This framework 110 is based on principles of good modeling practice, which are applicable to other physico-chemical or 111 adsorption processes of analogous systems described by mechanistic models.

112 This work is accompanied by open-access code (<u>https://github.com/UGentBiomath/IX-GMP</u>) and presents a

113 framework for model calibration based on good modeling practice. The work includes a description of a simple

- one-component IX model for fixed-bed operation and employs sensitivity and uncertainty analyses to verify
- the model structure and facilitate rigorous calibration of an identifiable parameter set. The resulting protocol
- is applicable to other IX modeling studies and allows for scenario analysis and optimization of the systemunder study. A description of the implementation of the presented framework can be found in the Appendix.

# 118 2. Materials and Methods

# 119 2.1 Model definition

120 We implemented a dynamic model describing the transport of ions through a fixed-bed IX column reactor. We selected the two one-dimensional partial differential equation (PDE) approach of Zhang et al. (2015) in order 121 to develop our model. The first PDE describes the liquid phase, while the second PDE describes the solid 122 123 diffusion. As the goal of this modeling study is to optimize the process, computational efficiency is a 124 significant factor. Consequently, we simplified the model with the following modifications: the simplified 125 model considers the same one-dimensional set of equations, comprising an advection-dispersion-reaction 126 partial differential equation (PDE) for the liquid phase, but proposes a linear driving force mass transfer 127 ordinary differential equation (ODE) for the solid diffusion, as proposed by LeVan and Carta (2008). Therefore, 128 we replaced the second PDE with an ODE.

129 Both models compute the concentration profile of ions in the liquid and solid phases along the column, 130 thereby predicting the breakthrough of these ions following saturation of the resin. A visual inspection of Figure 131 7 shows that our simplified model using parameters from Zhang et al. (2015) (designated as "uncalibrated" in 132 the figure) does indeed produce the anticipated breakthrough curve of the data presented by Zhang et al. 133 (2015). Therefore, the simplified model can be employed as a fast, accurate tool to improve the efficiency of 134 IX operations. This is made possible by the fast concurrent computation of both scales of the process, namely 135 the ion concentration in the liquid and solid phases, which are respectively referred to as the macro and 136 micro-scale by Zhang et al. (2015). Subsequently, both scales are translated into a single output, namely the 137 ion transport in the fluid along the column. Similarly, the solid phase can be analyzed for the purposes of 138 regeneration, although this is beyond the scope of the present work.

# 139 2.1.1 Model structure

Figure 1 depicts the IX process as a fixed-bed resin-packed column, illustrating the main variables involved in
the dynamic transport of ions along the column and through the liquid and solid phases.





Figure 1. Schematic representation of a typical fixed-bed IX column and description of the main variables of
 the 1-D model with appropriate initial and boundary conditions. The parameters are detailed in Table 1.

The evolution of the concentration profiles of a given ion in the column can be obtained by performing a mass
balance. This results in the PDE (1), which must be solved after the appropriate initial and boundary conditions
have been defined. Furthermore, the solution involves the discretization of the space and time domains:

(1)

148 
$$\frac{\partial C}{\partial t} = D_z \frac{\partial^2 C}{\partial z^2} - u \frac{\partial C}{\partial z} + \frac{\rho_b}{\varepsilon} \frac{\partial q}{\partial t}$$

149

150 In the above expression, C represents the concentration of ions present in the liquid phase at any given point. 151 The variable q denotes the concentration of ions in the solid phase. The term u is the fluid velocity, which is 152 assumed to be constant.  $D_z$  is the axial liquid dispersion coefficient.  $ho_b$  is the bulk bed density, while arepsilon153 denotes the bed porosity. In this context, the independent variables t and z are used to refer to time and 154 column height, respectively. Furthermore, the following assumptions have been made: a uniform velocity 155 profile and intraparticle diffusion in the column; and a controlling equilibrium between the liquid and solid 156 phases (negligible resistance to mass transfer in the boundary layer, as indicated by Zhang et al., 2015). The velocity can be calculated by assuming a constant inlet flow and a constant void bed cross-section in the 157 column, according to the following equation:  $u = \frac{Q}{A\varepsilon}$ . The last term in Equation (1) denotes the change in ion 158 concentration due to adsorption or desorption in the solid phase. Assuming a linear driving force between the 159 160 liquid and solid phases (LeVan and Carta, 2008), the second equation of the model can be expressed as an 161 ordinary differential equation at each discretized point within the spatial domain:

162 
$$\frac{\partial q}{\partial t} = k_{ldf}(q^* - q)$$

163

164 where  $k_{ldf}$  is the mass transfer coefficient and  $q^*$  is the concentration in the resin phase surface in equilibrium with the liquid phase concentration, C, which can be calculated using an equilibrium isotherm 165 166 model. Equations (1) and (2) must be integrated and solved concurrently.

(2)

(3)

(5)

167 The two most frequently utilized equilibrium isotherm models in literature are the Freundlich and Langmuir 168 models (O'Neal and Boyer, 2013). The Freundlich isotherm model (Freundlich, 1907) assumes multilayer 169 adsorption described by a power law:

 $q^* = K_f \cdot C^{n_f} = K_f \cdot C^{1/n}$ 

171

172 where  $K_f$  and  $n_f$  are parameters to be calibrated. The Langmuir model (Langmuir, 1918) assumes monolayer 173 adsorption, with  $q_{max}$  representing the maximum adsorption capacity, and  $K_L$  a second parameter to be 174 calibrated:

 $q^* = q_{max} \frac{K_L C}{1 + K_L C}$ 176 (4)

177 In this study, we used the Langmuir isotherm model due to the greater interpretability of its parameters. 178 Nevertheless, the Freundlich model has also been successfully applied to describe the equilibrium between 179 two phases (Sengupta and Pandit, 2011; O'Neal and Boyer, 2013). It should be noted that other, more 180 complex models exist; however, the inclusion of additional parameters complicates the calibration process 181 and is therefore not considered here.

182 The mass transfer coefficient  $k_{ldf}$  in Equation (2) is a function of the intraparticle diffusion coefficient,  $D_p$ , and 183 the particle radius,  $r_p$ , according to a pore or solid diffusion mechanism (LeVan and Carta, 2008). This can be 184 expressed as follows:

$$k_{ldf} = 15 \frac{D_p}{r_p^2}$$

186

The solid and liquid phases are related through the bed density given by  $\rho_b = \rho_p(1-\varepsilon)$ , where  $\rho_p$  is the resin 187 density and  $\varepsilon$  is the void fraction in the column left by the settled resin, also known as bed porosity. 188

#### 2.1.2 Model parameters and variables 189

190 Table 1 lists the variables considered for the present model structure, classified according to their function: 191 output variable, input variable, or parameter. The latter can be further subdivided into the following categories, 192 according to their function: design parameters, which include column dimensions and the physicochemical 193 properties of the resin; operational parameters, which are derived from the specific column operation and 194 subject to the inherent variability of the process; and equilibrium parameters, which could also be considered 195 operational but are here treated separately due to their importance in the considered model. The last column 196 indicates the source of the values: whether they were measured experimentally, calculated numerically, fixed 197 by the process or equipment specifications related to design decisions and reported in manufacturers' data

- 198 sheets, or obtained from the literature. Parameters for which there is no direct measurement or other means 199 of obtaining a value are estimated from output measurements in the calibration process. For each parameter, 200 the values reported in the literature are listed in Table 2, together with their ranges and sources. Even those 201 parameters which can be assumed to be constant for design reasons (manufacturer's data sheets, operating 202 rules) are considered uncertain for the purposes of the sensitivity analysis and the calibration, in order to 203 provide a comprehensive overview of potential model uncertainties. For instance, manufacturers frequently 204 offer data on the particle size distribution of resins. As a result, the inherent variability of this critical parameter 205 can be significant. Furthermore, only the average value is often considered as a model parameter. Therefore,
- 206 we regard the resin particle size as an uncertain parameter due to the uncertainty in measurement.
- The liquid phase concentration is frequently determined at the inlet ( $C_0$ ) and outlet (C) of the column. Direct measurement of the concentration profile within the column is not feasible without introducing multiple sample extractions, which would affect the total column volume. However, these profiles are predicted by the model. While the measurement of solid-phase concentrations is challenging, they can be calculated through a mass balance by means of Equation (2). The initial solid phase concentration,  $q_0$ , is assumed to be zero for fresh resin. As the original resin capacity is never fully restored, this initial condition could also be regarded as a variable in cases where regeneration is being investigated.
- Table 1. Variables and parameters considered in the simplified IX model. The symbols are in accordance with
   the commonly used nomenclature in the literature. For detailed values, see Table 2.

