Modeling Analytical Ultracentrifugation Experiments with an Adaptive Space-Time Finite Element Solution of the Lamm Equation

Weiming Cao* and Borries Demeler†
*Department of Applied Mathematics, The University of Texas at San Antonio, Texas; and †Department of Biochemistry, University of Texas Health Science Center at San Antonio, San Antonio, Texas

ABSTRACT Analytical ultracentrifugation experiments can be accurately modeled with the Lamm equation to obtain sedimentation and diffusion coefficients of the solute. Existing finite element methods for such models can cause artifactual oscillations in the solution close to the endpoints of the concentration gradient, or fail altogether, especially for cases where \( \omega^2 D \) is large. Such failures can currently only be overcome by an increase in the density of the grid points throughout the solution at the expense of increased computational costs. In this article, we present a robust, highly accurate and computationally efficient solution of the Lamm equation based on an adaptive space-time finite element method (ASTFEM). Compared to the widely used finite element method by Claverie and the moving hat method by Schuck, our ASTFEM method is not only more accurate but also free from the oscillation around the cell bottom for any \( \omega^2 D \) without any increase in computational effort. This method is especially superior for cases where large molecules are sedimented at faster rotor speeds, during which sedimentation resolution is highest. We describe the derivation and grid generation for the ASTFEM method, and present a quantitative comparison between this method and the existing solutions.

INTRODUCTION

In recent years, analytical ultracentrifugation has seen a resurgence as a method of choice for characterizing solution behavior of biological macromolecules and macromolecular assemblies. Sophisticated numerical analysis coupled with modern digital data acquisition technology offers a wealth of macromolecular information from this technique. The sedimentation process of a solute in an analytical ultracentrifugation cell is described by the Lamm equation (1),

\[
\frac{\partial C}{\partial t} = \frac{1}{r} \frac{\partial}{\partial r} \left( D \frac{\partial C}{\partial r} - s \omega^2 r C \right), \quad m < r < b, t > 0, \tag{1}
\]

where \( C(r, t) \) is the concentration, \( s \) and \( D \) are the sedimentation and diffusion coefficients, and \( \omega \) is the angular velocity. The terms \( m \) and \( b \) are the radii of the meniscus and bottom of the cell.

Using a nonlinear least-squares fitting algorithm, data from sedimentation velocity and approach-to-equilibrium experiments can be fitted to finite element solutions of the Lamm equation, and sedimentation and diffusion coefficients, partial concentrations, and to some degree, even equilibrium constants, can be obtained at high resolution. A number of approaches for solving the inverse problem of fitting moving boundary data to finite element solutions of the Lamm equation have been developed and implemented in software packages. Direct fitting of boundary data to fixed-mesh finite element solutions was originally proposed by Claverie et al. (2) and Todd and Haschemeyer (3). In the fixed-mesh approach, radial mesh points are uniformly distributed between the meniscus and the bottom of the cell. Software based on this solution was later developed by Demeler and Saber (4) and Schuck et al. (5). The approach of Stafford and Sherwood (6) fits time-difference data from experimental scans to time-difference data simulated with a fixed-mesh finite element solution. However, the fixed-mesh finite element method suffers from an artifactual broadening of the sedimenting boundary (so-called numerical diffusion). When such solutions are used to fit experimental data, the fitted diffusion coefficients tend to be smaller than the actual diffusion coefficients, especially when the concentration gradient is steep. Later, Schuck introduced an improved finite element solution, the so-called moving hat method (7). This method uses a moving frame of reference and discretizes the Lamm equation under the moving frame by using the standard finite element method. A similar idea relying on distorted grids in finite difference solutions of the Lamm equation has been applied in the earlier developments of sedimentation simulation techniques; see Cox and Dale (8) and references therein. The moving grid employed in the moving hat method is spaced exponentially in the radial direction and shifted at the speed of sedimentation. The moving hat method improves the accuracy of the numerical solution substantially compared to the fixed-mesh approach by Claverie. However, there are two issues not well addressed by the moving hat method. First, even though the gridpoints at the meniscus and at the bottom of the cell have to stay fixed, the moving reference frame used in the moving hat method would require these points to move like all other points in the grid. Therefore, the finite element discretization cannot be applied at the meniscus and at the bottom in the moving hat method. Schuck addressed this singularity by manually adjusting the values of the solution at the bottom.
and at the point next to the meniscus. Such post-processing is not based on the differential equation itself and may result in loss of accuracy. The second problem with the moving hat method is related to the fact that the grid spacing increases exponentially from the meniscus to the bottom, resulting in low radial mesh point density at the bottom of the cell where concentration changes can be very large, thus lacking the required resolution. It has been pointed out (4,7) that a lack of sufficient radial resolution at the bottom of the solution column results in oscillations in the finite element solution, especially for experimental conditions where $\omega^2/D$ is large. Such conditions are encountered when large molecules such as DNA molecules $>1$ kb, multi-enzyme complexes, and large assemblies like chromatin or virus particles are sedimented at moderately high rotor speed. Indeed, if the number of radial mesh points is not large enough, both the Claverie approach and the moving hat method will fail altogether.

Here we present an entirely new finite element method, termed the adaptive space-time finite element method (ASTFEM), for the numerical solution of the Lamm equation that incorporates the advantages introduced by the moving reference frame, and properly addresses the remaining issues in the moving hat method. In this method the Lamm equation is simultaneously discretized in the radial and the time domain (hence the name of space-time finite element method). This space-time discretization avoids the difficulty caused by the singularity of the moving grid at the meniscus and at the bottom in the moving hat method. To eliminate the oscillations around the cell bottom, we implement an adaptive grid locally in a very narrow region next to the cell bottom. The length of this narrow region is proportional to $D/(\omega^2)$. The number of the grid points placed in the narrow region is $\ln(D/(\omega^2))$, which increases very slowly with respect to $\omega^2/D$. This fine grid distribution around the cell bottom effectively eliminates the oscillation of the numerical solution. To minimize the numerical diffusion, we use the same moving mesh idea as in Schuck’s moving hat method in the region away from the bottom. The length of this region typically ranges between 90 and 99% of the cell length (see below). As a result, the ASTFEM enjoys the same accuracy with respect to numerical diffusion as the moving hat method, but avoids the inaccuracies due to the singularities at the meniscus and bottom of the cell, and completely eliminates oscillations at the bottom of the cell. At the same time is that the mass conservation of the Lamm equation is guaranteed automatically without the need for any post-processing. Overall, we conclude that the ASTFEM method proposed in this article is an accurate, efficient and robust method for the Lamm equation for sedimentation experiments. Furthermore, the concept of the space-time finite element discretization on adaptive moving grids can be extended to multiple interacting and self-association systems or models with concentration dependent sedimentation and diffusion coefficients.

