A fast stable discretization of the Constant–Convection–Diffusion–Reversible–Binding–Reaction (CCDRBR) equations

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A discretization scheme is introduced for a set of convection–diffusion–reaction equations with a non-linear reaction term describing reversible binding and a constant convection velocity for each reactant. This constancy of convection velocities allows a transformation to new spatial variables, which ensures the global stability of discretization. Convection–diffusion equations are notorious for their lack of stability, arising from the algebraic interaction of the convection and diffusion terms. Unexpectedly, our implemented numerical algorithm proves to be faster than computing exact solutions derived for a special case, while remaining reasonably accurate, as demonstrated in our runtime and error analysis.

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1. Introduction

1.1. Aims and overview

The aim of this paper is to develop a numerical algorithm for solving a system of convection–diffusion–reaction equations for a case of reversible binding reaction with constant convection velocities of each of the 3 reaction components (2 binders and their complex). These equations describe concentrations of the reaction components as functions of time and a spatial coordinate in kinetic separation methods, in which a reversible binding reaction is accompanied by separation of reaction components with constant convection velocities in a long and narrow tube (capillary or separation column). Practical kinetic separation methods have been developed for separation by capillary electrophoresis (methods of Kinetic Capillary Electrophoresis or KCE) [5,15,17,18,22,6,7,14,19,21] and size-exclusion chromatography (Kinetic Size-Exclusion Chromatography or KSEC) [3,2,6]. Though the convection is constant in kinetic separation, this constancy alone does not eliminate the instability of standard discretization schemes which convection–diffusion equations are notorious for [30], but it allows a change of variables which induces stability. The equations are introduced in Section 1.2 and discretized in Section 2.1.

The presented mathematical endeavor is fundamentally important for two directions of our ongoing scientific research concerned with kinetic separation: (1) the accurate computer simulation of experiments under properly setup experimental conditions; (2) the resolution of an experimental–computational inverse problem for determining kinetic rate constants of the reversible binding reaction. For these applications, a practical numerical solution method must be both stable and accurate, though for the inverse problem a low runtime of this direct solver is also critical.

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Previously, our simulations of kinetic separation were generated with COMSOL, which employs a streamline upwind Petrov–Galerkin method to solve a set of PDE describing mass transfer (see Sections 1.2 and 2.1.1). This method is neither globally stable, as evidenced by its output, nor fast enough for either testing scientific hypotheses, nor resolving the aforementioned inverse problem, described below. Furthermore, the use of “black box” simulation software, e.g. COMSOL, has obvious limitations, such as the lack of flexibility for modification or extension to a broader framework, like an inverse problem. By their very generality, such software is likely to be suboptimal for a specific task. Thus, the introduced numerical algorithm is not only an accurate simulation tool, but also an efficient direct solver serving a larger vision within the kinetic separation framework, as follows.

The kinetic separation system of convection–diffusion–reaction equations includes kinetic rate constants in the reaction term, which can be viewed as parameters that the solution functions of this system of partial differential equations depend on. The aim of the inverse problem is to find or approximate the parameters that induce an a priori given exact solution. If such a method can arbitrarily approximate an exact solution, then it can be applied to experimental data. The goal is to minimize an error function between the given solution and the approximating solutions, which must be generated for each set of test parameters as the optimization progresses. The generation of such parametric solutions is essentially akin to a function evaluation in an optimization iteration step, which must clearly be both efficient and accurate.

The determination of accuracy of a numerical solution is difficult in general, but for our model, it can be compared with an exact solution for a simplified case, derived in Section 2.2. The error is analyzed in Section 3.3, and the runtimes are compared in Section 3.2.

1.2. The physical model

The equations of our physical model can be deduced from the equations describing the conservation of mass with fast equilibrium of the reactants between the pores and the free volume in pressure-driven size-exclusion chromatography; this is the case of KSEC [3,6]. Below we show that they can also be deduced from the Nerst–Planck Equations, which express the conservation of mass of reactants moving under the influence of an electric field along a single spatial direction in a fluid medium; this is the case of KEC.