Name	Symbol	Unit	Туре	Source
Outputs				
Liquid-phase concentration	С	mmol/L		Computed by Eq. (1)
Solid-phase concentration	q	mmol/g		Computed by Eq. (2)
Inputs				
Inlet liquid-phase concentration	$C_0$	mmol/L		Directly measured*
Initial solid-phase concentration	$q_0$	mmol/g		Initial condition in Eq. (2)
Volumetric flow	Q	L/min		Manipulated/fixed*
Parameters				
Bed length	L	m	Design	Directly measured*
Bed diameter	D	m	Design	Column manufacturer*
Resin bead particle size	$r_p$	m	Design	Resin manufacturer*
Resin bead particle density	$\rho_p$	g/L	Design	Resin manufacturer*
Bed porosity (void fraction)	3	_	Operation	Literature*
Axial dispersion coefficient	$D_z$	m²/s	Operation	Literature*
Intraparticle diffusion coefficient	$D_p$	m²/s	Operation	Literature*
Maximum adsorption capacity	$q_{max}$	mmol/g	Equilibrium	Literature*
Langmuir constant	$K_L$	L/mmol	Equilibrium	Literature*

216 \*Determination of these parameters is uncertain and therefore could be considered for calibration.

217 The design parameters include the bed dimensions, which determine the volume available for both resin and

effluent in the column, as well as the particle size and the density of the solid phase (inversely proportional to

the specific volume occupied by the resin). The effective volume available for the liquid fraction is a function

- of the bed porosity, which is in turn determined by the compaction of the resin in the column and the swelling
- induced by the liquid. This parameter has been extensively studied, and a range of variability can be found in
- the literature depending on the packing shape and size (see Table 2). The actual liquid capacity of the bed is
- 223 calculated as follows:  $V = \varepsilon LA = \varepsilon L \frac{\pi}{4} D^2$ , with the height or length (L) of the bed directly measured, as it is

not always the case that the full column length is utilized. In the literature, the bed volume is typically referred to as a measure of the time elapsed during the process, calculated as the treated effluent volume per unit time divided by the effective bed volume. In addition, dispersion coefficients are frequently encountered in the literature in the form of empirical correlation functions of the Reynolds number and bed configuration. The determination of intraparticle coefficients is an experimental process, and they are therefore considered to be uncertain.

Table 2. Values and ranges of variability reported in the literature for the parameters listed in Table 1.

	Reported value	Variability range	Source
Design parameters			
Bed diameter, D	0.1 m	(±20%)	Zhang et al., 2015
Bed length, L	10 <sup>-2</sup> m	(±20%)	Zhang et al., 2015
Operational parameters			
Inlet P concentration, $C_0$	20 mmol/L	(±20%)	Zhang et al., 2015
Flow, Q	3.1 cm/min	(±20%)	Zhang et al., 2015
Porosity, $\varepsilon$	0.37	0.3–0.44 (±20%)	Yoshida et al., 1985
Resin size (diameter), $d_p = 2r_p$	7.5·10⁻⁴ m	3-12·10⁻⁴ m (±60%)	Sengupta and Pandit, 2011
Resin density, $\rho_n$	389 g/L	(±20%)	O'Neal and Boyer, 2013
Intraparticle diffusivity, $D_p$	5.3·10 <sup>-10</sup> m <sup>2</sup> /s*	(±20%)	Sengupta and Pandit, 2011
Maximum capacity, $q_{max}$	0.291 mmol/g*	(±20%)	O'Neal and Boyer, 2013
Langmuir constant, $K_L$	1.18 L/mmol*	(±20%)	O'Neal and Boyer, 2013

231 \* Values for synthetic fresh urine.

### 232 2.1.3 Model implementation and numerical solution

The model equations have been formulated as a set of one-dimensional differential equations, thereby enabling the model to be employed as a fast optimization tool for the IX process. It is assumed that the velocity profiles of the fluid across the column are uniform and that there is no significant existence of wide preferential channels due to the low diameter-to-length ratio (*D/L*) of the column.

In order to predict the one-dimensional spatial variation of pollutant concentration along the IX column, a
discretization of the column length was considered. The solution of the nonlinear system of equations can be
computationally intensive. Therefore, a discretization error analysis was conducted (see Appendix for a
detailed discussion) and an appropriate discretization step of 100 grid points and a time step of 0.1 seconds
was selected to achieve sufficient accuracy without substantial computational effort.

The model was implemented in Python 3 (Van Rossum and Drake, 2009) and makes use of available scientific packages *Numpy* (Harris et al., 2020), *Scipy* (Virtanen et al., 2020), *Pandas* (McKinney, 2010; The pandas development team, 2020), and *Matplotlib* (Hunter, 2007). The code will be made available at <u>https://github.com/UGentBiomath/IX-GMP</u>.

The equations were solved with the *scikit-finite-diff* package (Cellier and Ruyer-Quil, 2019) using the finite difference method and the method of lines for the spatial and temporal discretization of the PDE. Figure 2a illustrates the concentration profiles along the length of the column as a function of time in relation to the inlet concentration,  $C_0$ . The arrow indicates the typical temporal evolution of the profiles resulting from the transport of ions along the column and into the solid phase. The breakthrough curve (Figure 2b) reflects the evolving ion concentration in the effluent at the column outlet, illustrating the gradual depletion of the resin and the breakthrough of the ion, which is adsorbed onto the solid phase until the resin is completely saturated

and the breakthrough of the ion, which is adsorbed onto the solid phase until the resin is completely saturated.



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Figure 2. Concentration profiles (a) and breakthrough curve (b) obtained by solving the model equations (1) to (4) with the appropriate initial and boundary conditions, as specified by the scheme in Figure 1. The breakthrough time (highlighted) is defined as the time for 10% of initial concentration in the outlet.

### 257 2.2 Sensitivity analysis

The objective of the sensitivity analysis is to identify those parameters that exert the greatest influence on model output, with the aim of reducing the variability or uncertainty in the model output by accurately determining their values. By identifying which parameters are sensitive, it is possible to reduce the cost of experiments by focusing efforts on measuring those parameters (Saltelli et al., 2007). This methodology is based on principles applicable to a broad range of disciplines, as suggested by Saltelli et al.(2019).

### 263 2.2.1 Local sensitivity analysis

A local sensitivity analysis (LSA) quantifies which model parameters have a greater influence on the model output(s) for a given value of each parameter (Saltelli et al., 2007). The sensitivity function is defined as the change in the model output resulting from a small change or perturbation in the value of a single parameter. A numerical approximation of the sensitivity function, often referred to as the derivative method, is the firstorder forward finite approximation, which is normalized for the purpose of comparison:

269 
$$S_{\theta_i}^j = \frac{\partial y_j}{\partial \theta_i} \cdot \frac{\theta_i}{y_j} \approx \frac{y_j(\theta_i + \Delta \theta_i) - y_j(\theta_i)}{\Delta \theta_i} \cdot \frac{\theta_i}{y_j}$$

270

where  $y_j$  represents the model output,  $\theta_i$  denotes the parameter subject to variation, and  $\Delta \theta_i$  is the perturbation value. In the present study, a perturbation value of  $10^{-5} \cdot \theta$  was used for all parameters. The sensitivity function is normalized to accommodate disparate scales or magnitudes for both the output and the parameters. It is important to note that the results of a local sensitivity analysis may vary significantly depending on the specific location within the parameter space under investigation. Hence, the analysis may

(6)

be repeated at different locations to obtain further information, or a global sensitivity analysis may be performed (see section 2.2.2). Additionally, the calculation of sensitivity as outlined here can be applied to other mechanistic model descriptions, irrespective of the process, the presence of algebraic or differential equations, or the inclusion of non-linear terms.

