Coupled Lattice Boltzmann Modeling Framework for Pore-Scale Fluid Flow and Reactive Transport

Siyan Liu, Reza Barati,* Chi Zhang, and Mohammad Kazemi

ABSTRACT: In this paper, we propose a modeling framework for pore-scale fluid flow and reactive transport based on a coupled lattice Boltzmann model (LBM). We develop a modeling interface to integrate the LBM modeling code parallel lattice Boltzmann solver and the PHREEQC reaction solver using multiple flow and reaction cell mapping schemes. The major advantage of the proposed workflow is the high modeling flexibility obtained by coupling the geochemical model with the LBM fluid flow model. Consequently, the model is capable of executing one or more complex reactions within desired cells while preserving the high data communication efficiency between the two codes. Meanwhile, the developed mapping mechanism enables the flow, diffusion, and reactions in complex pore-scale geometries. We validate the coupled code in a series of benchmark numerical experiments, including 2D single-phase Poiseuille flow and diffusion, 2D reactive transport with calcite dissolution, as well as surface complexation reactions. The simulation results show good agreement with analytical solutions, experimental data, and multiple other simulation codes. In addition, we design an AI-based optimization workflow and implement it on the surface complexation model to enable increased capacity of the coupled modeling framework. Compared to the manual tuning results proposed in the literature, our workflow demonstrates fast and reliable model optimization results without incorporating pre-existing domain knowledge.

1. INTRODUCTION

Pore-scale numerical modeling for flow and reactive transport is crucial to understand the underlying mechanisms of physical-chemical interactions of numerous subsurface natural processes. Advanced waterflooding enhanced oil recovery (EOR) and CO₂ injection and sequestration are two typical subsurface reactive transport-associated scientific and engineering problems that involve complex physio-chemical processes. Meanwhile, they are by nature highly uncertain processes due to the high complexity level of geological settings, which brings more challenges to conducting high-resolution mechanistic modeling studies compared to the reactive transport problems that happen in an artificial structure or domain such as reaction beds or combustion engine cylinders. Take advanced waterflooding or low-salinity waterflooding EOR modeling as an example, multiple underlying mechanisms are potentially working together and affect the rock/mineral surface wettability, and hence improve the recovery, but the actual underlying mechanisms are not fully understood and still debatable. Meanwhile, the modeling approaches to simulate this process have simplified the problems without incorporating all physio-chemical processes in the pore-scale high-resolution fashion. To investigate the reactive transport and the relevant multiphysics phenomenon associated with the abovementioned scientific and engineering problems, a modeling framework capable of handling high-resolution pore-scale single-/multi-phase fluid flow with geochemical reactions and other known or potential new physics is crucial for achieving the desired research goals. The base of this type of modeling framework is the fluid flow model, which preferably can handle complex flow channel geometries, run in parallel CPUs/GPUs for performance, and incorporate other physio-chemical models.

Besides the conventional computational fluid dynamics (CFD) methods such as finite difference method, finite volume method (FVM), and finite element method (FEM), lattice Boltzmann method (LBM) is a unique modeling framework that is constructed with particle collision and streaming in regular lattice grids and solves macroscopic equations for the mesoscopic fluid flow kinetics. LBM is proved to be a flexible and practical framework for modeling pore-scale fluid flow and
multiphysics behaviors.\textsuperscript{11–13} In addition to the single-phase fluid flow model, multicomponent multiphase LBM models are attributed to another major component of the LBM family as the equivalent of other multiphase CFD methods, for instance, the level set method,\textsuperscript{15} front-tracking method,\textsuperscript{15} and volume-of-fluid method.\textsuperscript{16} The most popular multiphase LBM models used in this community are the Shan–Chen model,\textsuperscript{17,20} the color-gradient model,\textsuperscript{18} the free-energy model,\textsuperscript{19,20} and the He–Chen–Zhang model.\textsuperscript{21}

Regarding the pore-scale advection–diffusion and reactions, the LBM framework is flexible to adapt the conventional advection–diffusion equations and place the species concentration in scalar fields coupled with a fluid flow lattice.\textsuperscript{22} Various studies have focused on the LBM reactive transport modeling, for example, LBM chemical transport,\textsuperscript{23–25} reactive transport with mineral reactions,\textsuperscript{26} mineral dissolution and dissolution-induced rock mechanical property changes,\textsuperscript{27,28} CO\textsubscript{2} reactive transport,\textsuperscript{29} reactions between immiscible fluids,\textsuperscript{30} and convection and heat transfer-associated problems.\textsuperscript{31,32} More studies for the LBM multicomponent reactive transport models can be found in the review by Yoon et al.\textsuperscript{33}

On the other hand, to enhance the flexibility and capabilities of LBM models in solving complex fluid flow and reactive transport problems, numerous studies have focused on coupling other or external models to different applications. For instance, the proposed applications include but are not limited to the coupled models that incorporate FVM with LBM;\textsuperscript{33–35} multiscale coupling for heat, mass transfer, and reactions using a general reconstruction operator;\textsuperscript{30} mass transport,\textsuperscript{35} mineral dissolution and phase transition-associated multiphase reactive transport,\textsuperscript{36–38} and hydrodynamic property evolution with mineral dissolution.\textsuperscript{39,40}

Among those external solvers, PHREEQC\textsuperscript{41} developed by USGS and its variants IPHREEQ,\textsuperscript{42} and PHREEQC/CRM\textsuperscript{43} have been used extensively for reactive transport coupling due to their strong capabilities for solving complex reactions.\textsuperscript{44} IPHREEQ utilizes the Microsoft component object model (COM) interface to communicate with the outside code. Numerous studies have used IPHREEQ for reactive transport modeling, including LBM-based models.\textsuperscript{45–47} However, the data communication efficiency is limited for large-scale simulations. PHREEQC/CRM works as a C++ library at a lower level with a significantly improved data exchange speed, which is suitable for reactive transport coupling. There are a few notable applications coupled with PHREEQC/CRM used in reactive transport modeling.\textsuperscript{48–50}

In recent years, with fast-paced progress in data analytics, artificial intelligence (AI), and machine learning (ML), AI has been extensively implemented in solving various scientific and engineering problems. For subsurface pore-scale modeling specifically, AI/ML models are used for different applications, for example, microstructure synthesis and pore space reconstruction using generative adversarial network-based models,\textsuperscript{51,52} image segmentation for rock and flow,\textsuperscript{53–55} petrophysical property and other physical property estimation,\textsuperscript{56–58} prediction of fluid flow,\textsuperscript{59–61} and effective reaction rates.\textsuperscript{62} More applications of ML and geoscience-associated studies can be found in the thorough review studies by Tahmasebi et al.\textsuperscript{63} and Wang et al.\textsuperscript{64} According to their comprehensive review and many other studies, advanced AI/ML models have shown great capabilities in analyzing big data, complex pattern recognition and extraction, and knowledge discovery for physical science problems. AI/ML models are known for building surrogate models for property estimation. Meanwhile, the AI/ML models also demonstrate strong capabilities in assisting optimization of the numerical models.

However, the existing literature in pore-scale reactive transport usually suggests that, in the presence of multiple physical-chemical processes in porous media, the current models lack flexibility or extensibility to solve such complex conditions. For instance, in the advanced waterflooding EOR process, multiple reactions, including but not limited to equilibrium, kinetic, and surface complexation (SCM) reactions, happened simultaneously, which causes dynamic changing fluid–fluid, fluid–solid interaction, and electrokinetics on the rock surface. To comprehensively study and model such complex processes, it is critical to develop an effective framework to properly incorporate the aforementioned factors in the modeling process. In addition, the developed model should be able to scale up and go beyond the demonstration purposes in a parallel computing scheme to solve real-world problems. Furthermore, numerical fluid flow and reactive transport models usually involve tedious and time-consuming parameter tuning, which is computationally expensive. A good integrated model/parameter optimization tool that does not require too much human intervention would be greatly beneficial for obtaining optimum model and model parameters.

