# A Numerical Scheme for Particle-Laden Thin Film Flow in 2-D

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## Abstract

The physics of particle-laden thin film flow are not fully understood, and recent experiments have raised questions with current theory. There is a need for fully 2-D simulations to compare with experimental data. To this end, a numerical scheme is presented for a lubrication model derived for particle-laden thin film flow in two dimensions with surface tension. The scheme relies on an ADI process to handle the higher-order terms, and an iterative procedure to improve the solution at each timestep. This is the first paper to simulate the 2-D particle-laden thin film lubrication model. Several aspects of the scheme are examined for a test problem, such as the timestep, runtime, and number of iterations. The results from the simulation are compared to experimental data. The simulation shows good qualitative agreement. It also suggests further lines of inquiry for the physical model.

 $Keywords:\;$  adaptive timestepping, alternating direction implicit, coupled system, fourth order, particle-laden, thin film

# <sup>1</sup> 1. Introduction

In recent years, the problem of numerically solving gravity-driven thin film flow for clear fluids 2 has had ample work done in both one and two dimensions. However, the case when the film 3 contains particles suspended within it has received less attention, especially in two dimensions. 4 The evolution of a clear fluid down an inclined plane is modeled using a single partial differential 5 equation and numerical schemes have been derived using finite differences [15] and finite elements 6 [31]. For similar equations, such as spreading thin films, there are methods for finite elements in 7 one dimension [9, 10, 36] and for finite differences in two dimensions [34]. The incorporation of 8 particles into such a flow leads to another variable in the model, namely the particle concentration, g and an accompanying equation related to the evolution of the particles. The result is a system 10 of equations that requires a different approach from the clear-fluid case to formulate a practical 11 numerical scheme. 12

An active area of research in the last decade has been the development of numerical methods for higher-order thin film equations including complex fluids described by systems of equations. Related problems include methods for coupled systems of nonlinear parabolic equations [21, 25]. The scheme presented here is, in part, inspired by recent models for surfactants [33] and thin films [34]. We choose an Alternating Direction Implicit (ADI) scheme as a tractable method for implicit

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timesteps, because surface tension introduces a severe restriction on the timestep in the case of explicit schemes. While ADI schemes for numerically solving parabolic equations date back to the 1950's [26], their use in higher-order problems is rather new, e.g., [34], and not all that well-studied. However, the ease of parallelization makes such schemes a very viable choice for multiprocessor platforms. Since their inception, ADI schemes have been extended to handle parabolic problems with mixed derivative terms [2, 8, 23, 29], variable coefficients [14, 34], and high-order terms [34].

The ideas present in these schemes can be combined to create an efficient way to numerically 24 solve the particle-laden thin film flow equations. The nonlinearity and higher-order terms are han-25 dled in a similar manner to Witelski and Bowen [34], which dealt with thin-film equations, and 26 the remaining terms are treated as in Warner et al. [33], which devised a semi-implicit scheme 27 for surfactants. This combined approach is fine-tuned to draw out better efficiency, via adaptive 28 timestepping and an iterative procedure within each timestep. At the cost of the extra calcula-29 tions due to the iterative nature of the scheme, the timestep needed for stability can be improved 30 over recent methods. The result is an efficient method to simulate the continuum model in two 31 dimensions. 32

The full physics of particle-laden thin film flow is not well understood. Recent experiments, and their comparison to the model, have raised questions. We present such a comparison in this paper, where the results show qualitative agreement. In particular, by performing 2-D simulations, we are able to observe finger formation and compare directly with experiments. There is a need for accurate, fully 2-D simulations of the model for comparison with experimental data, such as in the case of mudslides and oil spills.

The paper is organized as follows: Section 2 presents the system of evolution equations for 39 the flow. In Section 3, the numerical scheme for this system is derived. Section 4 covers the 40 adaptive timestepping scheme implemented in the code. A complete explanation of the spatial 41 discretization is given in Section 5. The practicality and implementation of a moving reference 42 frame in the simulations are discussed in Section 6. Numerical simulations are presented in Section 43 7. We compare the results generated from the numerical scheme to an experiment using silicone 44 oil and glass beads in Section 8. Finally, in Section 9, we provide a discussion of the results and 45 future work. 46

## 47 **2. Model**

The results from experiments indicate that particle-laden thin film flows exhibit three distinct 48 regimes, based on the initial particle concentration and angle of inclination [35]. For low concentra-49 tions and angles, the particles settle to the substrate with clear fluid flowing over top of them. The 50 behavior after sedimentation is similar to clear-fluid experiments, such as those done by Huppert 51 [13]. High concentrations and angles cause a particle-rich ridge to emerge at the front of the flow. 52 Medium concentrations and angles lead to a particle concentration which appears to stay well-53 mixed throughout the duration of the experiment. Based on Cook [5], this behavior likely belongs 54 to one of the two previously mentioned regimes but may not have evolved to the point where this 55 distinction can be made. 56

The evolution equations for the flow are based on the regime where the inclination angle and particle concentration are both high enough to induce the formation of a particle-rich ridge. It is formulated in terms of the thickness of the film, h, and the particle concentration by volume,  $\phi$  (see Figure 1). The equations for modeling this regime were first derived in Zhou et al. [35]; re-derived in Cook et al. [6], using conservation of volume rather than mass; and modified in Cook et al. [7],



Figure 1: The coordinate system and variables considered in this problem.