**THE ADAPTIVE SPACE-TIME Finite Element Solution of the Lamm Equation**

Let $\delta t$ represent the size of the time-step, and let $t_n = n\delta t$ be the $n^{th}$ time-step. We multiply both sides of the Lamm equation by an arbitrary function $v(r,t)$ and integrate the result over the space-time slab $[m, b] \times [t_n, t_{n+1}]$. It follows from integration by parts

$$\int_{t_n}^{t_{n+1}} \int_m^b \left( \frac{\partial C}{\partial t} v + D \frac{\partial C}{\partial r} \frac{\partial v}{\partial r} - \omega^2 r^2 \frac{\partial^2 C}{\partial r^2} \right) \, dr \, dt = 0. \quad (2)$$

This is called the weak form of the Lamm equation. Any solution to the Lamm equation must satisfy Eq. 2, and any smooth function $C(r, t)$ satisfying Eq. 2 must also be a solution of the Lamm equation. We will derive the finite element solution based on this weak formulation.

To define an approximate solution to Eq. 2, we first divide the space-time slab $[m, b] \times [t_n, t_{n+1}]$ into a number of space-time elements. Each element is either a triangle or a quadrilateral. Let $N$ be the total number of points used in the $r$-direction, and suppose the grid points $r_j, 1 \leq j \leq N$ are already given (see next section for the distribution of the grid points). We connect $(r_j, t_n)$ to $(r_{j+1}, t_{n+1})$ for all $j = 1, \ldots, N - 1$. Then the slab $[m, b] \times [t_n, t_{n+1}]$ is divided into $N - 1$ quadrilaterals and two triangles (one at each end). For each $j$, let $c^n_j$ and $c^{n+1}_j$ be the approximate values of $C(r, t)$ at the grid points $(r_j, t_n)$ and $(r_{j+1}, t_{n+1})$, respectively. We define a continuous function $c(r, t)$ to approximate the exact solution $C(r, t)$ as follows.

Consider a quadrilateral element $K^n_j$ with vertices $(r_j, t_n), (r_{j+1}, t_n), (r_{j+2}, t_{n+1}), (r_{j+1}, t_{n+1})$. Let $\hat{K} = [0,1] \times [0,1]$ be the standard element in an auxiliary $\xi \eta$-coordinate system. Define the mapping $F^n_j : \hat{K} \rightarrow K^n_j$ as

$$\begin{align*}
\begin{cases}
r = r_j (1 - \xi)(1 - \eta) + r_{j+1} \xi (1 - \eta) + r_{j+1}(1 - \xi) \eta + r_{j+2} \xi \eta \\
t = t_n + \delta t \eta.
\end{cases}
\end{align*} \quad (3)$$

the computational costs for all three methods are essentially the same: At each time-step one needs to solve a triangular linear algebraic system of equations with a fixed coefficient matrix. Another important feature of the ASTFEM approach (see Fig. 1). This mapping will be used to introduce the finite element solutions for all $r$ and $t$ on $K^n_j$. Now, given four values $c^n_j, c^{n+1}_j, c^{n+1}_{j+1},$ and $c^{n+1}_{j+2}$, we define a bilinear function $\hat{c}(\xi, \eta)$ on the standard element $\hat{K}$ as
The Jacobian
\(v\) from \(t_1\) to \(t_2\) is into a quadrilateral element as \(jh\) \(v\) \(t\)

The finite element solution on \(t\) is defined by transforming a bilinear function on \(K\) with mapping \(F\):

\[
\hat{c}(\xi, \eta) = c^0_{j,1}(1 - \xi)(1 - \eta) + c^0_{j,2}(1 - \eta) + c^{n+1}_{j,2} \xi \eta + c^{n+1}_{j,1}(1 - \eta) \eta.
\]

(4)

It follows that \(\hat{c}\) takes the values \(c^0_{j,1}, c^0_{j,2}, c^{n+1}_{j,2}, c^{n+1}_{j,1}\) at the four vertices of \(K\), respectively. The approximate function \(c(r, t)\) on element \(K^n\) is defined as the "pull-back" of \(\hat{c}(\xi, \eta)\) under the inverse mapping of \(F\). More precisely, let

\[
\begin{bmatrix}
\xi \\
\eta
\end{bmatrix} = G^{-1}_j \begin{bmatrix}
r \\
t
\end{bmatrix} = \frac{r - r_j - (r_{j+1} - r_j) \eta}{(r_{j+1} - r_j) + (r_{j+2} - 2r_{j+1} + r_j) \eta} \begin{bmatrix}
r - t_j \\
t - t_j
\end{bmatrix}
\]

be the inverse mapping of \(F\). Then we define for \((r, t)\) in \(K^n\),

\[
c(r, t) = \hat{c}(\xi, \eta) = \hat{c}\left(c_j\left(\begin{bmatrix}
r \\
t
\end{bmatrix}
\right)\right),
\]

and \(c(r, t)\) takes the values \(c^0_{j,1}, c^0_{j,2}, c^{n+1}_{j,2}, c^{n+1}_{j,1}\) at the vertices \((r_j, t_0), (r_{j+1}, t_0), (r_j, t_{n+1}),\) and \((r_{j+1}, t_{n+1})\), respectively.