In this work, to address the issues mentioned above, a coupled modeling framework based on LBM fluid flow and advection–diffusion models using the PALABOS package is proposed.\textsuperscript{83} The framework incorporates the geochemical reaction solver PHREEQC to handle complex reactions within the porous medium. To gain access to the lower level of the reaction solver for better computational efficiency, the PHREEQC variant PHREEQCRM\textsuperscript{84} is selected in the coupling model, and it works as a C++ library that can be compiled together with the LBM flow solver. A coupling interface is developed to connect the two solvers by adding other capabilities, for example, the cell mapping system that handles different physical-chemical property fields within the flow domain. The framework enables a flexible setup for various chemical reactions both in the aqueous phase and on a solid surface and utilizes the unique capabilities of the two codes as much as possible. In addition, an AI-assisted automatic model optimization workflow is developed that integrates and works with the coupled numerical model. The optimization method is applied to the SCM adjustment for investigating low salinity waterflooding mechanisms. In this paper, Section 2 (Methodologies) includes the detailed model descriptions and coupling mechanisms and the AI-assisted optimization workflow. Four model validation cases in Section 3 (Model Validation and Results) are presented, including simple fluid flow, diffusion validation against the analytical solutions, reactive transport benchmark with calcite dissolution, and AI-optimized SCM results. Section 4 provides conclusions of the work and future research and development directions.

2. METHODOLOGY

2.1. Overview of the Proposed Modeling Workflow. To investigate the pore-scale fluid flow and physiochemical processes effectively and comprehensively, a numerical modeling framework that couples fluid flow models (LBM-based) and geochemical reaction solvers (PHREEQCRM) is proposed in this study. A tailor-made coupling interface connects the two major components for data communications.
The major development goal of the coupled modeling framework is to efficiently and effectively study various pore-scale reactive transport-associated scientific problems, while preserving the flexibility of the future extension in terms of multiphase flow and new add-on physical or chemical models. Figure 1 illustrates the overall workflow of the modeling framework.

2.2. LBM Fluid Flow Model. 2.2.1. LBM Fundamentals. LBM represents a generic CFD modeling framework. LBM is not only capable of modeling single-phase and multiphase fluid flow but also of building complex flow-based multiphysics systems such as the advection–diffusion–reaction system and complex fluid—fluid and fluid—solid interaction scenarios with high parallel computational efficiency.

One of the most widely used LBM models was proposed by Bhatnagar, Gross, and Krook (BGK) with the approximation of the non-linear collisional dynamics by introducing the collision operator as

$$\Omega_k = -\frac{1}{\tau} (f_i - f^{\text{EQ}}_i)$$

where $\tau$ stands for the relaxation time used to reach the local equilibrium and $f^{\text{EQ}}_i$ represents the particle distribution function at equilibrium. By assuming the existence of a local attractor and considering its impact on the collision process, the BGK model linearizes the collision term to reach local equilibrium using a fixed relaxation time and fluid viscosity. The BGK model utilizes single relaxation time $\tau$ in the evolution function (eq 9). Future research and model development will include the two-relaxation time or multi-relaxation time models for improved model stability purpose.

Fluid flow macroscopic quantities are calculated in the BGK model based on the instantaneous status of particle distribution function within a specific grid block. For example, in a D2Q9 model (2D and 9 velocities), the lattice density $\rho_{LB}(x,t)$ is obtained by summing the particle distribution function on all points within the given grid block

$$\rho_{LB}(x,t) = \sum_{i=0}^{8} f_i(x,t)$$

where $f_i(x,t)$ represents the particle distribution function for velocity $i$. Meanwhile, the lattice pressure $P_{LB}$ is recovered by

$$P_{LB}(x,t) = C_s^2 \rho_{LB}(x,t)$$

where $C_s$ is a constant value for the lattice speed of sound, which varies for different models. For instance, in the D2Q9 model, as mentioned, $C_s = 1/\sqrt{3}$, and the lattice velocity vector $u_{LB}(x,t)$ is obtained by

$$u_{LB}(x,t) = \frac{1}{\rho_{LB}(x,t)} \sum_{i=0}^{8} f_i(x,t) \varepsilon_i$$

where $\varepsilon_i$ represents the relative locations of the points within the specific grid block pointing to the adjacent grids, whose values are 0, 1, or $-1$; $i$ is the index for the points within the grid block for the given model; $x$ stands for the location of the grid block in the 2D or 3D simulation domain; and $t$ describes the time step in the simulation for time-dependent particle distribution function evolution.

The particle distribution function for the equilibrium status $f^{\text{eq}}_i$ is given based on the two local macroscopic lattice properties: lattice velocity $u(x,t)$ and lattice density $\rho_{LB}(x,t)$. To ensure the mass and momentum conservation during the collision and streaming process, specific equations and weights $\omega_i$ are required for different models. In the D2Q9 model, the equilibrium distribution functions are written as

$$f^{\text{eq}}_0(x,t) = \omega_0 \rho_{LB}(x,t) \left[ 1 - \frac{3}{2} u_{LB}(x,t)^2 \right]$$

$$f^{\text{eq}}_i(x,t) = \omega_i \rho_{LB}(x,t) \left[ 1 + 3 \varepsilon_i \cdot u_{LB}(x,t) + \frac{3}{2} (3 \varepsilon_i \cdot u_{LB}(x,t))^2 - u_{LB}(x,t)^2 \right]$$

where $\omega_i$ represents a series of weights of a given lattice model to maintain the isotropy of the fourth-order tensor of velocities and the Galilean invariance. The weight distribution of the points within a single lattice grid for three widely used lattice models is described in Table 1.
Table 1. Weights $\omega$, for Lattice Models D1Q3, D2Q9, and D3Q19

<table>
<thead>
<tr>
<th>Model</th>
<th>center</th>
<th>horizontal/vertical</th>
<th>diagonal</th>
</tr>
</thead>
<tbody>
<tr>
<td>D1Q3</td>
<td>2/3</td>
<td>1/6</td>
<td>0</td>
</tr>
<tr>
<td>D2Q9</td>
<td>4/9</td>
<td>1/9</td>
<td>1/36</td>
</tr>
<tr>
<td>D3Q19</td>
<td>1/3</td>
<td>1/18</td>
<td>1/36</td>
</tr>
</tbody>
</table>

Take the D2Q9 model used in this work as an example; $\omega_5 = 4/9$ in eq 5 describes the equilibrium function for the center point within the grid block, where the rest of the 8 points are given by eq 6 with $\omega_1 = 1/36$ or 1/9 depending on the geometrical position of the point with respect to the grid block. Figure 2 demonstrates the lattice arrangement and velocity vectors for a D2Q9 model. The center point is labeled as 0, and indexes 1, 2, 3, and 4 indicate the velocity directions to the center of four sides, where 5, 6, 7, and 8 point to the four corners of the lattice grid.

The $e_i$ is determined by

$$
\begin{align*}
&e_i = \begin{cases}
0, & i = 0 \\
\epsilon^i = \frac{2}{\sqrt{\delta}} \left( \cos \left( \frac{(i-1)\pi}{4} \right) , \sin \left( \frac{(i-1)\pi}{4} \right) \right), & i = 1,2,3,4 \\
\sqrt{2} \epsilon = \left( \cos \left( \frac{(i-1)\pi}{4} \right) , \sin \left( \frac{(i-1)\pi}{4} \right) \right), & i = 5,6,7,8
\end{cases}
\end{align*}
$$

Equation 7 is converted to the following form

$$
\begin{align*}
\epsilon_0 & = (0,0) \\
\epsilon_1, \epsilon_2, \epsilon_4 & = (\pm 1,0), (0, \pm 1) \\
\epsilon_{5,6,7,8} & = (\pm 1, \pm 1)
\end{align*}
$$

The overall BGK LBM model is given by the following evolution equation

$$
f^i_{\text{eq}}(x + e_i, t + 1) = f^i(x, t) - \Omega_c[ f^i(x, t) - f^i_{\text{eq}}[\rho_{\text{LB}}(x, t), u_{\text{LB}}(x, t)] ]
$$

where $\Omega_c$ is the relaxation frequency defined by $\Omega_c = 1/t$; the relaxation time $\tau$ indicates the time used for the system to reach equilibrium during each streaming-collision cycle, which is associated with the lattice kinematic viscosity $\nu_{\text{LB}}$

$$
\nu_{\text{LB}} = (\tau - 1/2)c_s^2
$$

A typical iteration cycle of an LBM BGK model starts with the initialization of the lattice velocity and lattice density to an equilibrium state based on the initial condition. Furthermore, the macroscopic quantities (e.g., lattice pressure, density, velocity) are obtained for the initial equilibrium state using eqs 2–4. Meanwhile, the calculated quantities for the current time step are used for computing the equilibrium function $f^i_{\text{eq}}$ based on eqs 5 and 6 (or the D2Q9 model in this case). Next, the collision and streaming processes based on the evolution eq 9 are executed. Specifically, the local particle distribution function $f_i(x, t)$ is altered by the current equilibrium function $f^i_{\text{eq}}$ within the relaxation time $\tau$ described in the evolution function. Followed by the collision step, the streaming step simply transfers the updated distribution functions to adjacent nodes, and a full iteration is completed. Boundary treatments are executed after the full cycle if applicable.