<sup>62</sup> adding in a shear-induced diffusion term to correct for an instability affecting  $\phi$ . The dimensionless <sup>63</sup> system [7] is

$$h_t + \nabla \cdot (hv_{av}) = 0, \tag{1}$$

$$(\phi h)_t + \nabla \cdot [\phi h (v_{av} + (1 - \phi) v_{rel}) - F_{diff}] = 0.$$
(2)

<sup>64</sup> The orientation for (1)-(2) is such that  $\mathbf{x}$  lies in the plane and is parallel to the direction of the <sup>65</sup> flow,  $\mathbf{y}$  is across the inclined plane and perpendicular to  $\mathbf{x}$ , and  $\mathbf{z}$  is normal to the plane.

<sup>66</sup> The volume-averaged velocity of the liquid and the particles together is

$$v_{av} = \frac{h^2}{\mu(\phi)} \nabla \nabla^2 h - D(\alpha) \left[ \frac{h^2}{\mu(\phi)} \nabla \left( \rho(\phi)h \right) - \frac{5}{8} \frac{h^3}{\mu(\phi)} \nabla \left( \rho(\phi) \right) \right] + \frac{\rho(\phi)}{\mu(\phi)} h^2 \hat{\mathbf{x}}, \tag{3}$$

where the terms in (3) come from surface tension, the effects of gravity normal to the inclined plane, and the effects of gravity parallel to the inclined plane.

The density of the fluid as a whole is  $\rho(\phi) = 1 + \Delta \phi$ ;  $\Delta = \frac{\rho_P - \rho_l}{\rho_l}$  is the difference in the densities between the particles and the liquid. The function  $\mu(\phi) = (1 - \phi/\phi_{max})^{-2}$  [17, 30] is the effective fluid viscosity, where  $\phi_{max}$  is the maximum packing fraction of particles, assuming the particles are spheres. For this problem, the maximum packing fraction has been empirically determined to be 0.58, while the theoretical value is 0.64 [32].  $D(\alpha) = (3Ca)^{1/3} \cot(\alpha)$  [3] is a modified capillary number, where Ca is the capillary number of the liquid and  $\alpha$  is the angle of inclination of the <sup>75</sup> plane on which the fluid is flowing ( $\alpha = 0$  corresponds to the plane being horizontal while  $\alpha = \pi/2$ <sup>76</sup> to vertical).

The settling velocity of the particles, relative to the velocity of the liquid, is a combination of three factors, assumed to be multiplicative,

$$v_{rel} = V_s f(\phi) w(h) \hat{\mathbf{x}}.$$
(4)

The coefficient  $V_s = \frac{2}{3}a^2\Delta$  in (4) is the Stokes settling velocity of a single sphere settling in a viscous liquid, where *a* is the dimensionless particle radius. A hindered settling function, in this case the Richardson-Zaki function  $f(\phi) = (1 - \phi)^5$  [28], accounts for the effect of sedimentation. The particles settling parallel to the substrate is modeled using a wall effects function,  $w(h) = A(h/a)^2/\sqrt{1 + (A(h/a)^2)^2}$  with A = 1/18. This function is an approximation to a method of images solution to a single sphere falling parallel to a vertical wall [12]. This has the property that it is near 0 for *h* small and near 1 for *h* large.

Since (3) contains higher-order terms but (4) does not,  $v_{rel}$  is not regularized. This leads to an instability affecting the particle concentration in numerical simulations [7]. To correct for this, a shear-induced diffusion term (5) was added in,

$$F_{diff} = \frac{3}{2}a^2 (3Ca)^{1/3} \hat{D}(\phi) \frac{h^2 \rho(\phi)}{\mu(\phi)} \nabla \phi.$$
(5)

This behavior can be seen in a 1-D example on the domain x: 0-50 with  $\Delta x = 0.05$ . The initial film thickness is a jump, from 1 to 0.05, smoothed by hyperbolic tangent. The initial particle concentration is taken to be  $\phi = 0.3$ . This simulation is similar to those described in Section 7, and a moving reference frame is used, as discussed in Section 6. By time t = 1000, the solution without the extra diffusion term has developed an instability (Figure 2) while the one with it is still stable (Figure 3).