The local sensitivity analysis is employed as an initial model check on the parameters' behavior, requiring minimal computational effort and offering insight into the extent to which varying parameters influence the model output. Furthermore, it identifies potential correlations between parameters and operational regions where parameter sensitivity is higher and correlation minimal. These regions are of particular interest for experimental data collection.

### 285 2.2.2 Global sensitivity analysis

A global sensitivity analysis (GSA) aims to quantify the relative importance of parameters in determining the variability of a model output over a wide range of parameter values (Saltelli et al., 2007). In contrast to a local analysis, a systematic exploration of the parameter space can provide a more comprehensive understanding of the overall influence of the different parameters on the model output, even in regions of the parameter space where the model exhibits anomalous behavior. This phenomenon arises when multiple parameters are modified simultaneously. To this end, sensitivity indices can be calculated in order to identify these effects.

First-order indices describe the main effect of parameters on the variance of model outputs. They can be calculated as the variability in model output resulting from the variation of a single parameter in isolation, relative to the total variability attributable to changes in all parameters, expressed as follows:

$$S_i = \frac{V[E(Y|X_i)]}{V(Y)}$$

296

where *V* represents the variability of model output *Y*, *E* denotes the expected value, and  $X_i$  is a specific parameter value. In contrast, total effects comprise the cumulative impact of a parameter on the model output, accounting for both first-order and higher-order effects stemming from interactions or non-linearities between parameters. A disparity between total and first-order effects indicates the existence of higher-order interactions:

(7)

(8)

302 
$$S_{T_i} = 1 - \frac{V[E(Y|X_{\sim i})]}{V(Y)}$$

303

304 The GSA is performed at a specific time point. In the present study, the time at which 10% breakthrough is 305 observed was selected as the most informative for GSA. The sensitivity indices for the 10% concentration 306 breakthrough times with respect to the model parameters listed in Table 2 were calculated by Sobol uniform 307 sampling using the SALib sensitivity analysis library (Herman and Usher, 2017). Furthermore, second-order 308 sensitivity indices were calculated and employed to ascertain the existing correlation between parameters. 309 The Sobol sampling of parameters consisted of 5,632 uniform values, selected within the ranges of variability 310 outlined in Table 2. A 20% variability was considered for all parameters, including  $r_p$ . Subsequently, a Monte 311 Carlo-type of simulation was employed to ascertain the variability in model output derived from the 312 simulation results corresponding to each parameter subset sampling. Parameter variability is propagated 313 through the model. Consequently, output variability is a determining factor of sensitivity and results in an 314 uncertainty in the calculated sensitivity indices. Output variability can be reduced by increasing the number of simulations and scales as  $1/\sqrt{N}$ . The primary disadvantage of GSA is thus its high computational cost, which is a consequence of the substantial number of model evaluations required. One advantage of an LSA over a GSA is that it can facilitate the preliminary screening and reduction of the parameter set prior to a more comprehensive global analysis. However, it is important to note that, in principle, a GSA can be applied to any type of model with an arbitrary number of parameters. Once the most sensitive model parameters have been identified, they can be estimated through the process known as model calibration. In order to achieve this, it is necessary to obtain experimental data.

# 322 2.3 Experimental data for calibration and validation

The experimental data used for the calibration of our model are derived from measurements of breakthrough curves for ion exchange of phosphorus recovery from fresh urine, as documented by O'Neal and Boyer (2015). The data set describes the breakthrough of phosphate ions from synthetic fresh urine as it traverses a fixedbed column containing a specific phosphate-selective HAIX-Fe resin. Further details regarding the column tests can be found in the referred work.

328 The authors acknowledge a limitation in the data set, namely the absence of measurements of sulphate and 329 other ions present in the urine effluent. These ions compete with phosphates in their adsorption onto the 330 resin; however, the analysis of competition and affinity effects is beyond the scope of the present work and 331 not captured by the model. Therefore, the influence of these ions on the predictive power of the model will not 332 be addressed. The required parameters for the modeling of this system are enumerated in Table 2, 333 accompanied by the sources from which they were derived. As illustrated by Figure 7, the predicted 334 breakthrough curve exhibits a high degree of agreement with the measured data. Further details can be found 335 in the Results and Discussion section.

### 336 2.4 Model calibration

The goal of model calibration is to identify the optimal set of parameter values that fit the experimental data, which can be considered as an optimization problem. The quality of the fit is quantified by an objective function, which is then minimized. The weighted sum of squared errors (WSSE) was selected as the objective function for model calibration to represent the distance between model prediction and experimental data:

341 
$$J(\theta) = \sum_{i=1}^{N} (\hat{y}_i(\theta) - y_i)^T W(\hat{y}_i(\theta) - y_i) = WSSE$$

342

where  $\hat{y}_i(\theta)$  represents the model prediction of the output  $y_i$  corresponding to the parameter set of values  $\theta$ ,  $y_i$  denotes the measurement of the output y, W is the square matrix of weights associated with different outputs or time moments, and N is the number of measurements. In this analysis, we assumed W to be the identity matrix, thereby assigning equal weight to each time point of the single measured model output.

(9)

There are numerous techniques for minimizing the objective function, including the well-known Nelder-Mead simplex method (Nelder and Mead, 1965). The Levenberg-Marquardt method (Levenberg, 1944; Marquardt, 1963) was selected as a robust and efficient method well suited for unconstrained nonlinear least squares fitting problems. Further details regarding the implementation can be found in Gavin (2019). A recent overview of calibration methods for computer simulation is provided by Sung and Tuo (2024).

Unconstrained optimization methods may suggest values for the parameters that are outside of their range of
 physical validity. Consequently, we also employed a robust constrained trust region method (Branch et al.,

354 1999) to evaluate the performance of constrained parameter subsets in comparison to that of the 355 unconstrained Levenberg-Marquardt algorithm. Both constrained and unconstrained methods are readily 356 available in the optimization library of the Scipy Python package (Virtanen et al., 2020). For the constrained 357 optimization method, the parameter values were bounded by their ranges of validity. The initial parameter estimates, along with their respective ranges of variability, are presented in Table 2. The relative tolerance, 358 which serves as a stopping criterion for the iterative optimization algorithm, was set to  $10^{-4}$  in order to reduce 359 360 the number of model evaluations. This value was selected following a verification process, during which it was 361 determined that it produced results that were not significantly different from those obtained with lower 362 tolerance. Furthermore, the scale or magnitude of each parameter was provided to the algorithm, thereby 363 reducing the number of iterations. The accuracy of the solution was evaluated by comparing the model output 364 to the experimental data provided by Zhang et al. (2015).

#### 2.5 Uncertainty of the parameter estimation 365

366 As Saltelli et al. (2019) observe, while numerous studies acknowledge the significance of sensitivity analysis 367 for model calibration, uncertainty quantification is frequently absent from calibration procedures, resulting in 368 deficient model assessments and an overestimation of their predictive capacity. The estimated parameters 369 have an associated uncertainty due to the presence of noise in the experimental data set used for calibration, 370 and the degree of uncertainty is dependent on the relative importance of this imperfect information in the model structure. Accordingly, parameter estimates are only meaningful when accompanied by their 371 372 confidence region, within which the actual true value is situated at a specified confidence level (Donckels, 373 2009). The confidence region can be based on an approximation of the contour of the WSSE objective function, 374 as this provides a measure of the fit to the experimental data. In the case of linear models, the contour is exact. 375 However, for nonlinear models, it is common practice to employ a linear approximation of the parameter 376 estimation covariance matrix, providing a lower bound for this region (Marsili-Libelli et al., 2003):

$$\{\theta: WSSE(\theta) \le c \cdot WSSE(\hat{\theta})\}$$

378

A linear approximation results in the Fisher Information Matrix (FIM), whose inverse is employed for the 379 380 approximation of the error covariance matrix, can be computed as follows (Dochain and Vanrolleghem, 2001):

(10)

(12)

381 
$$FIM = \sum_{i=1}^{N} S'_{\theta} \cdot W^{-1} \cdot S_{\theta} = \sum_{i=1}^{N} \left( \frac{\partial \hat{y}}{\partial \theta}(t_i) \right)' \cdot W^{-1} \cdot \left( \frac{\partial \hat{y}}{\partial \theta}(t_i) \right)$$
382 (11)