Define \( c = (c_1, \ldots, c_N) : \mathbb{R}_+^2 \rightarrow \mathbb{R}_+^N \) (where \( \mathbb{R}_+ := [0, +\infty) \)) as the spatiotemporal concentration of \( N \in \mathbb{N} \) convected ions over \((t, x)\) points, \( V : \mathbb{R}_+^2 \rightarrow \mathbb{R}_+^N \) as the velocities of their convection, \( D : \mathbb{R}_+^2 \rightarrow \mathbb{R}_+^N \) as their diffusion, \( R : \mathbb{R}_+^N \rightarrow \mathbb{R}_+^N \) as a reaction term between the ions, \( K \in \mathbb{N}^N \) as coefficients depending on electrical charges and hydrodynamic sizes of ions as well as fluid viscosity and having meaning and dimensions of ionic mobilities, and \( E : \mathbb{R}_+^2 \rightarrow \mathbb{R}_+^N \) as the electric fields influencing the motion of the ions, while denote with \( \cdot \) the Hadamard product of two vectors. To arrive at an intermediate form towards our equations, the Nerst–Planck equations [23] reduce to the following vector PDE

\[
\partial_t c + \partial_x (V \cdot c) = \partial_x (D \cdot \partial_x c) + R(c) + K \cdot \partial_x (E \cdot c)
\]

with appropriate initial and boundary conditions that ensure the existence and uniqueness of a concentration vector solution.

For the experimental model of KCE [5,22,15], the above equation reduces further due to certain features of the experimental setup. Specifically the fluid velocity \( V \in \mathbb{R}_+^N \) is constant, the diffusion coefficients are constant \( D \in \mathbb{R}_+^N \), and the electric field strength \( E \) is constant as well. Thus a constant vector \( v := V - K \cdot E \in \mathbb{R}_+^N \) is introduced, and its components can be thought of as a case of constant convection in a convection–diffusion equation. Therefore, they can be stated as

\[
\partial_t c + v \cdot \partial_x c = D \cdot \partial_x^2 c + R(c)
\]

again with appropriate initial and boundary conditions [25,20].

In the formulation of kinetic separation model, the reaction is reversible binding, giving the name Constant–Convection–Diffusion–Reversible–Binding–Reaction (CCDRBR) equations. In a simple reversible-binding reaction, there are three reactants and their concentration vector is denoted either as \( c = (L, T, C) : \mathbb{R}_+^2 \rightarrow \mathbb{R}_+^3 \) representing the experimental ligand, target, and complex [17]. The velocities and diffusion coefficients are denoted as \( v = (v_L, v_T, v_C) \) and \( D = (D_L, D_T, D_C) \) respectively. In the kinetic separation model with a reversible-binding reaction, the reaction term is

\[
R(c) = (-k_{on}LT + k_{off}C, -k_{on}LT + k_{off}C, k_{on}LT - k_{off}C) : \mathbb{R}_+^2 \rightarrow \mathbb{R}_+^3.
\]

Here \( k_{on}, k_{off} \in \mathbb{R}_+ \) are the rate constants of complex formation and dissociation respectively. In MASKE, the concentration of the second ion \( T \) is assumed to be constant, so the second equation is omitted and the reaction term becomes \( R(c) = (-k_{on}LT + k_{off}C, k_{on}LT - k_{off}C) \). The point of this simplification is to enable us to find an exact solution [17]. Another simplification for the same reason, is to keep all three concentration components and simplify the reaction term only as \( R(c) = (k_{off}C, k_{off}C, -k_{off}C) \), as in Section 2.2 [18].

1.3. Initial and boundary conditions

Since our goal is to give both a numerical and an exact solution to the above equations, the latter must be conveniently derivable from a Green function via function convolutions. So the initial and boundary conditions (IBC) of the CCDRBR equations are chosen to be idealizations of the initial concentration profiles measured in kinetic separation experiments.
The initial conditions for the CDDRBR equations specify the concentration profiles of the injected reactant plugs. A plug can be represented using various density functions \( \varrho : \mathbb{R} \to \mathbb{R}_+ \). \( \int_{-\infty}^{\infty} \varrho = 1 \). An identical substance amount is assumed within the plug – regardless of the density function used – to ensure that the amount of reacting molecules remains consistent for various injected concentration profiles. The total amount of molecules at concentration \( c(x) \) in a capillary of constant radius \( r \), at locations \( x \) within cylinders of infinitesimal height \( dx \), is given by the integral

\[
\int_0^\infty c(x) \pi r^2 \, dx.
\]