To define \(c(r, t)\) on the triangular element \(K^0\) with vertices \((r_1, t_0), (r_2, t_0),\) and \((r_1, t_{n+1})\), we use the standard triangular element \(\hat{T}\) with vertices \((\xi, \eta), 0 \leq \xi, \eta \leq 1, \xi + \eta \leq 1\) (see Fig. 2). The mapping \(F^0\) from \(K^0\) to \(K^n\) is

\[
\begin{align*}
r &= r_1(1 - \xi - \eta) + r_2 \xi + r_1 \eta \\
t &= t_0 + \delta t \eta
\end{align*}
\]

Define a linear function \(\hat{c}(\xi, \eta)\) on \(\hat{T}\) as

\[
\hat{c}(\xi, \eta) = c^0_{j,1}(1 - \xi - \eta) + c^{n+1}_{j,2} \xi + c^{n+1}_{j,1} \eta,
\]

and define the approximate function \(c(r, t)\) on \(K^0\) as the "pull-back" of the above under the inverse mapping of \(F^0\).

Similarly, on the triangular element \(K^n\) at the right end of the space-time slab, we can introduce the mapping \(F^0\) from \(\hat{T}\) to \(K^n\) and define the approximate function \(c(r, t)\) at time \(t_0\), we now derive a system of linear algebraic equations for the unknown values \(c^{n+1}_{j,1}, 1 \leq j \leq N\) at the next time-step \(t_{n+1}\). We ask for the weak form (Eq. 2) to hold for all test functions \(v_j\) defined in the last subsection, namely, that the approximate function \(c(r, t)\) satisfies

\[
\int_{t_0}^{t_1} \int_m^b \left( \frac{\partial c}{\partial t} v_j + D r \frac{\partial c}{\partial r} \frac{\partial v_j}{\partial r} - s a r^2 \frac{\partial c}{\partial r} \frac{\partial v_j}{\partial r} \right) dr dt = 0,
\]

(6)

Since for each \(j\) the test function \(v_j\) is non-zero only in \(K^n\) and \(K^{n+1}\) \(v_j\) is non-zero only in \(K^n\), the above equation can be reduced to

\[
\left( \int_{K^0} + \int_{K^{n+1}} \right) \left( \frac{\partial c}{\partial t} v_j + D r \frac{\partial c}{\partial r} \frac{\partial v_j}{\partial r} - s a r^2 \frac{\partial c}{\partial r} \frac{\partial v_j}{\partial r} \right) dr dt = 0.
\]

(7)

To evaluate the integrals over \(K^n\), we change the variables from \((r, t)\) to \((\xi, \eta)\) by the mapping from Eq. 3 so that the integrals will be over the standard element \(K\). The Jacobian \(J\) of the coordinate transform is

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where by Eq. 3,
\[\frac{\partial r}{\partial \xi} = (r_{j+1} - r_j)(1 - \eta) + (r_{j+2} - r_{j-1})\eta,\]
\[\frac{\partial r}{\partial \eta} = (r_{j+1} - r_j)(1 - \xi) + (r_{j+2} - r_{j+1})\xi,\]
\[\frac{\partial t}{\partial \xi} = 0,\]
\[\frac{\partial t}{\partial \eta} = \delta t.\]  

By the fact that the Jacobian \(\frac{\partial(\xi, \eta)}{\partial(r, t)}\) for the inverse transform is equal to the inverse matrix of \(J\), we have
\[\frac{\partial \xi}{\partial t} = 1, \quad \frac{\partial \eta}{\partial t} = 0,\]
\[\frac{\partial \xi}{\partial r} = \frac{\partial \xi}{\partial t} \left(\frac{\partial t}{\partial \eta}\right) = 1.\]

Now it follows from the chain rule of derivatives that
\[\frac{\partial c}{\partial t} = \frac{\partial c}{\partial \xi} \frac{\partial \xi}{\partial t} + \frac{\partial c}{\partial \eta} \frac{\partial \eta}{\partial t},\]
\[= \frac{\partial c}{\partial \xi} + \frac{\partial c}{\partial \eta} \delta t,\]
and by Eq. 4 that
\[\frac{\partial c}{\partial \xi} = (c^{\text{p+1}} - c^{\text{p}})(1 - \eta) + (c^{\text{p+1}} - c^{\text{p+1}})\eta,\]
\[\frac{\partial c}{\partial \eta} = (c^{\text{p+1}} - c^{\text{p}})(1 - \xi) + (c^{\text{p+1}} - c^{\text{p+1}})\xi.\]  

For the second integral in Eq. 7, we note the fact that
\[v_j = 1 - \xi \text{ on } K_n^a,\]
\[\frac{\partial v_j}{\partial r} = \frac{\partial v_j}{\partial \xi} \frac{\partial \xi}{\partial r} + \frac{\partial v_j}{\partial \eta} \frac{\partial \eta}{\partial r} = -1 \frac{\partial \xi}{\partial r} = -1.\]

Then the second integral in Eq. 7 can be reduced to
\[\iint_{K_n^a} \frac{\partial c}{\partial t} \frac{\partial v_j}{\partial r} \sqrt{\gamma} \, dr dt = -\delta t D \int_0^1 \int_0^1 \frac{\partial c}{\partial \xi} \frac{\partial v_j}{\partial \xi} \sqrt{\gamma} \, d\xi d\eta,\]
\[= M_{21} \left( c^{\text{p+1}} - c^{\text{p}} \right) + M_{22} \left( c^{\text{p+1}} - c^{\text{p+1}} \right),\]
where
\[M_{21} = -\delta t D \int_0^1 \int_0^1 (1 - \xi)(1 - \eta) r \, d\xi d\eta,\]
\[M_{22} = -\delta t D \int_0^1 \int_0^1 \eta \, d\xi d\eta.\]