382

383 where W is the covariance matrix of the measurement errors associated with the measured variables, and  $S_{\theta}$ 384 denotes the parameter sensitivity matrix of all outputs with respect to each parameter, as calculated by 385 Equation (5) for  $N_p$  parameters and N experimental time points. The approximated error covariance matrix 386 can be used to construct a confidence region for the parameter estimates,  $\delta_i$ , with a specified level of 387 confidence 1- $\alpha$  (Marsili-Libelli et al., 2003):

$$\delta_i = t_{N-N_p}^{\alpha/2} \cdot \sqrt{\sigma_{i,i}^2}$$

389

390 where t represents the  $\alpha/2$  quartile of the Student's t distribution for a given confidence level  $\alpha$  and  $N - N_p$ 391 degrees of freedom, with N denoting the number of data points,  $N_p$  the number of estimated parameters, and 392  $\sigma_{i,i}^2$  the variance of parameter *i* taken from the error covariance matrix. The diagonals of the covariance matrix 393 thus provide the variances of the errors in the parameter estimates, whereas the off-diagonal elements are 394 the covariances of the errors and offer a measure of the correlation between the different parameters. The 395 linear correlation between two parameters can be estimated based on the following equation:

$$r_{i,j} = \sqrt{\frac{\sigma_{i,j}}{\sigma_{i,i}^2 \cdot \sigma_{j,j}^2}}$$

397

398 The linear correlation is approximately -1 (negative) or 1 (positive) for pairs of parameters with a high degree of correlation, whereas a value of approximately zero indicates a low correlation. 399

(13)

400 Once the confidence region for the parameter values has been established, the Monte Carlo technique can 401 be employed to sample the parameters within the region and thereby obtain the expected variability in the 402 model output. A normal distribution was assumed for all parameters, and a total of 640 values were sampled 403 for use in the Monte Carlo simulations. The plotting of the most frequent values of the model output can assist 404 in establishing an uncertainty band around the mean value, thereby providing a confidence interval band for 405 the prediction of the breakthrough curve.

#### 2.6 Fitness comparison 406

407 The root mean square error (RMSE) was employed as a metric for assessing the goodness of fit, or the 408 discrepancy between the experimental data and the calculated breakthrough concentrations. The RMSE can 409 be calculated from the WSSE as follows:

410  

$$RMSE = \sqrt{\frac{\sum(y_i - \hat{y}_i)^2}{N}} = \sqrt{\frac{WSSE}{N}}$$
411  
(14)

411

412 The RMSE offers a straightforward and readily understandable representation of the overall model's error, 413 using the same units as the measured variable, even for unitless comparisons, and represents a more 414 accurate average of the distance between the data and the model prediction than the WSSE from the objective 415 function. On the other hand, the RMSE is sensitive to both outliers and overfitting; consequently, it diminishes 416 when additional parameters are incorporated into the model. To further evaluate the fitting quality, methods 417 beyond the use of error statistics, such as residual plots or statistical hypothesis tests for model comparison, 418 can be considered. Residual plots can more reliably display the even distribution of errors centered around 419 zero than error statistics, showing clear trends or biases, while statistical tests allow to compare between models of different complexity (Hu et al., 2024). 420

421 In order to facilitate a comparative analysis of model fitness for different numbers of calibrated parameters, it 422 is possible to consider criteria that balance the goodness of fit to experimental data with the number of model 423 parameters (Wang et al., 2024). This approach allows to penalize overfitting when increasing parameters are 424 incorporated into the model structure. One such established method is Akaike's Information Criterion or AIC 425 (Akaike, 1974):

426 
$$AIC = N \cdot \log\left(\frac{WSSE}{N}\right) + 2 \cdot N_p$$

(17)

In Equation (15), the first term will decrease for overparametrized candidate models due to overfitting, while 428 429 the second term will penalize the added complexity when more parameters are considered. A lower value of 430 AIC is indicative of a superior model, whereas a low WSSE value is indicative of overfitting. In instances where 431 the sample size is less than 40, a corrected form of AIC can be employed (Wang et al., 2024):

432 
$$AIC = N \cdot \log\left(\frac{WSSE}{N}\right) + 2 \cdot N_p + \frac{2N_p(N_p + 1)}{N - N_p - 1}$$
433 (16)

433

427

434 Alternatively, the Bayesian Information Criterion (BIC) imposes a greater penalty on excessive complexity in 435 overparametrized models to a greater extent than AIC when applied to large data sets (Schwarz, 1978):

$$BIC = N \cdot \log\left(\frac{WSSE}{N}\right) + N_p \cdot \log N$$

437

438 Both AIC and BIC can be applied to evaluate and select the most suitable model for a given set of experimental 439 data. The utilization of the corrected AIC form is recommended for data sets of limited size (Wang et al., 2024).

#### 2.7 Model validation 440

441 To confirm the predictive power of the calibrated model, a new data set was selected from the same column 442 breakthrough experiments as the calibration tests (see Section 2.3 for a description of the data). For 443 calibration, the parameter  $q_0$  in our model was assumed to be zero, corresponding to fresh resin. The second 444 cycle was chosen to validate the model and therefore the measured data correspond to a regenerated column. 445 Hence, we assumed a regeneration efficiency of 95% for the resin, which was calculated as the amount of 446 phosphorus recovered from the total adsorbed in the column, and the remaining amount was assumed to be 447 retained or fouled in the resin, as also reported by the authors (O'Neal and Boyer, 2015; Zhang et al., 2015). 448 From the adsorption data, fouling is estimated at 1.7 mg out of the total 31.9 mg adsorbed, resulting in a 5.3% 449 fouled amount or 94.7% regeneration efficiency. We then assumed an initial concentration profile in the resin, 450 parameterized by  $q_0$  in our model.

#### Results and Discussion 451

#### 3.1 Sensitivity analysis 452

#### 3.1.1 Local sensitivity analysis 453

454 The specific set of model parameters to be analyzed is presented in Table 2. The parameter  $q_0$  was omitted 455 from the list, as it is assumed to be equal to zero in this calibration data set. Figure 3(a) shows the variation 456 over time, expressed in bed volumes, of the relative sensitivity values for the breakthrough concentration with 457 respect to all parameters in the selected subset, with the most sensitive parameters highlighted. The less 458 sensitive parameter  $D_p$  is also highlighted as a reference. As can be seen, the sensitivity to changes in 459 parameter values is practically zero at the start of the column operation and increases exponentially when the 460 first breakthrough is detected, around two bed volumes. A peak in sensitivity is reached for all parameters 461 around four to six bed volumes and decreases again exponentially until the end of the operation when the resin is saturated. The occurrence of peaks for several parameters at approximately the same time is a clear indication of a correlation between these parameters. This correlation will make it more challenging to obtain reliable values when these parameters are estimated simultaneously. Since some sensitivities appear to change sign over time, an averaged sensitivity allows to compare the overall trend. Figure 3(b) shows the averaged local sensitivity values over 16 bed volumes for all parameters with their corresponding signs.



467

Figure 3. Local sensitivity analysis (LSA) of the breakthrough curve for the parameters listed in Table 1: time variation in bed volumes for a  $10^{-5}$  relative perturbation (a), and time average with sign (b). The most sensitive parameters ( $q_{max}$ ,  $\varepsilon$ ,  $\rho_p$ ,  $C_0$ ,  $r_p$ ) are highlighted, with  $r_p$  and  $D_p$  negatively correlated.

In descending order of sensitivity, the most sensitive parameters are  $q_{max}$ ,  $\varepsilon$ ,  $\rho_p$ , and  $C_0$ . Most of these parameters exhibit positive or negative sensitivity to varying degrees, indicating that they affect the model output in a consistent manner but with differing magnitudes. However, parameters  $r_p$  and  $D_p$  exert an inverse influence on breakthrough, as evidenced by their inverse correlation, illustrated in Figure 3(a) and in accordance with Equation (5). The particle size is more than twice as sensitive as the intraparticle diffusivity due to the quadratic influence, with the greatest effect observed around four bed volumes, where both parameters are most sensitive.