So for two concentration functions to give the same amount of molecules, the areas under them must be equal. To be consistent with our earlier work [15], this amount is set at \( \pi r^2 \hat{c} l \), which arises from the special case of rectangular concentration profiles of heights \( \hat{c} \in \mathbb{R}^N \) and plug length \( l > 0 \). To rescale a density function \( \varrho \) as another density, the proper transformation is \( (1/l)\varrho(x/l) \), which retains a unit area under the curve. To ensure the standard amount of molecules, the area under the initial concentration profile \( IC := c(0, \cdot) : \mathbb{R}_+ \to \mathbb{R}_+ \) must equal \( \hat{c} l \), implying the transformation \( \hat{c} \varrho(x/l) \).

This way our earlier table of initial and boundary conditions [15] can be generalized to arbitrary plug densities as shown in Table 1.

Note that \( x_{det} \) denotes the location of the experimental detector at the end of the capillary, and \( \vartheta \) denotes the Heaviside function

\[
\vartheta(x) = \begin{cases} 
1 & \text{if } x \geq 0 \\
0 & \text{otherwise.}
\end{cases}
\]

2. Solution of the equations

2.1. Numerical solution of the complete equations

2.1.1. Discretization

As a reminder, the CDDRBR vector PDE equation is

\[
\partial_t c + v \cdot \partial_x c = D \cdot \partial_{xx} c + R(c).
\]

The standard way to discretize the equations would be

\[
\frac{c(t + \Delta t, x) - c(t, x)}{\Delta t} + \frac{c(t, x + \Delta x) - c(t, x)}{\Delta x} = D \cdot \frac{c(t, x - \Delta x) - 2c(t, x) + c(t, x + \Delta x)}{\Delta x^2} + R(u)
\]

yielding the vector iteration scheme

\[
c(t + \Delta t, x) = \frac{D \Delta t}{\Delta x^2} \cdot c(t, x - \Delta x) + \left( 1 + \frac{v \Delta t}{\Delta x} - 2 \frac{D \Delta t}{\Delta x^2} \right) \cdot c(t, x) + \frac{D \Delta t}{\Delta x^2} \cdot \frac{v \Delta t}{\Delta x} \cdot c(t, x + \Delta x) + R(c(t, x)) \Delta t.
\]

This common scheme is unstable due to the fact that the coefficients can be negative for a sparse mesh, causing oscillations to emerge in this recursion, commonly referred to as instability [30,24,12]. The main issue is that the Péclet number arising from the presence of both convection and diffusion (or the ratio max(\( v \))/min(\( D \)) is too large (convection-dominated case), requiring a very dense mesh for ensuring stability. This kind of instability can be partly resolved for convection–diffusion equations by weighting the diffusion term as in streamline upwind Petrov–Galerkin methods [13,11,1,4]. Some programs have already been developed for modeling similar molecular migration behavior in other settings [28,26,8,16,27].
Instead of a general partial resolution, we may attempt to resolve the instability by virtue of constant convection in our equations. Petrov et al. and others [22,9,10] suggest improving stability by aligning the grid points with the directions of plug peak motion in spacetime, by choosing special individual step sizes $\Delta x_n = v_n \Delta t$. Upon further thought, we can eliminate the convection term entirely by first changing variables for each line of the equations as

$$y_n := x - v_n t, \quad \gamma_n(t, y_n) := c_n(t, x)$$

resulting in the transformed equations

$$\partial_t \gamma_n = D_n \partial^2 y_n \gamma_n + R_n(\gamma) \quad (n = 1, \ldots, N).$$

Then the iterations will be executed separately for each concentration variable, and we interpolate between the meshes to compute the reaction term

$$R_n(\gamma) = R_n(\gamma_1(t, y_1), \ldots, \gamma_N(t, y_N)) = R_n(c(t, x)).$$