The third integral in Eq. 7 can be changed into
\[\iint_{K_n^a} \sqrt{\gamma} \frac{\partial v_j}{\partial r} r \, dr dt = -\delta t \sqrt{\gamma} \int_0^1 \int_0^1 r \, d\xi d\eta,\]
\[= M_{31} c^{\text{p+1}} + M_{32} c^{\text{p+1}} + M_{33} c^{\text{p+1}} + M_{34} c^{\text{p+1}} + M_{35} c^{\text{p+1}} + M_{36} c^{\text{p+1}},\]
where
\[M_{31} = -\delta t \sqrt{\gamma} \int_0^1 \int_0^1 (1 - \xi)(1 - \eta) r^2 \, d\xi d\eta,\]
\[M_{32} = -\delta t \sqrt{\gamma} \int_0^1 \int_0^1 \xi (1 - \eta) r^2 \, d\xi d\eta,\]
\[M_{33} = -\delta t \sqrt{\gamma} \int_0^1 \int_0^1 (1 - \eta) \eta^2 r^2 \, d\xi d\eta,\]
\[M_{34} = -\delta t \sqrt{\gamma} \int_0^1 \int_0^1 \xi r^2 \, d\xi d\eta.\]

The above formulas cover all the integrals in Eq. 7 over \(K_n^a\). We can deal with the other integrals in Eq. 7 over element \(K_n^b\), \(K_n^c\), \(K_n^d\), \(K_n^e\), \(K_n^f\), \(K_n^g\) similarly. For the two triangular elements \(K_n^a\) and \(K_n^b\) at the end of the space-time slab, we need to use the mapping \(F_0^s\) and \(F_0^t\) defined in Eq. 5 to change the integrals into those over the standard triangle \(T\). We skip the details of the formulas, since they can be worked out in the same way as for the quadrilateral elements.

By putting all these integrals into Eq. 6, we obtain a system of linear algebraic equations
\[M_i C^{\alpha+1} = M_0 C^n,\]
where $C^n = [c_1^n, c_2^n, \ldots, c_N^n]^T$ is the vector of the known approximate values of $C$ at time $t_n$, and $C^{n+1} = [c_1^{n+1}, c_2^{n+1}, \ldots, c_N^{n+1}]^T$ is the vector of the unknown approximate values of $C$ at time $t_{n+1}$. $M_1$ and $M_0$ are the coefficient matrices assembled from the above integral formulas. Note that both of these two matrices are independent of the time-level $n$. Therefore, they need to be calculated only once for the entire simulation process. Also, since both $M_1$ and $M_0$ are tridiagonal, the system of equations in Eq. 14 can be easily solved. If we use the Gaussian elimination method to solve Eq. 14, then the total number of arithmetic operations needed is only $13N$ for each time-step. If we perform the LU-decomposition on $M_1$ (which costs only $5N$ operations) and store it in memory, then we only need 11N operations at each time-step.

Another important feature of the above ASTFEM scheme is that the mass conservation for the Lamm equation is guaranteed automatically without any post-processing. If we sum up the expressions in Eq. 6 for all $j = 1, 2, \ldots, N$, and notice that

$$
\sum_{j=1}^{N} (v_j r, t) = 1, \quad \text{on} [m, b] \times [t_n, t_{n+1}],
$$

we have

$$
\int_{t_n}^{t_{n+1}} \int_{m}^{b} \frac{\partial c}{\partial t} dr dt = 0,
$$

which implies the conservation of mass

$$
\int_{m}^{b} rc(r, t_{n+1}) dr = \int_{m}^{b} rc(r, t_n) dr.
$$

**ADAPTIVE GRIDS**

In this section we discuss the selection of an ideal distribution of grid points used for the ASTFEM approach described in the last section. First, we recall the behavior of the concentration function $C(r, t)$ in a typical velocity experiment. A notable feature is the rapid buildup of the concentration at the bottom of the cell. This sharp increase can cause numerical oscillations if insufficient radial grid points are used to describe the transition. It is well known that all existing numerical solutions for the Lamm equation, including Claverie’s classical finite element method and Schuck’s moving hat method, exhibit oscillations at the bottom of the cell if the step size is not small enough. This oscillation happens especially when the sedimentation speed $s$ or angular velocity $\omega$ is large or the diffusion coefficient $D$ is small. The oscillations can propagate back toward the meniscus and even cause a complete failure of the computation. Although an increase of the number of grid points will correct this problem, this will also increase the computational cost. Furthermore, a rule to determine an appropriate grid spacing for the fixed-mesh or moving hat method does not exist, and the user is forced to experiment to avoid such oscillations.

Our first aim in the application of adaptive grids is to increase the resolution locally around the cell bottom. The second goal for adaptive grids is to minimize the numerical diffusion typically accompanied with a discretization of sedimentation-diffusion equations. This is essential since numerical diffusion may smooth out the steep boundary of $C$ and predict a larger diffusion coefficient or smaller molecular weight. It is well known in the numerical analysis community (and as observed by Schuck; see (7)) that the numerical diffusion introduced can be greatly reduced by aligning the space-time element with the sedimentation speed (9). Under the moving frame, the Lamm equation is essentially free of numerical diffusion.

Based on this observation, we will define a grid that aligns with the sedimentation speed in most regions of the cell similar to the implementation in the moving hat method. At the same time, we will maintain a very small step size in the narrow region around the bottom. To this end, we divide the interval $[m, b]$ into three different regions: a regular region, a steep region, and a transition region between the steep and regular regions. The grid in the regular region is generated comparable to the grid in the moving hat method. The steep region at the bottom of the cell consists of closely spaced grid elements (the exact spacing is explained below), and the transition region between the two consists of elements whose grid spacing changes gradually from one end to the other to produce a smooth transition of the overall grid spacing.

First we determine the width of the steep region by using the equilibrium condition of the Lamm equation, which is given by

$$
C(r) = \frac{b^2 - m^2}{2} \frac{\nu e^{(r^2-b^2)/2}}{1 - e^{(m^2-b^2)/2}},
$$

where

$$
\nu = \frac{s^2}{D}.
$$

We set a threshold $C_*$ of $C$ to indicate the beginning of the steep region. That means we consider $[r_s, b]$ as the steep region with $r_s$ satisfying $C(r_s) = C_*$. It can be seen that

$$
r_s = b - \frac{1}{\nu b} \ln \left( \nu (b^2 - m^2)/(2C_*) \right).$$

The reason we choose the equilibrium condition to determine the width of the steep region is that the concentration $C(r, t)$ builds up gradually at the bottom, with the steep region expanding over time. Thus using the equilibrium condition (which corresponds to infinite long time) will include all regions exhibiting a steep concentration change during the entire simulation process.