478 Changes in sensitivities are also indicative of nonlinearity. In order to illustrate the impact of varying 479 sensitivities, it is possible to plot the effect of a change in different parameters on the model output. Figure 4 480 shows the impact of a 10% increase in four of the most sensitive parameters and its effect on the breakthrough 481 concentration. An increase in a parameter with negative sensitivity (e.g.,  $q_{max}$ ) will result in a delay in 482 breakthrough, as it has a negative influence on the concentration. Conversely, a positive sensitivity (e.g., with 483 respect to  $\varepsilon$  or  $C_0$ ) will result in an earlier occurrence of breakthrough for an increase in the parameter value. 484 Figure 4 also confirms that  $C_0$  is a less sensitive parameter than  $\varepsilon$ , indicating that a 10% increase in the latter 485 will have a more pronounced effect on breakthrough. An analogous analysis may be performed for the 486 remaining, less sensitive parameters. As can be observed, a change in the less sensitive parameter  $r_p$  is most 487 pronounced around four bed volumes, where the parameter exhibits the greatest sensitivity to changes, as 488 illustrated in Figure 3. However, the overall effect of this one parameter is minor in comparison to an increase 489 in the maximum adsorption capacity,  $q_{max}$ , as demonstrated by Figure 3: an increase in the bed height 490 (parameter L) would result in greater resin availability too, which in turn delays breakthrough. However, this 491 increase has overall less effect than a comparable change in the maximum adsorption capacity. This serves

492 to illustrate once more the nonlinear nature of the process and the importance of taking into account these493 effects for the estimation of parameters.



494

Figure 4. Effect of a 10% increase in the most sensitive parameters  $(q_{max}, \varepsilon, C_0 \text{ and } r_p)$  on the breakthrough curve: the maximum adsorption capacity  $(q_{max})$  has a marked negative impact on the breakthrough concentration, while the other three parameters have a positive effect, albeit with varying magnitudes.

Following Figure 3 and Figure 4, we can establish a local ranking of the parameter sensitivities for the model structure considered. It is important to note, however, that this is only a local ranking and should therefore be interpreted and used with care. In the case of nonlinear models, parameter sensitivities may vary considerably when evaluated in different regions of the parameter space. Accordingly, a global sensitivity analysis is conducted in the following section to ascertain an overall parameter ranking for the model calibration.

503 The results of the LSA serve two distinct purposes. First and foremost, the LSA results offer insights into the 504 parameter sensitivities as a function of time, thereby enabling the identification of an optimal time point for 505 conducting a GSA analysis. Based on the LSA analysis, the time at which a 10% breakthrough occurs is 506 identified as an informative time instant to perform the GSA. Secondly, the LSA results are employed to 507 conduct an initial screening of the parameters, whereby parameters exhibiting low sensitivity can be excluded 508 from a subsequent analysis. As the GSA analysis is considerably more computationally intensive, a 509 preliminary screening through LSA can significantly accelerate the global analysis. The parameters exhibiting 510 the lowest local sensitivity are  $K_L$ , Q,  $D_p$  and  $D_z$ . As the inlet flow rate Q is a variable that can be manipulated, 511 it was decided that it should be retained for a comprehensive analysis in order to ascertain its influence. 512 Furthermore, the isotherm parameter  $K_L$  is frequently calibrated in conjunction with  $q_{max}$ , and thus it was also 513 determined that it should be included in the GSA. However, in the event of computational constraints, this 514 category of parameters may be excluded from the subsequent analysis, given their negligible impact. It is 515 therefore possible to assign a fixed value within their range of variability to those parameters that have little

influence on the output and produce little variance, without compromising the estimation process for theremaining parameters.

518 A sensitivity analysis may also serve to identify the experimental conditions that will yield the most accurate 519 parameter estimation by maximizing the effect on the model output, which is the goal of optimal experimental 520 design or OED (Donckels, 2009). The objective may be to generate experimental data with high information 521 content, thereby further reducing the uncertainty of the parameter estimates. Nevertheless, the optimization 522 of the experimental design for parameter estimation lies beyond the scope of the study. The LSA analysis 523 conducted in the present study indicates that experiments where the breakthrough concentration surpasses 524 at least 50% of the initial concentration represent the most informative experiments, exhibiting the highest 525 parameter sensitivity.

### 526 3.1.2 Global sensitivity analysis

### 527 Parameter subset selection for GSA

528 Based on the LSA results, parameter  $D_z$  was excluded from the GSA: its quasi-zero sensitivity corresponds to 529 the common plug-flow assumption and the reason why the diffusion term in Equation (1) is often neglected in 530 practice. Figure 5 illustrates the Sobol sensitivity indices for 10% concentration breakthrough times across 531 the ten remaining parameters. Both the total and first-order sensitivity indices are plotted for each parameter, 532 accompanied by 95% confidence bands. The uncertainty in the calculation of the sensitivity indices, along 533 with the variability in the model output, can be reduced by increasing the number of simulations performed. 534 First-order sensitivity indices are analogous to local sensitivities; however, they are calculated over the entire 535 parameter space. As can be observed,  $\varepsilon$  and  $q_{max}$  are the parameters that exert the greatest influence on the 536 breakthrough curve based on both the total and first-order indices. Despite the broader uncertainty bands 537 when compared to the remaining parameters, these bands do not significantly overlap and are sufficiently 538 narrow to render the ranking clear. Consequently, it was determined that conducting additional simulations would not contribute a substantial difference. The maximum adsorption capacity,  $q_{max}$ , has a smaller effect 539 540 than anticipated by LSA when the full range of parameter values is considered. In contrast, the bed porosity, 541  $\varepsilon$ , has overall a more pronounced effect on the breakthrough concentration than anticipated by LSA. This is 542 due to the fact that a slight increase in the value of  $\varepsilon$  results in a reduction of the resin available in the column, 543 which in turn leads to a decrease in the overall adsorption efficiency. The other equilibrium parameter, the 544 Langmuir constant,  $K_L$ , has a negligible effect on the breakthrough concentration across the entire range of 545 values considered. Consequently, the considerably more sensitive equilibrium parameter  $q_{max}$  should be 546 given precedence in model calibration. Lastly, we selected an arbitrary cut-off value of 0.1 for parameters 547 exhibiting minimal global sensitivity. The design parameters, D and L, and the inlet flow, Q, have low indices, 548 indicating that these parameters could be excluded from the calibration process initially. Nevertheless, the 549 resin density and particle size parameters demonstrated a notable degree of local sensitivity, as illustrated in 550 Figure 3. Accordingly, the outcomes of the GSA should always be interpreted with caution.



551

Figure 5. Sobol global sensitivity indices for the parameters listed in Table 2: total-order indices (left), and firstorder indices (right). The analysis was performed for a 10% concentration breakthrough time. A variability of 20% for all parameters was considered. The two most influential or sensitive parameters are  $\varepsilon$  and  $q_{max}$ .

555 Figure 5 additionally illustrates the total-order sensitivity indices for each parameter. The observation that the 556 sum of the total and first-order indices is, respectively, 1.08 and 0.99, with an uncertainty indicated by the 557 95% confidence bands, suggests that this model exhibits some degree of nonlinearity. This is also evidenced 558 by the discrepancy between total-order and first-order effects, which suggests the presence of higher-order 559 interactions or correlations between parameters. The parameters exhibiting the most significant discrepancy 560 between total and first-order indices indicate the highest degree of interaction with other parameters. The 561 SAlib library further allows the calculation of second-order sensitivity indices with supplementary parameter 562 values sampled, thus necessitating additional computational resources and effort. Figure 6 depicts these 563 second-order sensitivity indices, which illustrate the interactions between pairs of parameters. The largest 564 interactions are observed for both isotherm parameters and the intraparticle diffusivity with the bed porosity, 565 the particle size, and the resin density. This indicates that all these parameters have a high degree of 566 correlation and may exert a particularly strong influence on the adsorption process. This strong correlation 567 between parameters was also evident in the single point of the parameter space depicted in Figure 3, with the 568 majority of parameters exhibiting a peak in sensitivity at approximately the same time instant, following 569 breakthrough and preceding saturation. A strong correlation complicates the simultaneous estimation of 570 multiple parameters, since correlated parameters are non-identifiable (Dochain and Vanrolleghem, 2001). 571 Nevertheless, an example of parameter exhibiting comparatively less correlation with the remaining 572 parameters is that of the intraparticle diffusion,  $D_p$ , with the notable exception of  $r_p$  and  $\varepsilon$ , with which it is 573 significantly correlated. Furthermore, the remaining parameters also exhibit some degree of interaction, 574 particularly the inlet concentration,  $C_0$ , although to a lesser extent.