We discretize the $(t, y_n)$-planes as $\Delta y_n := v_n \Delta t$ in order to align the $(t, y_n)$ grid points with the boundary conditions, which are now skewed in the transformed planes. So the iteration scheme becomes the following

$$\gamma_n(t + \Delta t, y_n) = \frac{D_n \Delta t}{\Delta y_n^2} \gamma_n(t, y_n - \Delta y_n) + \left(1 - \frac{2 D_n \Delta t}{\Delta y_n^2}\right) \gamma_n(t, y_n) +$$

$$+ \frac{D_n \Delta t}{\Delta y_n^2} \gamma_n(t, y_n + \Delta y_n) + R_n(\gamma) \Delta t.$$ 

The stability of the iteration is now ensured, since the $v$ terms in the earlier discretization are eliminated, so the condition for stability becomes $\Delta t > \max_0(2D_n/\Delta y_n^2)$ which is easy to satisfy for a large Péclet number

$$v_n \approx 10^{-3}, \quad D_n \approx 10^{-10} \Rightarrow \Delta t > 10^{-7}.$$ 

So with $t_{\text{max}} = 10^3$ s, this implies a limit of at most $10^{10}$ grid points in time (and corresponding $\Delta y_n$). If $D_n = 0$, then there is no limitation for refinement.

### 2.1.2. The solver algorithm

The algorithm presented below finds approximate values of the concentration vector solution $c$ of the CCRBR equations with $N \in \mathbb{N}$ lines, over a given grid $(t_i, x_j) \in \mathbb{R}^2$ $(i = 0, 1, \ldots, I, \quad j = 0, 1, \ldots, J)$ where $x_j = x_{\text{det}}$, with an equidistant temporal discretization $\Delta t$ and arbitrary values in space.

A different $x$-grid for each line of the equations is defined as $(t_i, x'_{n,j})$ where $x'_{n,j} := j v_n \Delta t$ $(j = 0, 1, \ldots, J_n)$ where $x'_{n,j} = x_{\text{det}}$ (by choosing an equidistant spatial discretization $\Delta x_n = x_{n,j+1} - x_{n,j} = v_n \Delta t$). The $y$-grid for the transformed equations is defined as $(t_i, y_{n,i,j})$ where $y_{n,i,j} := x'_{n,j} - v_n t_i$ $(j = 0, 1, \ldots, J_n)$ (implying that $\Delta y_n = \Delta x_n$ for all $i$).

Denote the sought concentration values by $c_{n,i,j} \approx c_n(t_i, x_{n,j})$, the values on the equidistant grid by $c_{n,i,j} \approx c_n(t_i, x'_{n,j})$, while the values on the transformed grids by $\gamma_{n,i,j} \approx \gamma_n(t_i, y_{n,i,j})$. Note that $\gamma_{n,i,j} = \gamma_{n,j}^i$, since $\gamma_n(t_i, y_n) = c_n(t, x)$ by definition, for $y_n = x - v_n t_i$.

The reaction vector must be evaluated at the same spacetime grid points, so for each line of the equation $n \in \{1, \ldots, N\}$ we interpolate the concentration values on the other spatial grids $\gamma_{n,i,j}$ $(k \neq n, \quad j = 0, 1, \ldots, J_k)$ onto the current $n$-th grid $(t_i, y_{n,i,j})$ $(j = 0, 1, \ldots, J_n)$, and denote the new values as $\gamma_{n,i,j} \approx \gamma_n(t_i, y_{n,i,j})$ (note that we can define $\gamma_{n,n,i,j} := \gamma_{n,i,j}$). The associated reaction values are denoted as

$$R_{n,i,j} := R_n(\gamma_{1,n,i,j}, \ldots, \gamma_{N,n,i,j}) \quad (j = 0, 1, \ldots, J_n).$$