In our numerical tests we chose $C_* = 1/N$ to slightly increase the steep region for larger $N$. However, this is not
a critical choice. Since \( C(r, t) \) is an exponential function of \( r \) around the bottom, different \( C_\ast \) values have only a minor effect on the position of \( r_\ast \), and any other fixed value of \( C_\ast \) will also work well.

Next, we consider the requirement to eliminate the oscillations in the steep region. To avoid oscillations, the grid size \( \delta x \) must be sufficiently small compared to the value of \( D/(\kappa w^2) \). It is well known that for the discretization of sedimentation-diffusion problems the step size \( \delta x \) should be chosen such that the local Peclet number—which is proportional to the sedimentation speed and the grid size, and inversely proportional to the diffusion coefficient—is smaller than 1 (see Morton (10)). In our case, this condition implies that

\[
\delta x < h_\ast = \frac{2D}{\kappa w^2 b} = \frac{2}{\nu b}
\]  

(13)

We may simply choose a uniformly distributed grid satisfying Eq. 15 for the steep region. However, noticing that the solution \( C \) is exponential as \( r \) approaches the bottom \( b \), it is preferable to select a grid that becomes finer and finer as \( r \to b \). To this end, we use a simple sine function to determine the grid position in the steep region. We let

\[ N_\ast = \left[ \frac{\pi}{2}(b - r_\ast)/h_\ast \right] + 1 = \left[ \frac{\pi}{4}\ln(\nu(b^2 - m^2)N/2) \right] + 1 \]

be the total number of grid points in the steep region, where \([ \cdot ]\) represents the integer part of the number in between. We define the grid points in region \([r_\ast, b]\) as

\[ y_j = r_\ast + (b - r_\ast)\sin \left( \frac{(j - 1)\pi}{2(N_\ast - 1)} \right), \quad j = 1, 2, \ldots, N_\ast. \]

It follows that the grid size is given by

\[ y_{j+1} - y_j \approx (b - r_\ast)\frac{\pi}{2(N_\ast - 1)}\cos \left( \frac{j\pi}{2(N_\ast - 1)} \right) \]

\[ \approx h_\ast \cos \left( \frac{j\pi}{2(N_\ast - 1)} \right), \quad j = 1, 2, \ldots, N_\ast - 1. \]

Therefore, the step size decreases gradually as \( j \) increases. The leftmost interval is approximately of size \( h_\ast \), whereas the rightmost interval is approximately of size \( (\pi/2N_\ast)h_\ast \), which is one order of \( N_\ast \) smaller than \( h_\ast \).

Next we determine the grid in the regular region. We consider \((m, r_\ast)\) as the regular region with

\[ r_\ast = r_\ast - \frac{b}{N_\ast - 1} \ln \left( \frac{b}{m} \right). \]

The reason to choose \( r_\ast \) as the right end of the regular region is to make the grid size around the right end of the regular region match the grid size around the left-hand side of the transition region. When \( r \) is large, \( r_\ast \) is close to \( b \) and most of the cell is regular (see Table 1). The grid points in the regular region are determined analogous to the moving hat method, namely,

\[ x_j = m(b/m)^{(j-3/2)/(N_\ast - 1)}, \quad j = 2, 3, \ldots, N_\ast, \]

where

\[ N_\ast = \left\lfloor (N - 1)\log_{b/m} \left( \frac{r_\ast}{m} \right) + \frac{3}{2} \right\rfloor. \]

Finally, we determine the grid points in the transition region between \( r_\ast \) and \( r_\ast \). Let

\[ N_\ast = \left[ \log_{b/m} \left( \frac{r_\ast - r_\ast}{h_\ast} \right) \right] + 1. \]

There are \( N_\ast \) grid points in the transition region,

\[ t_j = r_\ast - (j - 1)h_\ast, \quad j = 1, 2, \ldots, N_\ast. \]

Note that the step size in the transition region halves from left to right. The length of the rightmost interval in the transition region is \( h_\ast \), and the length of the leftmost interval in this region is \( 2^{N_\ast - 1}h_\ast \), which is approximately the length of the rightmost interval in the regular region.

Since the grid point \( x_{N_\ast} \) at the right boundary of the regular region may not match the point \( t_{N_\ast} \) at the left boundary in the transition region due to integer rounding in calculating \( N_\ast \), we need to adjust \( t_{N_\ast - 1} \) such that it is positioned halfway in between its two neighboring grid points.

The grid over the entire cell is composed of all the points \( y_j, t_j, \) and \( x_j \) determined in the above manner. The total number of points actually used is \( N_\ast + N_\ast + N_\ast \). This number is typically a small fraction of \( N \); see Table 1 for the values of \( N_\ast, N_\ast \), and \( N_\ast \) for two examples—one with a moderate and one with a very large \( \kappa w^2/D \) value.

A typical grid defined by the above method is shown in Fig. 3. The nice feature of this mesh is that for most of the region the grid distribution follows the moving frame of

<p>|TABLE 1| The number of grid points used for the ASTFEM for the simulation of a typical cell with ( m = 5.8, b = 7.2 ), and different ( s, \omega, ) and ( D ). |
|----------|--------|--------|--------|------|--------|--------|</p>
<table>
<thead>
<tr>
<th></th>
<th>( S )</th>
<th>( \omega )</th>
<th>( D )</th>
<th>( s )</th>
<th>( \omega )</th>
<th>( D )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N )</td>
<td>101</td>
<td>201</td>
<td>2001</td>
<td>101</td>
<td>201</td>
<td>201</td>
</tr>
<tr>
<td>( N_\ast)</td>
<td>11</td>
<td>14</td>
<td>16</td>
<td>16</td>
<td>14</td>
<td>14</td>
</tr>
<tr>
<td>( N_\ast)</td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>99</td>
</tr>
<tr>
<td>( N_\ast)</td>
<td>115</td>
<td>214</td>
<td>1010</td>
<td>130</td>
<td>231</td>
<td>999</td>
</tr>
<tr>
<td>( r_\ast)</td>
<td>7.169337</td>
<td>7.181502</td>
<td>7.191381</td>
<td>7.176244</td>
<td>7.188469</td>
<td>7.190767</td>
</tr>
<tr>
<td>( r_\ast)</td>
<td>7.194758</td>
<td>7.194472</td>
<td>7.193806</td>
<td>7.199993</td>
<td>7.199993</td>
<td>7.199992</td>
</tr>
</tbody>
</table>
reference, which is desirable for maintaining the right sedimentation speed and minimizing the numerical diffusion introduced by discretization. The grids near the cell bottom have a very small step size, which is important to suppress the oscillation and to describe the detailed structure near the bottom of the cell.