575

# 576 Figure 6. Second-order sensitivity indices. Parameters $\varepsilon$ , $q_{max}$ , $K_L$ , $D_p$ , $r_p$ show the largest interactions.

577

### Parameter subset selection for calibration

578 In light of the results of the GSA presented in Figure 5 and Figure 6, it becomes evident that only a limited 579 subset of parameters should undergo calibration. A subset of parameters for calibration can be selected 580 based on the most sensitive parameters. Consequently, the first parameter to be estimated would be  $\varepsilon$ , as 581 this exerts the greatest influence on the breakthrough concentration. Furthermore, it can be observed that 582  $q_{max}$  exerts a considerable impact on the breakthrough concentration. A calibration subset comprising the 583 two most sensitive parameters could be formed. Both  $\rho_n$  and  $r_n$  have low indices, yet Figure 3 indicates that 584 both parameters exhibit high local sensitivities. Accordingly, both parameters could also be included in a 585 calibration subset. The parameters  $D_p$  and  $K_L$  exert an overall negligible influence on the breakthrough 586 concentration and should therefore not be calibrated initially. Moreover, we assumed that the inlet 587 concentration,  $C_0$ , remains constant in this study. The design parameters L and D can be excluded, given their 588 low sensitivity indices and the fact that their values are frequently fixed for existing columns. With regard to 589 the manipulable variable Q, its value is typically fixed for operational reasons. Therefore, only the parameters 590  $\varepsilon$ ,  $q_{max}$ ,  $\rho_p$  and  $r_p$  will be considered for calibration.

### 591 3.2 Model calibration

In accordance with the findings of the GSA, we conducted a stepwise calibration for the model parameters, beginning with the calibration of the most sensitive parameter and subsequently incorporating an additional parameter into the calibration set. For instance, we initially calibrated  $\varepsilon$ , followed by the simultaneous calibration of  $\varepsilon$  and  $q_{max}$ . This process was then repeated for subsequent parameters. For each calibration exercise, the confidence intervals of the parameters were calculated. The RMSE, AIC (corrected form) and BIC values are provided as fitness criteria. The results are summarized in Table 3. 598 Table 3. Results of the parameter estimation for up to four parameters in different calibration subsets. The 599 values indicate the estimated parameter values, the corresponding 95% confidence intervals (CI, relative 600 percentage calculated with FIM), and RMSE, AIC and BIC as fitness criteria. Initial estimates given by  $\theta_0$ .

$N_p$ a	$N_{f}$ b	Min. °	Е	$q_{max}$	$ ho_p$	$r_p$	RMSE	AIC	BIC
1	9, 10	LM, TR	0.35 ± 1.8%				0.0233	-148	-147
	11, 12	LM, TR		0.323 ± 1.4%			0.0261	-144	-143
	10	TR			463 ± <b>25.8%</b>		0.0300	-136	-134
	9,8	LM, TR				3.4·10 <sup>-4</sup> ± <b>34.8%</b>	0.0857	<b>-96</b>	-95
2	31,38	LM, TR	<b>0.31 ± 30.6%</b>	0.243 ± 22.1%			0.0207	-151	-149
	25	LM	<b>0.30 ± 30.0%</b>		<b>248 ± 42.9%</b>		0.0208	-148	-146
	24	TR	<b>0.32 ± 32.5%</b>		<b>311 ± 48.0%</b>		0.0213	-147	-145
	15	LM, TR	0.35 ± 1.9%			<b>4.1·10</b> <sup>-4</sup> ± 8.1%	0.0207	-151	-149
	21	LM <sup>d</sup>		0.280 ± <b>10</b> ⁵ %	<b>517 ± 10</b> ⁵%		0.0261	-141	-140
	15	TR <sup>d</sup>		0.312 ± <b>10</b> ⁵ %	417 ± <b>10</b> <sup>5</sup> %		0.0261	-138	-137
	15	LM, TR		0.326 ± 1.4%		<b>4.3·10</b> <sup>-4</sup> ± 8.4%	0.0206	-151	-149
3	44	TR <sup>d</sup>	0.31 ± <b>11.7%</b>	0.258 ± <b>10</b> <sup>6</sup> %	344 ± <b>10</b> <sup>6</sup> %		0.0207	-148	-146
	35	LM	0.33 ± <b>10</b> <sup>2</sup> %	0.275 ± <mark>10</mark> ²%		4.0·10 <sup>-4</sup> ± <b>10</b> <sup>2</sup> %	0.0206	-148	-146
	39	TR	0.36 ± <b>10</b> <sup>2</sup> %	0.312 ± <b>10<sup>2</sup>%</b>		4.2·10 <sup>-4</sup> ± 10 <sup>2</sup> %	0.0205	-148	-146
4	71	TR <sup>d</sup>	0.35 ± <b>10<sup>7</sup>%</b>	0.301 ± <b>10<sup>7</sup> %</b>	389 ± <b>10<sup>7</sup>%</b>	4.2·10 <sup>-4</sup> ± <b>10<sup>7</sup>%</b>	0.0207	-145	-143
	35	TR <sup>d</sup>	0.35 <b>± 10<sup>7</sup>%</b>	0.301 ± <b>10<sup>7</sup> %</b>	466 ± <b>10<sup>7</sup>%</b>	4.2·10 <sup>-4</sup> ± <b>10<sup>7</sup>%</b>	0.0207	-145	-143
$\theta_0$			0.37	0.291	389	3.8·10 <sup>-4</sup>	0.0864	-	-
		LM	-	-	_	-			
		TR	0.30-0.44 (±20%)	0.233-0.349 (±20%)	311-467 (±20%)	(3.0-4.5)·10 <sup>-4</sup> (±20%)			

<sup>a</sup> Number of model parameters in the calibration subset.

<sup>b</sup> Number of respective function evaluations of the minimization algorithm.

<sup>603</sup> <sup>°</sup> Minimization algorithm. LM: Levenberg-Marquardt (unconstrained); TR: trust region (constrained).

<sup>d</sup> The broad CIs are attributable to the inability to estimate the covariance matrix (FIM matrix is singular).

605 The calibration of only the most sensitive parameter, namely  $\varepsilon$ , resulted in a 73% improvement in fit (RMSE of 606 0.0233 vs 0.0864) compared to the initial uncalibrated value of 0.37 given by  $\theta_0$  at the bottom of Table 3. The 607 95% confidence interval (CI) was estimated at ±1.8%, expressed as a relative percentage of the optimal value. 608 After approximately 10 to 20 model evaluations, both the Levenberg-Marquardt method (unconstrained 609 parameter values) and the trust reflective region method (parameter values are constrained to the bounds 610 indicated at the bottom of Table 3) yielded identical results. In comparison, the calibration of the following 611 less sensitive parameter,  $q_{max}$ , yielded a confidence interval of ±1.4%, which can be attributed to the higher 612 local sensitivity compared to  $\varepsilon$ . Moreover, the determined value of 0.323 is more closely aligned with the total 613 phosphate loading on the resin of 10.2 mg P/g or 0.329 mmol P/g, as reported by O'Neal and Boyer (2015). The 614 remaining, less sensitive parameters  $\rho_p$  and  $r_p$  yielded a comparatively inferior fit when calibrated separately, 615 as evidenced by the increased values for the confidence intervals and RMSE, AIC and BIC fitness criteria. This 616 observation suggests that models with poor calibration, characterized by uncertain or unidentifiable 617 parameters, may not exhibit significantly superior performance in comparison to those that are uncalibrated. 618 This phenomenon can also be attributed to the potential for overfitting, which can lead to a failure to capture 619 process dynamics, as evidenced by research on the training of a hybrid model, a combination of mechanistic 620 and data-driven models, where the uncalibrated mechanistic models outperformed calibrated models 621 (Verhaeghe et al., 2024). The presented framework offers a methodology to circumvent such cases.