The algorithm contains nested for loops, first according to time, then by each reactant. Therefore, it begins by evaluating the initial conditions from some initial condition function $c(0, x) = c(x)$. The left boundary condition is given by the function $\gamma_{n,i,0} \approx \gamma_n(t_i, y_{n,0,j}) = c_n(t_i, 0) = B_c(t_i)$ and the right boundary condition is of Neumann type $\partial_y \gamma_n(t_i, y_{n,i,J_n}) = 0$ implying that $\gamma_{n,i,J_n} = \gamma_{n,i,J_n-1}$. Note that we must also extrapolate the transformed concentration for $j = 1$ as

$$\gamma_{n,i,j-1} \approx \frac{\gamma_{n,i,j-2} + \gamma_{n,i,j}}{2} \Rightarrow \gamma_{n,i,j-2} \approx 2\gamma_{n,i,j-1} - \gamma_{n,i,j}.$$ 

Lastly, denote the iteration coefficients (a convex combination) as

$$A_n := \frac{D_n \Delta t}{\Delta y_n^2}, \quad B_n := 1 - 2A_n \quad (n = 1, \ldots, N).$$

The algorithm may call various kinds of interpolation subroutines. Our implementation uses cubic Hermite splines, resulting in an $O(1/t) = O(\Delta t)$ convergence in the $L^2$ error from the exact solutions, as demonstrated computationally in Section 3.3.
Algorithm 1 Numerical solver for the CCDRBR equations.

1: function NUMSOLCDR(v, D, R, t, x)
2:     for n = 1,...,N do
3:         for j = 0, 1,..., Jn do
4:             γn,0,j := ICn(x_n, j)
5:         end for
6:         c_n,0 := INTERPOLATE(γ_n, 0, x_n, x)
7:     end for
8:     for i = 0, 1,..., n - 1 do
9:         for n = 1,..., N do
10:            for k = 1,..., N, k ≠ n do
11:                γn,i,k := INTERPOLATE(γ_n,i,γ_n,i,n,γ_n,i,k)
12:            end for
13:            γn,i,n := BCn(t)
14:        end for
15:     end for
16:     for j = 1,..., Jn do
17:         R_n,i,j := R_n(t,y_n,i,j,γ_n,i,j)
18:         if j = 1 then
19:             γn,i+1,j := R_n(t,y_n,i,j,γ_n,i,j) + R_n,t,i,j,γ_n,i,j + R_n,γ_n,i,j,γ_n,i,j
20:         end if
21:     end for
22:     c_n,i+1 := INTERPOLATE(γ_n,i+1,γ_n,i,x_n, x)
23: end for
24: return c
25: end function

2.2. The exact solution for a simplification of NECEEM

We plan to explicitly solve a simplified case of the CCDRBR vector PDE equation

\[ \partial_t C + v \cdot \partial_x C = D \cdot \partial^2_x C + R(C) \]

with reaction function \( R(C) = (k_{off} C, -k_{off} C, k_{off} C) : \mathbb{R}_+^2 \rightarrow \mathbb{R}^3 \) where \( c = (L, T, C) \), via the method of fundamental solutions as demonstrated earlier for the case of a rectangular plug [18]. Hereby the exact solutions are derived for a plug represented by a Gaussian initial condition, defined via the following density function

\[ \mathcal{Q}_C(x) = \mathcal{Q}_C[\mu, \sigma^2](x) := \frac{1}{\sigma \sqrt{2\pi}} \exp \left( -\frac{(x - \mu)^2}{2\sigma^2} \right). \]

The NECEEM vector initial condition according to Section 1.3 is

\[ c(0, x) = IC(x) := \mathcal{Q}_C(x/l) \]

while the boundary conditions remain unspecified for now, for the sake of simplicity.

Introducing some fundamental solutions will aid us in our derivation of an explicit solution. It can be shown by substitution that the equation

\[ \partial_t F + v \cdot \partial_x F - D \partial^2_x F = -kF \]

with the initial condition \( F(0, x) = \delta(x) \), is solved by

\[ F[k](t, x) := \theta(t) e^{-k t} \mathcal{Q}_C[vt, 2Dt](x). \]

On the other hand, the fundamental solution satisfying

\[ \partial_t F + v \partial_x F - D \partial^2_x F = \delta \]

with a spatiotemporal Dirac delta function, is coincidentally the function \( F[0] \).