The general steps of the BGK LBM model can be described in Figure 3.

**2.2.2. LBM Advection–Diffusion Model.** A series of advection–diffusion types of problems can be formed within the LBM modeling framework, for instance, chemical transport,\textsuperscript{24,25} heat transfer,\textsuperscript{30,31} multicomponent reactive transport in porous media,\textsuperscript{25} and reactive transport associated solid dissolution modeling.\textsuperscript{25} Some of the studies coupled the LBM flow model with a third-party reaction solver to model the complex pore-scale reactive transport scenarios.\textsuperscript{30,31,93–96} More
related studies on LBM reactive transport-associated applications can be found in review papers.\textsuperscript{46,97}

A typical advection–diffusion formulation is shown below

$$\frac{\partial C}{\partial t} + \nabla \cdot (C \mathbf{u}) = \nabla \cdot (D \nabla C) + q$$

(11)

where $C$ represents a scalar field for quantities that can be transported by the flow while diffuse based on the concentration or temperature gradient, $\mathbf{u}$ stands for the flow or advection velocity vector, $D$ is the diffusion coefficient, and $q$ is an optional source term which can be used to represent the locally destroyed or produced chemical species (if $C$ is for chemical concentration field) or represents the consumption or production of heat (if $C$ is the temperature field). These behaviors can be attributed to the mechanisms such as chemical reactions. In terms of the diffusion process, the associated LBM formulation can be written as the following evolution function

$$f_{AD_i}(x + c_i \Delta t, t + \Delta t) - f_{AD_i}(x, t) = \Omega_i(x, t)$$

$$+ Q_i(x, t)$$

(12)

The $Q_i(x, t)$ represents the same meaning as the source term $q$ in eq 11 in a “discretized” form with respective to lattice space and time. For example, if we have an additional reaction solver to “produce” or “destroy” chemical species in each lattice grid at each lattice time step, this would be added in the “$Q$” source term of the LBM evolution equations to incorporate the advection–diffusion process.

The relaxation frequency is calculated as

$$\Omega_i(x, t) = \frac{1}{\tau_{AD_i}}[f_{AD_i}(x, t) - f_{AD_i}^{eq}(x, t)]$$

(13)

The concentration or temperature term $C$ is recovered by the equation below similar to the lattice density $\rho$ in the BGK LBM flow model

$$C = \sum_i f_{AD_i}$$

(14)

Meanwhile, the diffusion coefficient $D$ is obtained by the following equation in light of the lattice kinematic viscosity ($\nu$) in the BGK LBM flow model

$$D = C_s^2 \left( \frac{\tau_{AD}}{2} - \frac{\Delta t}{2} \right)$$

(15)

The associated lattice equilibrium distribution function is written as

$$f_{AD_i}^{eq} = w_i \left[ 1 + \frac{c_i \cdot u}{C_s} + \frac{(c_i \cdot \mathbf{u})^2}{2C_s^2} - \frac{\mathbf{u} \cdot \mathbf{u}}{2C_s^2} \right]$$

(16)

The advection and diffusion coupling can be achieved by explicitly linking the concentration or temperature field $C$ with another fluid flow lattice with a unified solving time step.

### 2.2.3. PALABOS LBM Solver

PALABOS is chosen as the fluid-solving model due to its advantages, including but not limited to parallel, coupling, and complex boundary treating capabilities. The PALABOS (parallel lattice Boltzmann solver) package is selected as the base of the modeling framework for the fluid flow component.

PALABOS\textsuperscript{98} is an open-source LBM-based fluid flow solver built with the C++ programming language, developed and maintained by the University of Geneva. It is capable of solving single/multiple-phase fluid flow and convection–diffusion type problems in open or closed geometry. It also features parallel computing capabilities by utilizing a few to thousands of CPUs to speed up the simulations. Meanwhile, PALABOS provides flexible application programming interfaces (APIs) to extend the existing capabilities, such as modeling fluid–solid interactions by coupling a large-scale atomic/molecular massively parallel simulator,\textsuperscript{99} or simulating blood cell behaviors by coupling structural solver npFEM.\textsuperscript{100,101}

Regarding the extensive APIs, PALABOS supports convenient non-local operations by “data processors” in addition to in-place LBM particle collision and streaming operations. Data processors enable robust outside algorithm implementation on either lattices or additional layers of data fields. Examples of this type of implementation include advection–diffusion coupling, multiphase flow, and solid–fluid interactions.

### 2.3. Geochemical Reaction Model

#### 2.3.1. Geochemical Reaction Solver

Similar to the selection of the fluid flow solver, the widely used open-source geochemical reaction solver PHREEQC (pH-Redox-Equilibrium)\textsuperscript{102} developed and maintained by the United States Geological Survey (USGS) is selected in this work as the geochemical solver. PHREEQC is a scripting input-based reaction solver written in C and C++ languages for equilibrium, kinetic, ion-exchange, surface complexation reactions, solid solutions, and 1D reactive-transport solving capabilities.

In addition to the original version of PHREEQC, which is equipped with a graphical user interface, other variants are developed to be able to couple with thirty-party codes. For example, IPhreeqc\textsuperscript{44} utilizes the Microsoft COM interface for communicating with outside models, which is widely used in modeling reactive transport.\textsuperscript{103,104,105} Another variant PHREEQCRM\textsuperscript{106} is used in this study, enabling most of the low-level APIs to be exposed to developers for deep code integrations while preserving a flexible scripting input system. Meanwhile, the reactions can be solved in parallel by utilizing mature interfaces such as OpenMP and MPI schemes. The implementation of PHREEQCRM library includes FEFLOW,\textsuperscript{59} USGS PHAST,\textsuperscript{57,58} and modeling ionic transport.\textsuperscript{100}

#### 2.3.2. PHREEQCRM Geochemical Reaction Models

One of the appealing features of PHREEQCRM is the “batch reactor” functionalities, which allow one or multiple geochemical reactions to be calculated within a single grid block at a time. Also, it is possible to enable different types of reactions flexibly by using their scripting input system. The available reaction types (keywords) include: (1) solutions, (2) Equilibrium -, Phases, (3) exchange, (4) surface, (5) Gas Phase, (6) Solid Solutions, and (7) kinetics. Before executing different reactions, PHREEQC takes user-defined reactions and initializes the “batch reactor” to equilibrium with the initial chemical components.\textsuperscript{35} In this study, three reaction types were considered: solutions, kinetics, and surface for aqueous phase equilibrium reaction, kinetic reactions, and surface complexation reactions (SCM) on the solid surface.

A discretized 2D/3D simulation domain contains thousands or millions of grid blocks that are considered parallel batch reactors in PHREEQCRM. They are independent of one another. The transport of chemical species is executed by a third-party flow model (PALABOS LBM model in this study) and moves in or out of species within a reaction cell. PHREEQCRM solves the equilibrium reactions and returns the updated species concentrations back to the flow model.\textsuperscript{102}
2.4. Model Coupling Interface. A coupling interface is developed to connect the LBM flow solver PALABOS and the geochemical reaction solver PHREEQCRM in a dynamic fashion to solve pore-scale reactive transport problems. Both of the solvers are written in the C++ programming language, with APIs exposed to users to implement new functionalities, so the target of this task is to build an interface that is easy to use while maintaining high communication efficiency.

2.4.1. Data Pre-/Post-Processing and Script Input. The data preprocessing includes a series of functions for preparing the simulation data for fluid flow or chemical reaction domains. For instance, one of the functions is used to process the micro-CT scanned images to the simulation model readable ASCII/binary format data structure. Other functions are used for simulation parameters' determination based on the model constraints. Also, a unified XML format-based script reading system is developed to preprocess the user inputs. The user input script includes various parameters for PALABOS and PHREEQCRM as well as other auxiliary features. Meanwhile, the code loads the ".pqi" PHREEQC input script with detailed chemical reaction definitions at the model initialization stage. The data post-processing includes various help functions to execute the main simulations, result analyses, and visualizations.