622 Subsequently, a second parameter was incorporated into the calibration set. This allowed for the 623 simultaneous estimation of both parameters, beginning with their initial values as presented at the bottom of Table 3. The estimation of both  $\varepsilon$  and  $q_{max}$  resulted in a further reduction in the root mean square error (RMSE) 624 625 to 0.0207, representing an 11% decrease. However, both minimization methods yielded considerably low 626 values in comparison to those reported in the literature, and the quality of the estimation also declined, as 627 evidenced by the increased confidence intervals. This can be attributed to the significant correlation between 628 both parameters, which was calculated to be as high as 99.8%, as illustrated in Figure 3. Moreover, estimating 629 both  $\varepsilon$  and  $\rho_p$  did not yield enhanced results, as this last parameter is considerably less sensitive than  $q_{max}$ . 630 This was demonstrated in both Figure 3 and Figure 5, and is further corroborated by the calculated correlation 631 coefficient, which reached a value of 99.8%. However, when estimating both  $\varepsilon$  and  $r_p$  simultaneously, despite 632 the calculated correlation of 44.5%, the estimated value and uncertainty for  $\varepsilon$  remained largely unchanged. 633 Furthermore, the calculated 95% confidence interval for this second parameter was 8.1%, indicating that both 634 parameters can be estimated with a high degree of confidence. In comparison, the simultaneous calibration 635 of both  $q_{max}$  and  $r_p$  resulted in an uncertainty of 1.4% and 8.4%, respectively. An alternative calibration subset 636 comprising parameters  $q_{max}$  and  $\rho_p$  yielded unreliable estimates, likely due to their correlation.

637 As illustrated in Table 3, the calibration of additional parameters results in considerably larger confidence 638 intervals for all parameters, consequently leading to a notable increase in the uncertainty associated with the 639 estimated values. Such overparameterization does not result in a significant increase in the fit to the 640 experimental data, as the introduction of additional, less sensitive parameters necessitates substantially 641 larger alterations in their values to produce a change in the objective function. Moreover, this also dramatically 642 increases the number of required iterations for convergence of the minimization algorithm. As a result, the 643 estimation is rendered ill-conditioned. For instance, this is substantiated by the relatively higher (less 644 negative) values of AIC of -148 and -147, with the lowest value of -151 corresponding to the calibration of only 645 two parameters. The BIC criterion confirms these results. These findings are in line with values for RMSE and 646 AIC reported in recent modeling studies for breakthrough prediction (Hu et al., 2024). The simultaneous 647 calibration of three or four parameters results in a further deterioration in the precision of the estimation, as 648 evidenced by the considerably broader confidence intervals for all parameters, including those that were 649 previously estimated with a high degree of accuracy (11.7% for  $\varepsilon$  vs 1.8% when it is estimated together with 650  $q_{max}$  and  $\rho_p$ ). This phenomenon occurs when the selected minimization method yields disparate solutions 651 within the parameter space, with less sensitive parameters estimated at the expense of the certainty in the 652 value of the most sensitive ones. Furthermore, the estimation of less sensitive parameters provides only 653 minimal information, resulting in unreliable estimates due to a poor approximation of the covariance matrix 654 (Vugrin et al., 2007). Consequently, the sensitivity ranking from Figure 5 provides a rough indication of 655 calibration order for the most sensitive parameters. However, correlation and nonlinear interactions between 656 parameters should be taken into account to decide the final subset. This is further substantiated by the data 657 presented in Table 3, which displays the size of the confidence regions as a measure of uncertainty. Finally, 658 we note that this exhaustive analysis can be applied to models of any process or system with multiple 659 parameters.

Figure 7(a) illustrates the fit of the model predictions to the experimental data of Zhang et al. (2015), showcasing both calibrated and uncalibrated parameter values. The calibrated values for parameter  $q_{max}$ from Table 3 are slightly higher than the initial reported values in the literature, resulting in a rightward shift of the breakthrough curve with respect to the uncalibrated model. Figure 7(b) depicts the residuals, defined as the absolute difference between the experimental data and both the calibrated and uncalibrated model predictions. As can be observed, calibration essentially reduces the residuals in the zone where the calibrated parameters are most sensitive, namely after breakthrough, around 4-10 bed volumes. This is the case for all 667 parameter subsets, although beyond two calibrated parameters, the residuals are almost indistinguishable. 668 The calibration of additional parameters does not significantly improve the fit to the experimental data, due to 669 the phenomenon of overfitting or overparameterization, which underlines the usefulness of a screening 670 procedure to select the best parameter subset for calibration and increases the reliability of the estimation. 671 Furthermore, the uncalibrated model displays a tendency to overpredict the breakthrough concentration, 672 which is addressed in the calibration. However, all instances exhibit a comparable limitation in accurately 673 reproducing the initial time instants of the experimental data. This is a consequence of the low sensitivity of 674 the parameters in this zone, which renders the model less powerful in its ability to capture this aspect of the 675 curve for the utilized data. As a result, this limitation of the model structure, revealed by the local sensitivity 676 analysis, is attributable to the choice of model rather than to the calibration procedure itself. Further 677 improvements in the context of different model structures can be achieved by employing model selection 678 techniques.



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682 The optimal model, according to the ranking determined by the lowest value of both the Akaike Information 683 Criterion (AIC) and the Bayesian Information Criterion (BIC), as displayed in Table 3, is obtained by calibrating 684 only two of the most sensitive parameters. These may be, for example, the bed porosity,  $\varepsilon$ , and the particle 685 size,  $r_p$ , or alternatively, the maximum adsorption capacity isotherm parameter,  $q_{max}$ , and the particle size,  $r_p$ , 686 since  $\varepsilon$  and  $q_{max}$  are correlated. This example illustrates the importance of exercising caution when 687 estimating parameters with low sensitivity, as for subsets of more than two parameters, there exist infinite 688 combinations of parameters that produce the same fit, which significantly reduces the predictive capacity of 689 the model. Hence, these results confirm that only the most sensitive model parameters should be included 690 in the calibration process. In practice, many authors perform calibration without a previous analysis of the 691 model structure or provide parameter values without uncertainty estimates. Therefore, a robust calibration

692 protocol, where sensitivity and correlation of parameters are evaluated prior to calibration, is essential to693 ensure the development of a reliable model with minimal uncertainty and optimal predictive power.

### 694 3.3 Uncertainty analysis

695 In addition to the quality of the parameter estimation provided by the confidence regions from Table 3, an 696 uncertainty analysis of the model with respectively two and three calibrated parameters is presented in Figure 697 8. The calibrated model output of the breakthrough curve is enclosed by a 95% confidence band. As illustrated, the uncertainty in the prediction is minimal at the initial and final stages of the operation but increases after 698 699 breakthrough at approximately 2-4 bed volumes and subsequently decreases near the saturation point at 700 around 10-12 bed volumes. Consequently, the width of the uncertainty band is dependent upon the quality of 701 the model calibration step and thus determines the reliability of the model prediction. For two calibrated 702 parameters ( $q_{max}$  and  $r_p$ ,  $N_p = 2$ ), the uncertainty band is narrow and closely surrounds the calibration data 703 set, indicating a highly accurate prediction. However, while the initial instants of the process were not 704 accurately captured by the model, the prediction of breakthrough closely follows the calibration data. Further reasoning was provided in Section 3.2. In the case of calibrated parameters  $q_{max}$  and  $\varepsilon$ , the uncertainty band 705 706 becomes between 16% and 60% broader at 10% breakthrough concentration due to the higher uncertainty 707 resulting from the correlation of these two parameters, which is propagated to the model output. For three 708 calibrated parameters ( $N_p = 3$ , as illustrated in Figure 8), the uncertainty band surrounding the breakthrough 709 prediction is markedly broader, thereby demonstrating the inferior calibration. The lowest uncertainty is thus 710 obtained with the calibration of parameters  $q_{max}$  and  $r_p$ . This finding indicates that conducting sensitivity and 711 uncertainty analyses concurrently enhances the reliability on the predictive power of the model, and 712 integration of both analyses is advantageous for the modeling task.