Thus to find the third concentration component \( C \) for some initial condition function \( C(0, x) = IC(x) \), we must convolve it with the first fundamental solution above, resulting in

\[ C(t, x) = (IC \ast F_C[k_{off}](t, \cdot))(x) \]

where \( F_C \) is defined similarly as above, with parameters \( v_C \) and \( D_C \).

In order to get the other two concentration components \( L \) and \( T \), we break them up into an equilibrium term (the solution of the \( k = 0 \), \( F(0, x) = IC(x) \) case above) and a dissipation term (the solution with right-hand side \( k_{off} C \), resulting in...
\[ L(t, x) = (ICL * FL(t, \cdot))(x) + k_{off}(C * FL(t, \cdot))(t, x) \]
\[ T(t, x) = (ICT * FT(t, \cdot))(x) + k_{off}(C * FT(t, \cdot))(t, x). \]

Since the above formulas include several function convolutions (single and double integrals), it is desirable to simplify our derivation with the following well-known identity
\[ \varphi_G[\mu_1, \sigma_1^2] * \varphi_G[\mu_2, \sigma_2^2] = \varphi_G[\mu_1 + \mu_2, \sigma_1^2 + \sigma_2^2]. \]

Next, notice that for some \( \mu_0, \sigma_0 > 0 \) and \( \mu := \mu_0l, \sigma := \sigma_0l \) we have
\[ IC(x) = \bar{c} \varphi_G[\mu_0, \sigma_0^2](x/l) = \bar{c}l \varphi_G[\mu, \sigma^2](x). \]

Employing the above general solution formulas for this special initial condition, we get
\[ C(t, x) = (IC * FC[t_{off}](t, \cdot))(x) = I\bar{C} \left( \varphi_G[\mu, \sigma^2] * \left( \vartheta(t)e^{-k_{off}C}vLt, 2DCt \right) \right)(x) = \]
\[ = I\bar{C} \vartheta(t)e^{-k_{off}C} \varphi_G[\mu + vLt, \sigma^2 + 2DCt](x). \]

With the above, we may now derive the other concentration components as follows
\[ L_{eq}(t, x) = (ICL * FL(t, \cdot))(x) = I\bar{L} \varphi_G[\mu + vLt, \sigma^2 + 2DLt](x) \]
\[ L_{dis}(t, x) = k_{off}(C * FL(t, \cdot))(t, x) = k_{off} \int_{-\infty}^{+\infty} (C(\tau, \cdot) * FL(t - \tau, \cdot))(x) \, d\tau = \]
\[ = k_{off}I\bar{C} \int_{-\infty}^{+\infty} \vartheta(\tau)e^{-k_{off}C} \varphi_G[\mu + vL\tau, \sigma^2 + 2DC\tau] \vartheta(t - \tau) \varphi_G[vLt - (vC - vL)\tau, \sigma^2 + 2DL(t - \tau)](x) \, d\tau = \]
\[ = k_{off}I\bar{C} \int_{0}^{t} e^{-k_{off}C} \varphi_G[\mu + vLt + (vC - vL)\tau, \sigma^2 + 2DLt + 2(DC - DL)\tau](x) \, d\tau \]
\[ L = L_{eq} + L_{dis} \]

and \( T_{eq}, T_{dis}, T \) are defined similarly.

Despite the extensive simplification of the exact solutions for the Gaussian case above – otherwise containing even more computationally expensive convolution integrals – we will show in Section 3.2 that somewhat surprisingly, the numerical method of the previous section is much faster according to our computational tests. Not only is it faster, but it is also capable of solving the CCRBR equations for any reaction function and initial–boundary conditions, with a reasonable error according to Section 3.3.

3. Computational results

3.1. The direct solver package

The numerical solver algorithm of Section 2.1.2 has been implemented in MATLAB for the KCE equations, though can be easily generalized to more reactants and any reaction mapping. The package is available on GitHub [29]. Currently the package contains only the direct solver for KCE, but the inverse solver built on it is also under development, serving our research described in the introduction.

The direct solver sub-package, contains an implementation of the numerical algorithm that is functional for all initial and boundary conditions, and plug types described in Section 1.3. Exact solutions have only been derived and implemented for the simplified NECCEEM case in Section 2.2 with Gaussian and rectangular plug types [18], and for MASKE [17] with any plug type.