2.4.2. Model Initialization. The model initialization stage includes two steps: (1) simulation domain and reaction-type assignment; (2) initial equilibrium of the flow and reaction models. In the first step, the coupled model establishes the simulation domain from loaded porous media geometry information and creates cells mappings from the initial fluid—solid distribution. In addition to setting up the inlet/outlet flow and chemical flux boundary conditions, the "Bounce-Back" or "No-Flow" conditions are assigned within the "solid" grid blocks in LBM flow and diffusion lattices. Furthermore, the associated "solid" cells in PHREEQCRM are treated as "inactive" for geochemical reactions.

In the second step, the PHREEQCRM executes the initial sets of reaction calculations based on the user-defined equations in those reaction "active" cells, then obtains one or more scalar fields with the initial species concentration. It is worth noting that only the kinetic and surface complexation reactions on a type of cells called "solid interface cells" (see more details in Section 2.4.3, Cell Mapping Model) are enabled, where no flow exists but diffusion is enabled for mass transfer between the solid surface and outside aqueous phase cells.

On the other hand, the model initializes the fluid flow lattice from the initial and flow boundary conditions and obtains the initial velocity distribution. The initial velocity distribution fields were coupled with one or multiple concentration fields from PHREEQCRM, and the main simulation is ready to launch.

2.4.3. Cell Mapping Model. A unified cell mapping model was proposed to handle the fluid—fluid and fluid—solid interactions and geochemical reactions within different cell types. A dynamic mapping system is built on top of the porous media geometry, which initializes before the main simulation. The system includes four cell types:

(1) Fluid cells: cells represent the void space in the porous media
(2) Fluid interface cells: cells belong to "fluid cells" and are positioned at the outer surface layer cells of the "fluid cells"
(3) Solid cells: cells without any fluid flow and reaction enabled, defined from the initial input geometry
(4) Solid interface cells: cells belong to and occupy the outer surface layer of the "solid cells"

The fluid cells host the LBM advection—diffusion and the PHREEQC equilibrium reactions. No flow or reactions happen within the solid cells. However, the equilibrium, kinetics, and surface complexation reactions on the outer layer (solid surface cells) are enabled. Meanwhile, the solid dissolution and precipitation processes are implemented within solid interface cells. The solid interface layer is still considered as "solid" due to the disabled fluid flow, but the diffusion is enabled for information exchange between them and aqueous cells since the solid might be partially dissolved. Another type of cells called fluid interface cells are also defined to handle future potential applications when complex fluid—fluid interactions are involved and in cases where the pore-scale geometry changes due to mineral precipitations.

Table 2 summarizes the advection—diffusion and reaction availability within the proposed four cell types.

<table>
<thead>
<tr>
<th></th>
<th>fluid cells</th>
<th>fluid interface cells</th>
<th>solid cells</th>
<th>solid interface cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>fluid flow</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>diffusion</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>equilibrium reaction</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>kinetic reaction</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>dissolution</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>precipitation</td>
<td>no</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>surface complexation</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>yes</td>
</tr>
</tbody>
</table>

*The solid cells are inactive, whereas fluid cells only host advection—diffusion—reaction in the aqueous phase, and the interface cells are located at the boundary of the solid—fluid interfaces, hosting the fluid—solid interaction-associated processes except for the fluid flow.*

2.4.4. PALABOS Data Processor. PALABOS has a flexible and powerful built-in data processor function, utilizing the C++ template programming features. The data processor allows the user to create flexible operations between lattices, scalar fields, and tensor fields in parallel. This function provides the foundation for coupling other models within PALABOS or from the third-party code. Various customized data processors are developed to couple the chemical concentration fields with the fluid flow lattice. In this work, a cross-lattice data processor is built to connect the fluid flow velocity field lattice and multiple advection—diffusion field lattices. Hence, in each simulation time step, the displacements caused by fluid flow are imposed on the concentration fields for all chemical species in the aqueous phase.

2.4.5. PHREEQCRM Basic Function Feature. PHREEQCRM shares a majority of the functionalities and features with the standalone version of PHREEQC, including a basic function input interpreter. Under specific formatting standards, the interpreter loads the user-defined input script (.pqi file). In addition to the pre-defined keywords followed by different reaction equations, basic mathematic operations and arithmetic functions can be defined in the script for a single reaction cell in PHREEQCRM. Furthermore, the defined operations are optimized in parallel similar to the reaction solver. Thus, customized analytical or numerical analysis on concentration
fields can be done efficiently without spending a significant amount of data communication overhead due to data transfer. For instance, in this work, the calculation of the zeta potential \( \zeta \) is achieved using a PHREEQCRM basic function executed after the general internal geochemical reactions are solved during each iteration. \( \zeta \) is obtained by ref 103

\[
\zeta = \psi e^{-md}
\]

(17)

where \( d \) is the distance between the slipping plane and the stern layer in the electrical double layer model, \( \psi \) stands for the surface potential, and \( \kappa \) represents the inverse of the Debye length which is obtained by

\[
\kappa^{-1} = \sqrt{\frac{\epsilon_\lambda \epsilon_0 k T}{2 N_a e^2 I}}
\]

(18)

where \( N_a \) is the Avogadro’s number; \( \epsilon \) is the charge of an electron; \( I \) represents the ionic strength; \( \epsilon_\lambda \) and \( \epsilon_0 \) are the temperature-dependent relative permittivity of the solution and the vacuum permittivity \( (8.8541878128 \times 10^{-12} \text{ F m}^{-1}) \), respectively; \( k_b = 1.38064852 \times 10^{-23} \text{ J K}^{-1} \) is the Boltzmann constant; and \( T \) is the temperature in Kelvin.

2.5. AI-Assisted SCM Model Optimizations. The SCM describes solute adsorption from the aqueous phase to the mineral surface. It is also considered one of the most explainable models for the fluid–solid interactions between the crude–oil–brine–rock (COBR) interfaces during advanced waterflooding EOR processes. One of the challenges in building the SCM models for COBR investigations is the adjustment of the SCM reaction constants \( (\log K) \) based on the experimental measured zeta potentials \( (\zeta) \). To validate the SCM model integration for the coupling code, an SCM model based on the experimental data by Tetteh et al. \( 105,106 \) is developed and optimized for the brine/calcite interface. Instead of estimating and calibrating the \( \zeta \) within a bulk fluid solution, a more realistic 2D simulation domain is created with a centered spherical calcite grain surrounded by solutions with various salinities. Then, the PALABOS–PHREEQCRM coupled solver is used to obtain the surface properties such as \( \zeta \), surface potentials, and Debye length. The simulated \( \zeta \) over the surface of the calcite grain is averaged and used as the target to compare with the experimentally measured ones and re-adjust the reaction constants if they do not match. Since the directions for adjusting the sets of reaction constants are unknown, and each one of the numerical simulations takes time, an optimization framework is developed to assist the parameter adjustment tasks. The framework connects the geochemically coupled PALABOS–PHREEQCRM simulator with a multi-layer perceptron (MLP) neural network (NN). A series of pre-simulated model parameters with the calculated \( \zeta \) are prepared as the pre-train data for NN. The NN and simulator communicate with each other interactively to update the log \( K \) and match the target \( \zeta \) as much as possible. The conventional mean squared error (MSE) loss function and the stochastic gradient descent (SGD) optimizer are used during the interactive training and optimization process.

Figure 4 depicts the overall workflow of the proposed optimization framework.

3. MODEL VALIDATION AND RESULTS

In this section, the model setup and validation results for the proposed framework are presented. The following model validation experiments are included: (1) an LBM single-phase fluid flow in a 2D channel used to verify basic fluid flow; (2) an LBM diffusion in a 2D channel experiment to verify basic Fick’s diffusion; (3) a reactive transport in a 2D channel experiment with a dissolving circular-shaped calcite grain (this case is used to verify the reactive transport coupling and surface reactions/calcite dissolution kinetics); and (4) a static SCM model in a 2D domain with artificial neural network (ANN) optimization (this case is used to verify the SCM modeling capabilities integrated in the framework) as well as the ANN optimization feature.

3.1. LBM Single-Phase Fluid Flow in a 2D Channel. The first validation experiment designed is a 2D Poiseuille flow scenario, as shown in Figure 5. A 2D rectangular-shaped simulation domain is created with a 300 \( \times \) 100 lattice resolution, which represents a real-world physical dimension of 6 cm \( \times \) 2 cm rectangle. An artificial pressure gradient \( (2 \times 10^{-8} \text{ [Pa]}) \) is applied across the channel with an inlet on the left and an outlet on the right. The no-flow boundary conditions are imposed on the top and bottom walls. Water is assumed as the fluid used in this experiment at a standard temperature of 25 °C, and it has a density of 997 [kg/m³], viscosity of 8.891 \( \times \) 10^{-4} [Pa-s], and kinematic viscosity of 8.917 \( \times \) 10^{-7} [m²/s]. The fluid flow is
assumed to be a steady-state laminar flow, and the viscous dissipation and gravity effects are neglected.