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Figure 8. Uncertainty analysis of the breakthrough curve for a 95% confidence in the prediction interval. The uncertainty bands for two and three calibrated parameters are generated by the corresponding confidence intervals indicated in Table 3. The lowest uncertainty is obtained with two parameters ( $q_{max}$  and  $r_p$ ).

# 717 3.4 Model validation

Once the model has been calibrated with the accurate determination of the most sensitive parameters, the predictive power of the model can be tested against a validation data set. Figure 9 depicts the predicted breakthrough curve of an independent data set for the IX treatment of fresh urine, with a 6% higher influent concentration (713 mg/L). This is compared to the experimental data reported by O'Neal and Boyer (2015).



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Figure 9. Model validation on an independent data set from O'Neal and Boyer (2015). The inset illustrates that
 the initial time period corresponding to 10% breakthrough is accurately captured by the calibrated model.

725 As demonstrated in the inset, the initial time period of the experiment is accurately represented by the model, 726 providing a precise prediction (with an error of 1.4%) of the 10% breakthrough time, which occurs at 727 approximately 4.1 bed volumes (see Table 4). For the calibration set, the predicted 10% breakthrough time 728 exhibited strong agreement with the measured value of 3.8 bed volumes, with an error of 7.8%. To further 729 validate the calibrated model, a second and third cycle of operation were used to predict breakthrough 730 concentration after regeneration of the resin. An approximation of the initial concentration profiles in the resin 731 was calculated based on the fouling information reported by Zhang et al. (2015). The comparison between the 732 data and the model prediction is shown in Figure 10. The predicted breakthrough and saturation times are 733 summarized in Table 4.

Table 4. Breakthrough and saturation times (in bed volumes) from the model prediction and experimental data.
The first cycle corresponds to the calibration data set, the second and third cycle to the validation data set.
The last line corresponds to an independent data set with 6% higher influent concentration.

	Breakthrough time, $t_{10}$			Saturation time, $t_{90}$		
	Data	Model	Rel. error	Data	Model	Rel. error
First cycle (calibration, $C_0 = 672 \text{ mg/L}$ )	3.8	4.1	7.8%	8.8	9.0	2.1%
Second cycle (validation, 6% fouled)	3.2	3.2	1.0%	10.3	8.1	21.5%
Third cycle (validation, 54% fouled)	2.4	2.2	6.7%	9.3	7.2	22.6%
Higher inflow (validation, $C_0 = 713$ mg/L)	4.0	4.1	1.4%	9.3	8.9	5.4%

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Figure 10. Model validation for the second and third operation cycles following resin regeneration as reported
by Zhang et al. (2015). The calibration (first cycle) from Figure 7 is shown as reference. The 10% breakthrough
times for the second and third cycles were accurately predicted, while saturation times were overpredicted.

742 Figure 10(a) illustrates the breakthrough curve of a second cycle for the same influent with a 6% fouled resin. 743 As expected, the breakthrough time decreases as a consequence of resin fouling. Moreover, the saturation 744 time, defined as the time required for a breakthrough concentration of 90% of the initial concentration, also 745 decreases with the accumulation of fouling. These trends were also observed in the prediction of the third 746 cycle, depicted in Figure 10(b), with good agreement between prediction and experimental data. However, the 747 discrepancy between the predicted and actual values at the saturation point is more pronounced in 748 comparison to the breakthrough times. The potential causes of this discrepancy include interactions between 749 the adsorbed ions onto the resin, as postulated by O'Neal and Boyer (2015), which could result in an increased 750 adsorption capacity when the resin is close to saturation. This may, in turn, result in a delayed breakthrough 751 of the ions. It seems plausible that this trend may persist following additional regeneration cycles. However, 752 further analysis is necessary to substantiate this hypothesis, which is beyond the scope of the present study. 753 In addition, a more comprehensive model for resin regeneration is required to further optimize the process.

# 754 4. Conclusions

We present a framework for calibrating a dynamic model, such as for ion exchange (IX) fixed-bed column
operation, based on good modeling practice that can be used as a reference for future modeling studies and
practical model implementations.

- The local and global sensitivity analyses allowed us to identify the design and operational parameters that contribute most to the prediction of breakthrough curves. Specifically, the local sensitivity analysis (LSA) revealed which time intervals during IX operation provide the most information for model calibration, thus allowing the selection of the most informative time instances for further analysis as well as enabling an initial screening to identify highly sensitive parameters at low computational cost. Subsequently, the global sensitivity analysis (GSA) allowed us to select a limited subset of parameters for calibration.
- Our approach showed that calibrating multiple parameters is not invariably optimal. Only two parameters, namely the maximum adsorption capacity isotherm parameter and the resin particle size, require comprehensive calibration to achieve an accurate prediction of the breakthrough curve. Moreover, our findings demonstrated that the inclusion of additional, less sensitive and correlated parameters results in a reduction in the reliability of the parameter estimates, since the parameters become less identifiable (as illustrated by the AIC and BIC model selection criteria).
- We demonstrated the effect of parameter estimation uncertainty on the model output by propagating
   the parameter uncertainty through the model, which showed that the inclusion of unsensitive or highly
   correlated parameters in the calibration significantly increased the uncertainty of the prediction.

The model was validated using three different breakthrough experiments. In light of these findings, we proposed a robust calibration procedure, based on good modeling practice, that encompasses both sensitivity and uncertainty analyses, and therefore provides a basis for process optimization. We applied our calibration procedure to the IX process with the aim of improving the accuracy of breakthrough prediction. The framework is presented in a manner that allows for its application to analogous process settings.

# 779 Acknowledgements

We would like to express our gratitude to Benjamin Claessens for his input on the model development at the
 start of the project, and to Ivaylo Hitsov Plamenov for sharing his expertise on the ion exchange process.

# 782 Funding

This work was supported by Ghent University. MYS was supported by the Swiss National Science Foundation(P500PT\_211132).

# 785 Appendix



### 786 Discretization analysis with respect to numerical parameters



### 795 Description of the implementation of sensitivity and uncertainty analyses

- Algorithm 1. Detailed description of the implementation process for the calibration framework.
- 797 Inputs: parameter values, initial and boundary conditions, discretization steps.
- 798 Outputs: breakthrough concentration, C; breakthrough time,  $10\% C_0$ ; saturation time,  $90\% C_0$ .
- 799 Main steps:

800	1.	Model	definition
801		a.	Model structure, including the number of equations (algebraic, ODEs, PDEs) to solve.
802		b.	Variables and model parameters. Uncertain parameters are candidates for calibration.
803		с.	Numerical solution of the model, with appropriate initial conditions and boundary conditions,
804			and selected time and space discretization steps.
805	2.	Local s	ensitivity analysis
806		a.	Selection of parameters and output variable for analysis, with perturbation value $\Delta  heta_i$ .
807		b.	Calculation of the sensitivity function and average sensitivity $S_i$ with Eq. (6).
808	3.	Global	sensitivity analysis
809		a.	Selection of parameter subset according to LSA. Selection of output variable for analysis.
810		b.	Parameter sampling (Sobol) according to specific ranges (percentage or observed values).
811		с.	Monte Carlo simulation with $N$ runs for each set of sampled parameters from the subset.
812		d.	Calculation of first-order and total-order Sobol sensitivity indices according to Eq. (7) and (8).

813 e. (Optional) Calculation of second-order Sobol indices for non-linear interactions. 814 4. Model calibration 815 a. Selection of parameters to calibrate from sensitivity analysis (reduced calibration subset). 816 b. Selection of optimization algorithm, stopping criteria (tolerance, max. no. of function evaluations) and objective function, Eq. (9). 817 818 c. Estimation of selected parameters. 819 d. Calculation of Fisher Information Matrix (FIM) from sensitivity function according to Eq. (11). 820 e. Calculation of confidence intervals from covariance matrix, for confidence level  $\alpha$ , Eq. (12). 821 Calculation of correlation between pairs of parameters from covariance matrix, Eq. (13). f. 822 5. Uncertainty analysis 823 a. Sampling (assumed normal) according to confidence intervals of  $N_p$  calibrated parameters. 824 b. Monte Carlo simulation with N runs for the  $N_p$  sampled calibrated parameters. 825 c. Calculation of uncertainty band for 95% percentile of model output. 826 6. Model validation

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