Therefore, in order to compare the computational runtimes and errors between the numerical and exact solutions, we are restricted to either of the above two simplified methods. We chose simplified NECCEEM with a Gaussian plug for the analysis below, since its equations are closer to the full kinetic separation equations than those of MASKE, consisting of only two lines. Furthermore, the solution formulas can be minimized computationally as much as possible, by eliminating expensive convolution integrals which arise for other density functions. This way we achieve a lower bound on the computational complexity of the exact case, which can be compared with that of the numerical one. Specifically, three single convolution integrals are eliminated for \( C, L_{eq}, T_{eq} \) and two double convolution integrals are reduced to single ones for \( L_{dis}, T_{dis} \), as shown in Section 2.2.

The inverse problem of our research requires fast and accurate generation of solutions to the CCRBR equations for varying \( k_0, k_{off} \) values, in order to model experimental signals. Thus, we analyze our direct solver in terms of both runtime and error below.
3.2. Runtime analysis

In order to compare the runtime of our numerical solver (Section 2.1.2) to that of computing the exact solutions of simplified NECEEM (Section 2.2), we ran the solver for each temporal mesh size \( I \) five times, and plotted the average of runtimes in Fig. 1. We are only interested in varying \( I \) and not \( J \) (the spatial mesh size), because the experimental signals are available only at the detector location \( x_{\text{det}} \), restricting the inverse problem to this single location in space. Test runs were done with an AMD A8-5550M processor, 8.00 GB of RAM, 64-bit Windows 10, and MATLAB R2016a, with the parameters

\[
\begin{align*}
    k_{\text{on}} &= 3500 \text{ m}^3/\text{mol}, \\
    k_{\text{off}} &= 0.035, \\
    v &= (3.3, 5, 4) \times 10^{-3} \text{ m}^2/\text{Vs}, \\
    D &= (7, 7, 7) \times 10^{-11} \text{ m}^2/\text{s}, \\
    \bar{c} &= (1.86, 16.9, 3.14) \times 10^{-6} \text{ mol/m}^3, \\
    l &= 0.005 \text{ m}, \\
    t_{\text{max}} &= 90 \text{ s}, \\
    x_{\text{det}} &= 0.2 \text{ m}.
\end{align*}
\]

The runtimes seem to closely follow quadratic and linear relationships for the numerical and exact solvers, respectively, with high coefficients of determination \( R^2 \). This is expected for the exact solver, which simply evaluates the formulas of Section 2.2. The quadratic polynomial for \( T_N(I) \) makes sense, since the algorithm of Section 2.1.2 must necessarily solve for all the nodes on the spacetime mesh to provide the values at the only relevant location \( x_{\text{det}} \). The transformed spatial meshes all depend on the temporal mesh linearly \( \Delta y_n = v_n \Delta t \), implying that the spacetime mesh sizes must depend quadratically on \( I \).

It is important to observe that the ratio of runtimes \( r(I) \) decreases for increasing \( I \), and follows a power law with an exponent of \(-0.761\) (though the ratio of a linear and a quadratic polynomial should asymptotically follow a hyperbola with an exponent of \(-1\)). This implies that the gain in runtime using the numerical solver is much larger for lower mesh sizes, the difference ranging between 1–2 orders of magnitude below \( I = 2000 \). For our end goal of an efficient inverse solver, a mesh size of \( 1000 \leq I \leq 2000 \) is sufficient.

Thus, we have demonstrated that surprisingly the numerical solver has a significantly lower runtime than even an optimally low runtime of the exact solution formulas. For this to be relevant, the error between the solutions must be reasonable.

3.3. Error analysis

The exact solution formulas for simplified NECEEM (Section 2.2) provide a standard of comparison for the numerical solution values, since the formulas for \( C, L_{\text{eq}}, T_{\text{eq}} \) are exact, while the formulas for \( L_{\text{eq}}, T_{\text{eq}} \) can be approximated accurately with some quadrature method (we used the \texttt{integral} function of MATLAB, with a \texttt{RelTol} value of \( 10^{-6} \)).