**Remark**

By Taylor’s expansion of \( C(r, t) \) it is apparent that the discretization error is bigger in regions where the concentration gradient is steep, and the error is small where the solution is relatively flat. To obtain a small overall approximation error, the grid size should be small in regions with large solution curvature, but can be large elsewhere. For a typical concentration function \( C(r, t) \) large curvatures do not only exist near the bottom of the cell, but can also exist around the moving boundary. Therefore, an ideal grid distribution should also have higher grid density around the boundary. However, since the solution boundary is moving, this requires adjustment of the grid distribution throughout the simulation, and the coefficient matrices in the linear algebraic system in Eq. 12 need to be recalculated. This will offset the computational savings offered by the adaptive methods. Therefore, we do not consider this option here.

**Claverie’s finite element method**

We first evaluated the finite element method based on a uniform fixed mesh as described by Claverie (2). The time-step sizes are chosen according to the formula given by Schuck (7),

\[
\delta t = \ln(b/m)/\omega^2 s(N - 1). 
\]

Our simulation example used the following parameters:

**Example 1**

\[
m = 6.5, \ b = 7.2, \ s = 10^{-12}, \ D = 2 \times 10^{-7}, \ 60,000 \ \text{rpm}. 
\]

This model approximates a 0.4-kilobase DNA molecule sedimenting through a 7-mm solution column in 1 h. All our simulations were initialized with unity concentration across the entire solution column. We tested four grid densities, \( N = 101 (\delta t = 25.90s), 201 (\delta t = 12.95s), 401 (\delta t = 6.48s), \) and \( 1001 (\delta t = 2.59s) \). In Figs. 4–6 we show the numerical solution \( c(r, t) \) in different regions and with various \( N \) and \( \delta t \). From these graphs it can be seen that:

1. In the region near the bottom of the cell the oscillation of the numerical solution \( c(x, t) \) depends mainly on \( \delta x = (b - m)/(N - 1) \), and not on \( \delta t \) (see Fig. 4).
2. In the region near the meniscus, the oscillation occurs only at the beginning of the simulation. The magnitude of the oscillation depends mainly on the time-step size \( \delta t \) (see Fig. 5). Also, if rotor acceleration is taken into account, the oscillation near the meniscus can be markedly reduced.
3. In the center of the cell, the accuracy of the solution is determined by both \( N \) and \( \delta t \) (see Fig. 6).
These observations clearly indicate that, to obtain a highly accurate numerical solution, $\delta x$ must be very small near the bottom of the cell, and $N$ and $\delta t$ must be adjusted simultaneously. This is our motivation for the design of the adaptive grids.

**Schuck’s moving hat method**

Next, we evaluate the performance of Schuck’s moving hat method. The grid for time level $t_n$ used in the moving hat method is given by

\[
x_i = m, x_N = b,
\]

\[
x_j = m\frac{b}{m}^{(j-3/2)/(N-1)}, \quad j = 2, 3, \ldots, N-1.
\]

The same grid is used for the next time-step $t_{n+1}$, but the entire grid is shifted right by one point, with the last point deleted. See Fig. 7 for a typical grid used in the moving hat method. The basic idea in the moving hat method is first to introduce a time-dependent coordinate transform $r' = r(r, t)$ and change the Lamm equation into a differential equation in $(r', t)$-variables, and then to discretize this equation with the standard finite element method. The exponential grid distribution from Eq. 16 sets the grid-speed equal to that of the sedimentation speed. In this case, the Lamm equation is free of sedimentation under the moving coordinate system. There are two issues not well addressed by the moving hat method:

1. The step size $\delta x$ determined by Eq. 16 is too large around the cell bottom (even larger than in the rest of the cell), although a smaller step size is actually necessary for this region to accurately resolve the exponential buildup of the concentration. Therefore, the numerical solution by the moving hat method exhibits oscillations around the cell bottom if $N$ is not sufficiently large (see Figs. 8 and 9, A, C, and E).

2. The coordinate transform used in the moving hat method is singular at the meniscus and at the bottom. This is because at these points the grid is required to move according to the sedimentation velocity, even though these positions are fixed and have to remain at the meniscus and at the bottom. Therefore, the finite element discretization cannot be applied to the two points at the meniscus and the bottom.

Schuck dealt with this problem by imposing two additional conditions:
1. The value $c_{2n+1}$ is the weighted average of $c_{1n+1}$ and $c_{3n+1}$.
2. The value $c_{2n}^{n+1}$ is determined by requiring that the total mass at $t_{n+1}$ is equal to that at $t_n$.

Although these two conditions are consistent with the continuous equation, they are not based directly on the finite element discretization, i.e., they are a type of post-processing, whose effect is not determined by the properties of the Lamm equation. Nevertheless, the moving hat method is much more accurate than the fixed grid finite element method, even though it does not fix the oscillation problems near the bottom of the cell associated with the standard method by Claverie.

**Comparison of ASTFEM with the moving hat method**

Next, we evaluated the performance of the ASTFEM method based on the adaptive grid solution described earlier, and compared it to the moving hat method. Since the moving hat method is much more accurate than the standard finite element method by Claverie et al. (2), we do not present the comparison with the fixed-mesh method.