To verify the solutions, the analytical solutions are derived based on the mass and momentum conservation equations

\[ \nabla \cdot \vec{u} = \frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} = 0 \]  

(19)

\[ \rho \left( \frac{\partial \vec{V}}{\partial t} + \nabla \cdot \vec{V} \right) = \mu \nabla^2 \vec{V} - \nabla P + \rho g \]  

(20)

After simplifying the momentum equation with the integration of boundary conditions, the below equation is obtained for the velocity profile

\[ u(y) = -\frac{H^2 dp}{2\mu \Delta x} \left[ 1 - \left( \frac{y}{H} \right)^2 \right] \]  

(21)

where \( y \) is the coordinate across the width of the channel, \( H \) is the width of the channel, \( \mu \) is the fluid viscosity, and \( dp/dx \) represents the assigned pressure gradient along the length of the channel. The average velocity is computed as

\[ u(\text{avg}) = -\frac{H^2 dp}{3\mu \Delta x} \]  

(22)

From eq 22, the maximum velocity calculated at the center is \( 1.87436 \times 10^{-8} \) [m/s] and the average velocity is \( 1.24970 \times 10^{-8} \) [m/s]. The same physical properties are applied and converted to lattice units, and the LBM simulation is conducted. The pressure gradient is represented by the lattice density gradient map shown in Figure 6.

\[ \text{Figure 6. Lattice density map generated from the given pressure gradient.} \]

The 2D velocity profile obtained from LBM simulation after the steady state is reached, as shown in Figure 7.

The 2D velocity distribution is evaluated from the LBM simulation, and the computed maximum \( 1.24371 \times 10^{-8} \) [m/s] and average \( 1.024970 \times 10^{-8} \) [m/s]. The same physical properties are applied and converted to lattice units, and the LBM simulation is conducted. The pressure gradient is represented by the lattice density gradient map shown in Figure 6.

\[ \text{Figure 7. Lattice velocity distribution induced by the pressure gradient of the fluid flow channel.} \]

\[ \text{Figure 8. LBM simulated velocities (red dots) against the analytical solutions (blue curve).} \]

\[ \text{Figure 9. Model setup for the LBM diffusion validation in a 2D channel.} \]

creating a fluid flow inlet from the left end, a single chemical species with a constant concentration \( c_0 = 1.0 \) is imposed. The right-hand side of the channel is assumed to be infinitely long \( (c_{inf} = 0) \), and the boundary conditions and boundary effects are not considered for top and bottom walls.

The following assumptions are made in our model setup: (1) there is no boundary effects such as surface adsorption to influence the diffusion; (2) the process is considered isothermal such that the rate of diffusion is not affected by the temperature; (3) the process is considered isotropic diffusion, which is the same in every direction and is characterized by a single diffusion coefficient \( D \); and (4) the flow is one-dimensional Fick’s diffusion with a constant diffusion coefficient \( D = 10 \times 10^{-6} \) [m/s]. The analytical solution of the concentration profiles is solved from Fick’s second law
\[
\frac{dc}{dt} = D \frac{\partial^2 c}{\partial x^2} 
\]  
(23)

The corresponding analytical solutions for the semi-infinite channel with a constant species concentration on one side is
\[
c(x, t) = c_0 \text{erfc} \left( \frac{x}{2 \sqrt{Dt}} \right) 
\]  
(24)

where \( c_0 \) is the constant concentration (\( c_0 = 1 \)) on the left terminal, \( D \) represents the diffusion coefficient, \( t \) is the time, \( x \) is the location from the left terminal, erfc stands for the complementary error function, and \( c(x,t) \) describes the concentration of the distance \( x \) from the left terminal at time \( t \). 7 time points \(( t = 1, 20, 50, 100, 150, 200, 250 \text{ s} )\) are assigned as the testing time for the concentration profile comparison evaluations.

A pure concentration field is created for single species in the LBM model without implementing any fluid flow in the channel. The lattice diffusivity is obtained from the physical diffusion coefficient and the domain lattice time. Figure 10 illustrates the concentration distribution at the initial stage \(( t = 1 \text{ s} )\) from the LBM simulation. The 1D concentration profile is extracted from the center line across the domain as shown in Figure 10 at the given time steps.

Figure 11 demonstrates the species concentration distribution at the given time points and the evaluation of the concentration profile at the center cross line at those time steps.

The comparison results are shown in Figure 12. Good agreement is observed between the analytical solutions (curves) and LBM simulated results (dots).

The errors are measured quantitatively regarding three metrics: mean absolute error (MAE), MSE, and coefficient of determination \( (R^2) \) which are defined in Appendix A. The evaluation results are shown in Table 3. The LBM simulated concentration profiles are in very good agreement with the analytical solutions at all time points except at \( t = 1 \text{ s} \). This can be attributed to the fluctuations at the beginning of the simulation, and the error becomes significantly smaller as the time steps go by.

3.3. Coupled Model for Advection—Diffusion—Reaction Transport in a 2D Channel with Calcite Dissolution Kinetics. The model validation results for single-phase fluid flow and pure diffusion from Sections 3.1 and 3.2 create a solid foundation for further reaction coupling. In this section, the advection—diffusion—reaction capabilities of the proposed framework are presented and verify the model performance by using the benchmark problem proposed by Molins et al.\(^{107}\) This reactive transport benchmark problem is designed to evaluate the dynamic concentration fields in a 2D channel with a circular-shaped calcite grain located at the center. The average reaction rate is evaluated from the concentration differences between the inlet and the outlet. The simulation domain (Figure 13) is a rectangular-shaped 2D flow channel with a length of 0.1 cm and a width of 0.05 cm. A representing lattice grid is created with a resolution of 256 × 128. A circular-shaped calcite grain with a diameter of 0.02 cm is placed at the center of the channel. A constant velocity \((0.12 \text{ cm/s})\) flow boundary condition is imposed at the left terminal. Meanwhile, an outflow condition is assigned to the outlet at the right terminal, indicating that no pressure gradient exists, or any driving force/resistance applied at the outlet. The top and bottom walls are non-reactive and impermeable and have no-flow boundaries.

In the initial stage, the flow channel is assumed to be filled with hydrochloric acid \((\text{HCl})\) with a concentration of \( 10^{-2} \text{ [mol/L]} \), which results in a \( \text{pH} = 2 \). After the injection starts, the solution with a concentration of HCl is pushed into the channel continuously. Meanwhile, the irreversible heterogeneous reaction starts to dissolve the calcite based on the stoichiometric equation
\[
\text{CaCO}_3(s) + \text{H}^+ \rightarrow \text{Ca}^{2+} + \text{HCO}_3^{-} 
\]  
(25)

and the kinetic reaction rate factor \((rf)\) for the calcite dissolution is dependent on the \( \text{H}^+ \) concentration
\[
rf = k_{\text{H}^+} \gamma_{\text{H}^+} c_{\text{H}^+} 
\]  
(26)

where \( k_{\text{H}^+} \) is the rate constant with units of \([\text{mol} \cdot \text{cm}^{-2} \cdot \text{s}^{-1}]\), \( \gamma_{\text{H}^+} \) is the activity of \( \text{H}^+ \) with units of \([\text{cm}^3 \cdot \text{mol}^{-1}]\), and the \( c_{\text{H}^+} \) is the concentration of \( \text{H}^+ \) with units of \([\text{mol} \cdot \text{cm}^{-3}] \) or \([\text{mol} \cdot \text{L}^{-1}] \).

The detailed parameters of the benchmark problem are shown in Table 4.