To show that the numerical solutions generated by Algorithm 1 are “reasonable”, we demonstrate that the \( L^2 \)-error from the exact values vanishes with an increasing temporal mesh size \( I \). Fig. 2 shows that the error decreases for increasing temporal mesh size according to similar power laws for the three concentration components. Taking the weighted average of the three exponents according to the coefficients of determination \( (R^2 \text{ values}) \) gives \(-0.9798 \). Thus we conjecture that all three error components are inversely proportional to the temporal mesh size for large \( I \), meaning they are precisely of order \( O(1/I) = O(\Delta t) \).

The convergence of Algorithm 1 for increasing mesh size is self-evident from its design. Clearly, the error in the solution \( \Delta u \) arising from the derivative approximations, and the multimesh interpolation \( \Delta u \) vanishes as the spacetime mesh size increases. For an infinitesimally fine mesh, the error is necessarily zero. Thus the above comparison in Fig. 2 – between the exact and simulated solutions in the simplified case – is not intended to be a proof of convergence, but merely a
rate analysis of the de facto vanishing of the error with respect to mesh size. It gives an idea of how “reasonable” this convergence is.

Nevertheless, this computational error analysis of the simplified system bears relevance to the original equations. The two differ in the reaction terms: (1) original: $-k_{on}LT + k_{off}C$; (2) simplified: $k_{off}C$, but the solutions differ negligibly. As illustrated in Fig. 3 for the simplified solution, the peaks are separated in a way that the following hold

\[ LT \gg 0 \text{ when } C \gg 0; \quad LT \approx 0 \text{ when } C \approx 0; \quad LT \ll C \text{ when } C \gg 0. \]

Consequently, the following approximations hold

\[ -k_{on}LT + k_{off}C \approx 0 \text{ when } C \approx 0; \quad -k_{on}LT + k_{off}C \approx k_{off}C \text{ when } C \gg 0. \]

So we may conclude heuristically, that the above rate analysis of the error in the simplified case, is closely aligned with the original.
4. Concluding remarks

4.1. Summary and future directions

A stable algorithm has been introduced for the efficient generation of accurate solutions to a set of convection–diffusion equations, with constant convection velocities and a non-linear reaction term describing a reversible binding reaction. Stability has been demonstrated algebraically, due to the change of variables to new spatial meshes, for each line of this system of partial differential equations. Both efficiency and accuracy have been demonstrated as well.

The direct solver package corresponding to this work [29] may be extended in two potential directions: the implementation of new initial and boundary conditions, perhaps with a generic classification system; or the derivation of new exact solutions for either other simplifications of the original equations, or perhaps the explicit solution of the complete equations. According to our runtime analysis, however, the computation of closed form solutions is not likely to be more efficient, even in an extremely simple Gaussian case which gives a lower bound on computational complexity, implying the permanent significance of our solver.

The relevance of this algorithm to our further work on an experimental inverse problem has also been highlighted. Indeed, this is the direction our solver package is being extended in, now enabled by the fast computation of parametrized solutions with an error measured from an experimental signal, which is minimized via an optimization algorithm.

4.2. Experimental remarks

The presented discretization method was developed primarily for the KCE and KSEC equations [15,6], which describe only the migration of, and interaction between three species: the ligand L, the target T, and the complex C. The KCE equations do not incorporate a variety of other “background” species and reactions – such as components of the acid–base equilibration in the background electrolyte in KCE, which occur during electrophoresis of these three species. In practice, our method utilizes experimentally measured velocities of the three species, and plug lengths corresponding to them. Such parameters are sufficient for the description of the relevant migration–reaction phenomena, only if the respective experiment is planned and carried out properly. Most importantly, in KCE experiments, the concentrations of the analytes should be well below the concentrations of the ions in the background electrolyte. In turn, in KSEC experiments, the equilibration between the pores and the free volume should be fast. Beneficially, these conditions are easily satisfied for KCE and KSEC, as analyte concentrations are typically at least four orders of magnitude below the background electrolyte concentration and the sub-nanometer size of pores allows fast equilibration between the pores and the free volume.

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References