To illustrate the difference between the two methods we chose the following experimental conditions corresponding to a 1-kilobase DNA molecule in random coil conformation sedimenting in a 1.4-cm column over 1.5 h:

*Example 2*

\[ m = 5.8, \ b = 7.2, \ s = 1.562 \times 10^{-12}, \]
\[ D = 1.279 \times 10^{-7}, \ 50,000 \text{ rpm}. \quad (17) \]

In this case, the time-step was calculated according to Eq. 16 to be $\delta t = 50.49$ for $N = 101$ points. In Figs. 8 and 9 we plot the numerical solution obtained from the moving hat method and the ASTFEM method with various $N$ and $\delta t$. We observe from these figures that:

1. For the moving hat method, if $N$ is sufficiently large (e.g., $N \geq 400$ for Example 1, above), the oscillation around the bottom can be eliminated, and the solution is quite accurate. When $N$ is moderate, the oscillation occurs around the bottom, and it does not propagate back into the cell by much. But the buildup of the concentration at the cell bottom has been considerably broadened (Figs. 8 C and 9 C), and the accuracy in the bottom region is very poor. When $N$ is small, e.g., $N = 100$ for Example 2, the oscillation can travel back rapidly toward the meniscus of the cell and destroy the solution profile completely.

2. For the ASTFEM method, the solutions are not only very well resolved around the bottom, but also maintain a high accuracy around the boundary at all times. Most importantly, they are free from oscillation around the cell bottom for both examples with $N = 100$. We even tested some unrealistic cases with $D = 10^{-14}$ and $N = 100$ and found no oscillation at the cell bottom, and the solution remains highly accurate. Thus, unlike the moving hat method, the ASTFEM solution with the adaptive grids proposed in this article never breaks down and is
extremely robust. The issue of conservation of mass is also handled remarkably well by ASTFEM. Even for a small number of grid points the method maintains the total mass within no less than seven-decimal-digits accuracy without any post-processing during the entire simulation.

To compare the two methods quantitatively, we choose as the reference solution a numerical solution obtained using Claverie’s fixed-mesh finite element method with a very large number of radial grid points and a very small $\delta t$ ($N = 10^4$ and $\delta t = 0.259$ for Example 1, and $N = 10^4$ and $\delta t = 5.049$ for Example 2). Then we measure the difference between the moving hat or ASTFEM solutions and the reference solution. Since in all experimental data analysis the data near the meniscus and near the bottom of the cell are excluded from the fit, we consider here only data points from the solution column that are slightly inside the meniscus and the bottom of the cell. The largest difference can be noticed at the bottom of the cell ($C$ and $D$), where the moving hat method is poorly conditioned. The oscillations near the meniscus are reduced for the ASTFEM method ($A$ and $B$), and can be significantly reduced for both solutions when slow rotor acceleration is modeled in the solution.

FIGURE 8 Concentrations obtained by the moving hat method ($A$, $C$, and $E$) and the ASTFEM method ($B$, $D$, and $F$) with various $N$ and $\delta t$ for Example 1 ($m = 6.5$ cm, $b = 7.2$ cm, $s = 10^{-12}$, $D = 2 \times 10^{-7}$, 60,000 rpm, 9000 s). ($A$ and $B$) The concentrations near the meniscus. ($C$ and $D$) The concentrations near the bottom of the cell. ($E$ and $F$) The concentrations on the entire cell. The largest difference can be noticed at the bottom of the cell ($C$ and $D$), where the moving hat method is poorly conditioned. The oscillations near the meniscus are reduced for the ASTFEM method ($A$ and $B$), and can be significantly reduced for both solutions when slow rotor acceleration is modeled in the solution.
\[ \| c(t) - c_{\text{ref}}(t) \|_{\infty} = \max_{r_a \leq r \leq r_b} |c(r, t) - c_{\text{ref}}(r, t)| \]
and the \( L^2 \) error (which is equivalent to the square-root-mean error in the discrete case)

\[ \| c(t) - c_{\text{ref}}(t) \|_{L^2} = \left( \frac{1}{r_b - r_a} \int_{r_a}^{r_b} |c(r, t) - c_{\text{ref}}(r, t)|^2 \, dr \right)^{1/2}. \]

We plot in Fig. 10 the errors versus time \( t \) for different numerical methods. From these results we observe that both ASTFEM and the moving hat method (when it is convergent) exhibit second-order convergence properties (12), i.e., if the step size \( \delta x \) and \( \delta t \) are reduced by a factor of 2, the error is reduced by a factor of 4. With about the same number of grid points, ASTFEM is always more accurate than the moving hat method. In particular, when \( D \) is small, the moving hat method fails altogether, but ASTFEM remains quite accurate. If we include the entire cell from meniscus to bottom for the error comparison, then ASTFEM is much more accurate than the moving hat method, since the latter cannot resolve the solution sufficiently at the bottom.

*FIGURE 9* Concentrations obtained by the moving hat method (A, C, and E) and the ASTFEM method (B, D, and F) with various \( N \) and \( \delta t \) for Example 2 \((m = 5.8 \, \text{cm}, b = 7.2 \, \text{cm}, s = 1.562 \times 10^{-12}, D = 1.279 \times 10^{-7}, 50,000 \, \text{rpm}, 6000 \, \text{s})\). (A and B) The concentrations near the meniscus. (C and D) The concentrations near the bottom of the cell. (E and F) The concentrations on the entire cell.
Guideline for choosing $N$

A practical question in the numerical solution of the Lamm equation is how many grid points one should use to get a sufficiently accurate approximate solution. A satisfactory answer to this question clearly depends on the following factors: the measure of what constitutes an acceptable error, the portion of the solution column to be included for the finite element fit of the experimental data, and the parameters for the Lamm equation. For our error comparison we consider here the $L^2$-norm, and include only 90% of the solution column in middle for the accuracy check, and report on the dependence of the accuracy on $N$, $s$, $D$, and $\omega$.

First, it is clear that the effects of $s$ and $\omega$ are not independent of each other. It is the product of $s\omega^2$ which determines the behavior of the solution of the Lamm equation. Therefore, we fix the rotor speed at 50,000 rpm, and examine the effects of various values of $s$.