The evolution of the concentration profile is evaluated according to the methods from Molins et al.\(^{107}\) with the assumptions that several impact factors need to be considered: (1) the velocity distribution influenced by the inlet flow rate and the calcite grain; (2) the instantaneous equilibrium reactions in the fluid phase; (3) the kinetic reaction rate on the calcite surface; and (4) the multicomponent diffusion in the fluid phase as well as the diffusive boundary layer defined on the calcite surface. In addition, the chemical reactions are assumed not to influence the fluid flow. The average reaction rate \((R)\) is calculated as
\[
R = \frac{Q(c_{\text{out}} - c_{\text{in}})}{\xi A} 
\]  
(27)

where \( \xi \) is the stoichiometric coefficient, \( A \) stands for the reacting surface area of the calcite, \( c_{\text{in}} \) represents the universal inlet concentration from the boundary condition, and \( c_{\text{out}} \) is evaluated as the flux-weighted-average concentration across the outlet boundary
\[
c_{\text{out}} = \int u \cdot ds / Q 
\]  
(28)

where \( Q \) is obtained from the integration across the outlet
\[
Q = \int u \cdot ds 
\]  
(29)
To calculate the reaction rate regarding the volumetric properties, a height of 1 cm is attributed to the 2D simulation domain for the benchmark problem.

Figure 14A demonstrates the steady-state velocity distribution from the LBM fluid flow lattice domain; the red color represents relatively higher velocity located at the top and bottom surface of the calcite grain.

Figure 14B shows the Ca\(^{2+}\) concentration distribution at the same moment with respect to Figure 14A. The formation of the Ca\(^{2+}\) concentration thin layer is observed around the grain as a result of the calcite dissolution, and the teardrop-shaped overall concentration pattern indicates the impacts of the flow velocity and direction.

Similarly, Figure 15A,B illustrates the H\(^{+}\) concentration evolution at the start of the simulation (t = 0.2 s) and after reaching a steady state at around t = 1.5 s. The H\(^{+}\) concentration distribution pattern demonstrates the H\(^{+}\) consumption during the calcite dissolution processes, which matches the Ca\(^{2+}\) concentration distribution pattern shown in Figure 14B.

<table>
<thead>
<tr>
<th></th>
<th>MAE error</th>
<th>MSE error</th>
<th>R(^2) score</th>
</tr>
</thead>
<tbody>
<tr>
<td>T = 1 s</td>
<td>0.0030</td>
<td>0.0003</td>
<td>0.9772</td>
</tr>
<tr>
<td>T = 20 s</td>
<td>0.0006</td>
<td>2.1829 \times 10^{-6}</td>
<td>0.9999</td>
</tr>
<tr>
<td>T = 50 s</td>
<td>0.0004</td>
<td>5.4259 \times 10^{-7}</td>
<td>0.9999</td>
</tr>
<tr>
<td>T = 100 s</td>
<td>0.0003</td>
<td>1.9075 \times 10^{-7}</td>
<td>0.9999</td>
</tr>
<tr>
<td>T = 150 s</td>
<td>0.0003</td>
<td>1.1620 \times 10^{-7}</td>
<td>0.9999</td>
</tr>
<tr>
<td>T = 200 s</td>
<td>0.0004</td>
<td>4.2959 \times 10^{-7}</td>
<td>0.9999</td>
</tr>
<tr>
<td>T = 250 s</td>
<td>0.0009</td>
<td>3.1512 \times 10^{-6}</td>
<td>0.9999</td>
</tr>
</tbody>
</table>

Figure 11. 2D concentration distribution from LBM simulation at t = 20, 50, 100, 150, 200, and 250 s.

Figure 12. Concentration profile comparison between LBM simulation and analytical solutions at t = 1, 20, 50, 100, 150, 200, and 250 s.
The average effluent H⁺ concentration evolution at the right terminal of the channel directly from the concentration field from the LBM model is evaluated. Equation 27 is used to estimate the average reaction rate, and the results are compared according to Mollins et al. The results are shown in Figure 16 for the outlet average effluent H⁺ concentration evolution from the beginning of the simulation until the steady state at 1.5 s. The overall trend from PALABOS–PHREEQCRM agrees with the other five codes despite the initial bump. The small bump is caused by the equilibrium reaction results from the initialization stage. Specifically, we believe that the PHREEQCRM generates slightly higher H⁺ concentrations (about 1.4%) at the assigned temperature of 40 °C; even we explicitly specified pH = 2.0. However, the effluent concentration reaches the desired equilibrium when the calcite dissolution kinetic reaction dominates the process, although the initial condition for the vortex method is different (pH = 7.0) compared to other methods, and the slightly different initial conditions do not affect the later and final solutions.

Meanwhile, Figure 17 shows the average dissolution rate over time for the PALABOS–PHREEQCRM framework and the other five codes. The dissolution rate curve and the trend of the proposed model demonstrate good agreement among the other five codes. In addition, PALABOS–PHREEQCRM has a slightly higher rate at the endpoint of this comparison with a rate of $4.59 \times 10^{-8}$ (mol·cm⁻²·s⁻¹); however, it is comparable to the other “Lattice–Boltzmann” approach $4.57 \times 10^{-8}$ (mol·cm⁻²·s⁻¹) described in the benchmark problem (Table 5). This may be due to the slightly increased surface area attributed to the nature of the LBM-based methods that use “staircase”-shaped pixels/voxels to replace smooth boundary curves.

### 3.4. Static SCM Model and ANN Optimizations

In this section, the model validation for the SCM model and associated optimization using an ANN are presented. The SCM model is built based on the experimental work from Tetteh et al., to investigate the electrokinetics on limestone surfaces when various brine compositions and salinities are applied. The rock sample is taken from Indiana limestone, which contains 99% calcite, and the solid is assumed to be 100% calcite in the modeling process for simplicity. Tetteh et al. synthesized the brines based on the water compositions from the Lansing Kansas City (LKC) group and then diluted the brine to create low salinity and seawater samples. In addition, they synthesized five single salt solutions to represent the low salinity brine with the same ionic strength. Table 6 shows the detailed brine composition.

The associated zeta potentials ($\zeta$) are measured at the brine/calcite interfaces at both 25 °C (atmospheric conditions) and 40 °C (LKC reservoir conditions).

Extensive investigations have been done to study the calcite surface complexation reaction models and the reaction constants. The proposed reactions and reaction constants are summarized in Table 7. To keep consistency with the experimental work by Tetteh et al., the same calcite surface site density (4.95 sites/nm²) and calcite-specific surface area (1 m²/g) are deployed. Meanwhile, the $P_{CO_2} = 10^{-3.4}$ atm condition is used in the SCM model with calcite as equilibrated phases.

To validate the SCM model built on top of the PALABOS–PHREEQC framework, a 2D static model is designed instead of a simpler batch reactor shown in Figure 18. Brines with various compositions are filled in the closed system, and a pure calcite grain is placed at the center. To simplify the process, the equilibrium reactions are assumed to be instantaneous within the aqueous phase. The surface cells of the calcite grain (the “solid interface cells”) are set active to kinetic and calcite surface complexation reactions.

### Table 4. Simulation Parameters for the Reactive Transport Benchmark Problem

<table>
<thead>
<tr>
<th>parameters</th>
<th>symbol</th>
<th>value</th>
<th>units</th>
</tr>
</thead>
<tbody>
<tr>
<td>fluid density</td>
<td>$\rho$</td>
<td>1</td>
<td>g cm⁻³</td>
</tr>
<tr>
<td>kinematic viscosity</td>
<td>$\nu$</td>
<td>$10^{-2}$</td>
<td>cm²·s⁻¹</td>
</tr>
<tr>
<td>diffusion coefficient</td>
<td>$D$</td>
<td>$10^{-3}$</td>
<td>cm²·s⁻¹</td>
</tr>
<tr>
<td>inlet velocity</td>
<td>$u_{in}$</td>
<td>0.12</td>
<td>cm·s⁻¹</td>
</tr>
<tr>
<td>width of the channel</td>
<td>$\omega$</td>
<td>0.05</td>
<td>cm</td>
</tr>
<tr>
<td>specific grain reactive area</td>
<td>$S$</td>
<td>200</td>
<td>cm⁻¹</td>
</tr>
<tr>
<td>rate constant</td>
<td>$kH^+$</td>
<td>$10^{-4.05}$</td>
<td>mol·cm⁻²·s⁻¹</td>
</tr>
<tr>
<td>activity coefficient</td>
<td>$\gamma H^+$</td>
<td>1000</td>
<td>cm³·mol⁻¹</td>
</tr>
<tr>
<td>inlet concentration (pH = 2)</td>
<td>$C$</td>
<td>$10^{-5}$</td>
<td>mol·cm⁻³</td>
</tr>
<tr>
<td>Reynolds number</td>
<td>$Re = u_{in} \omega \nu^{-1}$</td>
<td>0.6</td>
<td></td>
</tr>
<tr>
<td>Péclet number</td>
<td>$Pe = u_{in} \omega D^{-1}$</td>
<td>600</td>
<td></td>
</tr>
</tbody>
</table>

“Simplified from ref 107.”
After the model initialization, PHREEQCRM calculates the SCM model-related properties at the grain surface cells, such as the surface charge, Debye length, and zeta potential ($\zeta$). The average surface $\zeta$ is calculated and compared to the experimentally measured values. Furthermore, the SCM reaction constants defined in the PHREEQCRM input script file are adjusted to improve the matching. Instead of manual tuning, an ANN is developed to optimize the reaction constants automatically.

The ANN is built with a relatively simple MLP network for this work on top of the TensorFlow 2 framework. The network itself has three hidden layers with 50, 100, and 50 neurons, respectively. "ReLU" activation functions are assigned to each hidden layer, and a dropout layer with a dropout rate of 0.2 is attached after each hidden layer to alleviate potential overfitting issues. The weights of the layers are initialized with random normal distribution before training. The adaptive moment and learning rate estimation (Adam) optimizer and the standard MSE are used during the network training. The pre-trained data are shuffled within the data loader before being fed in the main training loop with a batch size of 1024. The network inputs contain 9 features including 8 simulated zeta potential values from the coupled numerical model plus the temperature. The outputs are 6 SCM reaction constants that we would like to optimize such that the resulted zeta potentials can match the experimental measurements. The input data are normalized using “standardization” scaling which scales the inputs separately by subtracting the mean and dividing by the standard deviation, such that the data distribution will have a mean of 0 and a standard deviation of 1 for better fitting performance.

At the beginning of the optimization, parallel simulations are performed on a wide array of parameter (six reaction constants) combinations and generate the corresponding $\zeta_{SIM}$ values for ANN pre-training. The data generated for pre-training containing about 20,000–40,000 samples “warm up” the ANN in the first few training epochs, which leads to a general desired

Figure 14. (A) Velocity distribution under the steady-state condition; (B) 2D Ca$^{2+}$ concentration distribution in the steady state; the higher Ca$^{2+}$ concentration region surrounding the grain indicates the stable dissolution of CaCO$_3$ on the grain surface.
direction to converge. Note that the ANN takes \( \zeta_{\text{SIM}} \) as inputs and predicts reaction constants \( \log K \). The \( K_i \) is used to represent \( \log K \) values for simplicity. Furthermore, the interactive optimization is performed by (1) obtaining a set of \( K_i \) from ANN predictions; (2) assigning the ANN predicted \( K_i \) to PALABOS-PHREEQCRM and running coupled numerical simulations for all 8 solution cases under given temperature conditions; the resulting 8 \( \zeta_{\text{SIM}} \) values are used to compare the experimental measurements (\( \zeta_{\text{EXP}} \)) to compute the loss function; (3) representing the differences between \( \zeta_{\text{SIM}} \) and \( \zeta_{\text{EXP}} \) by the MSE loss

\[
\text{MSE} = \frac{1}{n} \sum_{i=1}^{n} (\zeta_{\text{EXP}} - \zeta_{\text{SIM}})^2
\]

where \( n \) is the number of ANN predictions, 8 is used since there are 8 \( K_i \), \( \zeta_{\text{EXP}} \) is the experimentally measured zeta potential and \( \zeta_{\text{SIM}} \) represents the numerical model simulated zeta potentials; and (4) updating the ANN weights based on the MSE loss; (5) using the updated ANN to predict a new set of \( K_i \) and enter the

**Figure 15.** (A) 2D H\(^+\) concentration distribution map; the color contour illustrates the beginning stage \( t = 0.2 \) s of the consumption of H\(^+\) due to calcite dissolution kinetic reaction on the solid surface. (B) 2D H\(^+\) concentration distribution map in the steady state \( t \geq 1.5 \) s.

**Figure 16.** Averaged effluent H\(^+\) concentration evolution from this study and the other five codes.\(^{107}\)
Table 6. Brine Compositions Used in This Study

<table>
<thead>
<tr>
<th>brine</th>
<th>Ca(^{2+})</th>
<th>Mg(^{2+})</th>
<th>Na(^+)</th>
<th>K(^+)</th>
<th>Cl(^-)</th>
<th>SO(_4^{2-})</th>
<th>IS (mol/L)</th>
<th>TDS, ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.04-NaCl</td>
<td>0</td>
<td>0</td>
<td>920</td>
<td>0</td>
<td>1418</td>
<td>0</td>
<td>0.04</td>
<td>2338</td>
</tr>
<tr>
<td>0.04-KCl</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1564</td>
<td>1418</td>
<td>0</td>
<td>0.04</td>
<td>2982</td>
</tr>
<tr>
<td>0.04-CaCl(_2)</td>
<td>534</td>
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<td>0</td>
<td>945</td>
<td>0</td>
<td>0.04</td>
<td>1480</td>
<td></td>
</tr>
<tr>
<td>0.04-MgCl(_2)</td>
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<td>0</td>
<td>945</td>
<td>0.04</td>
<td>1269</td>
<td></td>
</tr>
<tr>
<td>0.04-Na(_2)SO(_4)</td>
<td>0</td>
<td>0</td>
<td>613</td>
<td>0</td>
<td>0</td>
<td>1281</td>
<td>0.04</td>
<td>1894</td>
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<tr>
<td>FWS</td>
<td>11,000</td>
<td>2800</td>
<td>48,000</td>
<td>500</td>
<td>101,913</td>
<td>260</td>
<td>3.27</td>
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<tr>
<td>SWS</td>
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<td>560</td>
<td>9600</td>
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<td>52</td>
<td>0.65</td>
<td>32,895</td>
</tr>
<tr>
<td>LSW</td>
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<td>34</td>
<td>585</td>
<td>6</td>
<td>1243</td>
<td>3</td>
<td>0.04</td>
<td>2006</td>
</tr>
</tbody>
</table>

*The concentrations are in units of ppm prepared by Tetteh et al.\(^{105}\) FWS: formation water; SWS: seawater; LSW: low salinity water.

Figure 17. Averaged reaction/dissolution rate evolution from this study and the other five codes.\(^{107}\)

Table 5. Average Steady-State Dissolution Rate from This Study Compared to Other Codes Described by Reference 107

<table>
<thead>
<tr>
<th>code</th>
<th>surface area (cm(^2))</th>
<th>grain volume (cm(^3))</th>
<th>average rate (mol cm(^{-2}) s(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>theoretical</td>
<td>0.0628</td>
<td>3.14 \times 10^{-4}</td>
<td>4.59 \times 10^{-8}</td>
</tr>
<tr>
<td>this study</td>
<td>0.0628</td>
<td>3.14 \times 10^{-4}</td>
<td>4.59 \times 10^{-8}</td>
</tr>
<tr>
<td>OpenFOAM-DBS</td>
<td>0.0628</td>
<td>3.14 \times 10^{-4}</td>
<td>4.59 \times 10^{-8}</td>
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<td>lattice–Boltzmann</td>
<td>0.0628</td>
<td>3.14 \times 10^{-4}</td>
<td>4.59 \times 10^{-8}</td>
</tr>
<tr>
<td>vortex</td>
<td>0.0628</td>
<td>3.14 \times 10^{-4}</td>
<td>4.59 \times 10^{-8}</td>
</tr>
</tbody>
</table>
optimization framework can offer significant advantages over the manual tuning method in an automatic fashion.