Here we consider two scenarios, the first is for a fixed $D$ and a varied $s$, and the second is for a fixed $s$ and a varied $D$. In Fig. 11 we plot the $L^2$ error versus time for various sedimentation coefficients. Note that for different sedimentation coefficients the total time needed to reach the equilibrium state is different. Hence the error curves dip at different times. Furthermore, it can be seen from Fig. 11 that for different $s$, the accuracy is different mainly at the beginning of the process. Once the solution profile is established, the solution accuracy is almost the same for different $s$. This can be explained by the fact that the Lamm equation is sedimentation-free under the moving computation grids. Therefore, the solution accuracy is almost independent of the sedimentation coefficient.

To examine the accuracy with respect to the diffusion coefficient $D$, we average the $L^2$ error at each time-step $t$ to get an averaged global $L^2$ error. More precisely, we consider

$$\|c - c_{\text{ref}}\| = \left(\frac{1}{T(r_b - r_a)} \int_0^T \int_{r_a}^{r_b} |c(r,t) - c_{\text{ref}}(r,t)|^2 \, dr \, dt\right)^{1/2},$$

where $T$ is the duration of the experiment. The square of this measure is equivalent to the variance for the samples consisting of all the $c(r, t)$ values. In Fig. 12 we plot the
averaged $L^2$ error versus the reciprocal of the square root of the diffusion coefficient. This graph indicates a linear relationship between the error and $1/\sqrt{D}$.

Combining the above observations, we conclude that away from the meniscus and the bottom the error of the ASTFEM solution is roughly a linear function of $1/\sqrt{D}$, and a quadratic function of $1/N$, and independent of $s$ or $\omega$, i.e.,

$$\| c - c_{\text{ref}} \| \approx \text{const.}/(N^2 \sqrt{D}).$$

In practice, one may estimate the error by using the Richardson extrapolation technique. More precisely, start with a small $N$, say $N = 100$. Compute the ASTFEM solution using $N$ points. Then compute another ASTFEM solution using $2N$ points, and by Eq. 18 we have

$$\| c_{2N} - c_{\text{ref}} \| \approx \frac{1}{4} \| c_N - c_{\text{ref}} \| .$$

Hence we have from the triangle inequality that

$$\| c_N - c_{\text{ref}} \| \leq \| c_N - c_{2N} \| + \| c_{2N} - c_{\text{ref}} \| \approx \| c_N - c_{2N} \| \frac{1}{4} \| c_N - c_{\text{ref}} \| ,$$

which implies

$$\| c_N - c_{\text{ref}} \| \leq \frac{4}{3} \| c_{2N} - c_N \| .$$

This can predict quite accurately the range of the actual error. Listed in Table 2 is the comparison of the actual error against a reference solution (obtained by using 1001 points), and the difference $|c_N - c_{2N}|$ for $N = 101$. Clearly this extrapolation technique can be used to get an estimate of the actual error and determine a proper $N$.

CONCLUSION AND DISCUSSIONS

In the previous sections we studied the behavior of three numerical solutions of the Lamm equation, and proposed an adaptive space-time finite element solution method. The features of this ASTFEM approach include:

1. It is free from the oscillation typically observed at the bottom of the cell with Claverie’s finite element method and Schuck’s moving hat method in experiments with large molecules and/or high rotor speed. Indeed, there is no observable oscillation for the ASTFEM solution with only $\sim 100$ points for any experimental parameter combination we have tried, suggesting that the ASTFEM solution provides superior robustness.
2. ASTFEM is much more accurate than Claverie’s fixed-mesh or Schuck’s moving hat method in the region near the cell bottom. It is also always more accurate than these two methods in the other regions of the cell.
3. ASTFEM automatically guarantees mass conservation of the Lamm equation without any post-processing.
4. The computational cost of ASTFEM is the same as for Claverie’s and Schuck’s method. All of them need to solve a tridiagonal linear system of equations with a fixed

\[ \begin{array}{c|cccc}
\text{D} = 1e - 7 & D = 2e - 7 & D = 4e - 7 & D = 8e - 7 \\
\hline
\| c_N - c_{\text{ref}} \| & 1.01579e - 03 & 6.54426e - 04 & 4.36885e - 04 & 2.99125e - 04 \\
\| c_N - c_{2N} \| & 1.00326e - 03 & 6.63717e - 04 & 4.51976e - 04 & 3.17079e - 04 \\
\end{array} \]
coefficient matrix for each time-step. Therefore, we conclude that ASTFEM is superior to both Claverie’s and Schuck’s methods.

The significance of the improvements in accuracy and concomitant efficiency are important for several reasons. The numerical solution of the Lamm equation by finite element methods is computationally relatively expensive. Nonlinear least-squares fitting of experimental sedimentation data with such solutions is an iterative process that often requires many repetitions of function evaluations. Statistical analysis of fitting results with Monte Carlo requires several thousand evaluations. Thus, improvements in efficiency are amplified in fitting applications and statistical analysis. In addition, the robustness displayed by the ASTFEM method is important for the stability of unconstrained fitting sessions where parameters may assume values under which the moving hat or fixed-mesh method will simply fail, but ASTFEM solutions will remain well-conditioned.

The idea of space-time finite element discretization can be extended directly to the cases of interacting and self-association systems, and cases for concentration dependency of \( s \) and \( D \). One of the advantages of the ASTFEM approach is that the mass conservation is satisfied automatically in these cases. We will report the numerical solution and results for interacting and concentration-dependent solutes in a forthcoming publication.

It is noted that we only attempted to use adaptivity around the cell bottom. Mesh adaptation to the concentration boundary would require the update of the coefficient matrix of the linear system, which increases the computational work substantially. However, for cases where the diffusion or sedimentation coefficients are dependent on the concentration itself, the coefficient matrix needs to be updated from time to time anyway, and then such adaptation will not introduce additional work to the solution procedure. This is a topic under current investigation.

The ASTFEM method is programmed in the UltraScan software, which is available for free download from our website (http://www.ultrascan.uthscsa.edu) for all major computer platforms (see also Demeler (13)).

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