Table 7. Surface Complexation Reaction Constants for the Brine/Calcite Interface Proposed in the Literature

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;CaOH + H^+ ↔ &gt;CaOH2^+</td>
<td>11.5</td>
<td>12.2</td>
<td>12.9</td>
<td>11.85</td>
<td>11.85</td>
<td>11.85</td>
</tr>
<tr>
<td>&gt;CaOH2^+ + SO_4^2- ↔ CaSO_4 + H_2O</td>
<td>2.89</td>
<td>2.89</td>
<td>2.1</td>
<td>2.1</td>
<td>2.1</td>
<td>2.1</td>
</tr>
<tr>
<td>&gt;CaOH + HCO_3^- ↔ &gt;CaCO_3^- + H_2O</td>
<td>5.6</td>
<td>4.9</td>
<td>3.32</td>
<td>5.8</td>
<td>4.28</td>
<td>5.8</td>
</tr>
<tr>
<td>&gt;CO_3H ↔ &gt;CO_3^- + H^+</td>
<td>-5.1</td>
<td>-4.9</td>
<td>-4.9</td>
<td>-5.1</td>
<td>-5.1</td>
<td>-5.1</td>
</tr>
<tr>
<td>&gt;CO_3H + Ca^{2+} ↔ &gt;CO_3Ca^{2+} + H^+</td>
<td>-1.7</td>
<td>-2.8</td>
<td>-3.16</td>
<td>-2.6</td>
<td>-2.6</td>
<td>-4.4</td>
</tr>
<tr>
<td>&gt;CO_3H + Mg^{2+} ↔ &gt;CO_3Mg^{2+} + H^+</td>
<td>-2.2</td>
<td>-2.2</td>
<td>-3.17</td>
<td>-2.6</td>
<td>-2.6</td>
<td>-4.4</td>
</tr>
</tbody>
</table>

4. CONCLUSIONS, DISCUSSION, AND FUTURE WORK

In this paper, a numerical modeling framework is proposed that integrates LBM-based fluid flow and advection–diffusion models with geochemical reaction models to study pore-scale physiochemical processes. The framework and the demonstrated initial stage of numerical validations paved the way for modeling complex and realistic subsurface scientific and engineering problems such as advanced waterflooding and CO_2 sequestration. Meanwhile, the framework is flexible and can be broadly applied in different pore-scale reactive transport simulations. In this work, PHREEQC is used as the geochemical reaction solver due to its capabilities of solving multiple reaction types as well as being able to couple third-party codes for multiphysics modeling. One of the variants of PHREEQC called PHREEQCRM is selected for the model building since it exposes low-level API for coupling purposes and works as a C++ library. Meanwhile, by using PHREEQCRM, much faster reaction solving and data communication between the two packages are obtained. In addition, a cell-mapping mechanism is developed within the framework to handle different reaction types at various cells, which allows the implementation of complex reactions such as SCM models on a solid surface. To validate the integrated modeling framework, a series of numerical experiments are designed to demonstrate the capabilities of this framework, ranging from simple 2D channel fluid flow and 2D diffusion to a 2D reactive transport benchmark problem. Furthermore, SCMs are added to the tests and validated against the SCM models with experimental measurements proposed in the literature. Meanwhile, a workflow using ANN to automatically assist the SCM model optimization is proposed. It successfully improves the SCM model tuning while
illustrating the great potential to optimize more complex scenarios or models.

In Section 2 (Methodologies), the background and formulation of the LBM method in fluid flow and advection–diffusion problems are introduced at first, and the LBM-based PALABOS package is discussed briefly. Second, the geochemical reaction solver PHREEQC and its capabilities as well as the variant PHREEQCRM selected in the modeling framework are discussed. Furthermore, the development details of the coupling interface are demonstrated. Specifically, the model/data preprocessing and initialization that provide essential steps for the coupling are discussed. The cell-mapping mechanisms
handle various reaction types within different mediums that allow modeling multiphysics in complex geometries. The PALABOS data processor and PHREEQC/RM basic function features provide great capabilities and flexibility to transfer information between different physical or chemical fields while preserving parallel computing performance for built-in or customized functions. At last, the ANN-based automatic numerical model optimization workflow is illustrated and is applied to the SCM model tuning using the model framework.

In Section 3 (Model Validation and Results), the numerical validations are designed and performed for the model in various aspects. First, in the 2D LBM fluid flow-only validation experiment based on a channel Poiseuille flow problem, the proposed model shows good agreement with analytical solutions with a small error (around 0.5%). Furthermore, the code is verified for the 1D diffusion problem in a 2D channel, and the time-dependent concentration profiles are evaluated with a small error (around 0.5%). The code is verified for the time-dependent averaged effluent H+ concentration is evaluated, and the averaged reaction/calcite dissolution rate as the metrics is estimated for comparison. The results demonstrate good agreement with the other five codes described in the benchmark problem. The last model validation focuses on the SCM model verification for complex physical–chemical processes using an advanced waterflooding-associated mechanistic study as an example. In addition to the SCM model integration, an ANN-assisted automatic SCM model tuning workflow is proposed that dynamically interacts with the numerical model to optimize the reaction constants. The results show good zeta potential estimations from SCM reactions compared to experimental data. Meanwhile, the ANN optimized SCM model parameters improve the overall matching performance while significantly reducing human labor in the manual model adjustment.

It is worth noting that the proposed coupling framework has its limitations at this initial stage of study. For example, we made assumption in our initial modeling practice that the relatively high concentrations of chemical species (Ca\textsuperscript{2+} in this case) do not reversely affect the fluid transport process due to intermolecular interactions, such that the LBM fluid flow is purely driven by pressure differences and affected by the no-flow boundaries at the walls. We plan to include the intermolecular interaction effects in the future modeling approaches.

Another limitation of this modeling framework is that we have not fully explored and utilized the lower-level source code and functionalities of PHREEQC/RM; thus, we do not have full control of some implementation details to meet our desired needs, for example, the initialization behavior with a small concentration bump described in Section 3.3. Future work would include the coupling improvements with in-depth integration of PHREEQC/RM for customized modeling needs and adjustments.

Meanwhile, all the numerical experiments are designed and proceed in a 2D domain, and we target pore-scale ranging from sub-micrometers/micrometers to centimeters (rock core or a portion of rock core). The numerical validation in a 2D symmetrical channel indicates the applicability of a 3D symmetrical domain such as a cylindrical pipe. We reserve the 3D modeling scenarios for future studies. It is worth noting that we intended to validate the coupled model in a relatively simple pore geometry instead of a realistic pore network due to the limitation of the computational resources. We treated this work as a “proof of concept” type of study, which paved the way for further investigations on more complex and realistic pore structures by incorporating a computing cluster with abundant computational resources.

## APPENDIX

### Evaluation Metrics

In this study, we use MAE, MSE, and coefficient of determination \( R^2 \) to evaluate the model validation results and the ANN prediction performance. MAE is defined as

\[
MAE = \frac{1}{n} \sum_{i=1}^{n} |Y_{i}^{\text{obs}} - Y_{i}^{\text{pred}}|	ag{31}
\]

\[
MSE = \frac{1}{n} \sum_{i=1}^{n} (Y_{i}^{\text{obs}} - Y_{i}^{\text{pred}})^2
\]

\[
R^2 = 1 - \frac{\sum_{i=1}^{n} (Y_{i}^{\text{obs}} - \bar{Y}_{\text{obs}})^2}{\sum_{i=1}^{n} (Y_{i}^{\text{obs}} - \bar{Y}_{\text{obs}})^2}	ag{33}
\]

where \( Y_{i}^{\text{obs}} \) stands for the \( i \)th observation, \( \bar{Y}_{\text{obs}} \) is the mean value of the observations, and \( Y_{i}^{\text{pred}} \) represents model or ANN predictions.

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### Notes

The authors declare no competing financial interest.
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NOMENCLATURE

CFD computational fluid mechanics
LBM lattice Boltzmann method
EOR enhanced oil recovery
Ω LBM collision operator
τ LBM relaxation time
\( f_k \) lattice distribution function
\( f_{eq} \) lattice equilibrium distribution function
\( \rho_{LB} \) lattice fluid density
\( P_{LB} \) lattice pressure
\( c_i \) lattice velocity
\( \nu_{LB} \) lattice kinetic viscosity
\( \omega_i \) weight coefficient for i direction
SCM surface complexation model
COBR crude–oil–brine–rock
\( \zeta \) zeta potential
\( H \) width of the flow channel
\( u(y) \) flow velocity profile along y-axis
\( p \) pressure
\( \mu \) fluid viscosity
\( nx \) number of lattice nodes along the x-axis
\( ny \) number of lattice nodes along the y-axis
\( c \) species concentration
\( c_0 \) constant species concentration at boundary
\( c_{inf} \) species concentration at the end of the assumed infinitely long channel
\( D \) diffusion coefficient
\( r_f \) reaction rate factor
\( k \) rate constant
\( \gamma \) activity coefficient
\( Re \) Reynolds number
\( Pe \) Péclet number
\( \xi \) stoichiometric coefficient
\( A \) reacting surface area
\( R \) average reaction rate
NN neural network
ANN artificial neural network
MLP multi-layer perceptron
SGD stochastic gradient descent
MAE mean absolute error
MSE mean squared error
\( R^2 \) coefficient of determination
\( Y_{obs} \) observation values
\( Y_{pred} \) prediction values
\( K_r \) reaction constant log K
FWS/SWS/LSW formation water/seawater/low salinity water

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