Figure 2: The numerical solution of  $\phi$  at time t = 1000 without shear-induced diffusion. By this time, an instability has developed.



Figure 3: The numerical solution  $\phi$  at time t = 1000 with shear-induced diffusion (5). The solution is still stable due to the extra term.

Equation (5) accounts for horizontal diffusion of particles in the fluid caused by horizontal gradients of  $\phi$  and was derived based on results from Leighton [19] and Leighton and Acrivos [20]. The term  $\hat{D}(\phi) = (1/3)\phi^2 (1 + (1/2)e^{8.8\phi})$  is a dimensionless diffusion coefficient.

#### 98 3. Numerical Scheme

In the case of a gravity-driven clear fluid flow, the model reduces to a single equation [3] for the film thickness, h,

$$h_t + (h^3)_x + \nabla \cdot \left(h^3 \nabla \nabla^2 h - D(\alpha) h^3 \nabla h\right) = 0.$$
(6)

Solving (6), and similar problems, numerically in one and two dimensions has been done using several different methods [1, 15, 22, 31, 34]. Including particles in the physics not only adds a second equation, but couples it to the equation for the film thickness. The particle-laden case has been solved numerically in one dimension with methods such as forward Euler with upwind differencing [35] and the Lax-Friedrichs method [6] when the high-order terms are omitted, and backward Euler with centered differencing [35] when the terms are included.

This system of PDE's in two dimensions poses numerical difficulties beyond those present in the 107 clear-fluid problem. For both the clear and particle-laden cases, fully explicit schemes typically have 108 the problem that an  $O(\Delta x^4)$  timestep, assuming  $\Delta x = \Delta y$ , is needed for stability. One solution is 109 to use an implicit scheme. For the clear-fluid and similar problems, the nonlinearity combined with 110 an implicit scheme amounts to solving the problem at each timestep using an iterative process, such 111 as Newton's method, to converge to the solution [34]. For the particle-laden case, using an implicit 112 scheme typically requires that both equations be solved simultaneously, using an iterative process 113 to account for the nonlinearity. This results in a linear algebra problem with twice the number 114 of unknowns and a matrix that is twice as large in each dimension, compared to the clear-fluid 115 problem. Therefore, solving the particle-laden case leads to larger linear algebra problems to solve 116

at each timestep and the matrix from Newton's method will have a more complex structure thanfor clear fluids.

The goal of the scheme presented here is to circumvent some of the aforementioned difficulties. The advantages of this approach, over a purely explicit scheme or implicit with Newton's method, is that the timestep is more lenient than for a fully explicit scheme and the linear algebra problem that results from the implicit part of the scheme is reduced to a series of smaller banded matrix solves, which can be done efficiently and independently for each equation.

The numerical scheme that we employ for the particle-laden thin film flow problem is inspired by the schemes presented in Witelski and Bowen [34] for higher-order parabolic PDE's and Warner et al. [33] for surfactants. In Witelski and Bowen, an ADI scheme is derived for solving the nonlinear PDE known as the thin film equation,

$$h_t + \nabla \cdot \left( f(h) \nabla \nabla^2 h \right) = 0. \tag{7}$$

The ADI scheme for (7) is backward Euler in time and uses approximate values of h in the nonlinear 128 and mixed-derivative implicit terms. It is suggested to start with approximations, such as time-129 lagged values, for evaluating these terms and calculating the numerical solution at the timestep. 130 Then use this solution for the new approximate values within the same timestep and recalculate. 131 This results in an iterative scheme at each timestep. However, for solving the thin film equation, 132 it was noted that the iterations did not provide a noticeable improvement. Warner et al. use 133 this method for a coupled system of nonlinear PDE's relating to surfactants. They handle the 134 higher-order terms implicitly using Crank-Nicholson, and apply ADI to this. The remaining terms, 135 which are at least second-order in space, are treated explicitly. For the nonlinearity and mixed-136 derivative terms, the values are time-lagged and the problem is solved only once per timestep. In 137 the simulations,  $\Delta x = \Delta y = \pi/100 \approx 0.0314$  required a timestep of  $O(10^{-5})$ . 138

Our approach is to handle applicable terms implicitly, using ADI, and treat the remaining terms explicitly, as we show below. Iterations within each timestep allow for a larger  $\Delta t$  to be taken at the cost of some extra calculations. In general, the increase in the size of the timestep outweighs the extra computational work, as shown in Section 7.

For equation (1), the terms

$$\nabla \cdot \left(\frac{h^3}{\mu(\phi)} \nabla \nabla^2 h + \frac{\rho(\phi)}{\mu(\phi)} h^3 \hat{\mathbf{x}}\right) \tag{8}$$

can be handled implicitly. This is because the spatial derivatives on these terms are applied to h. Including the first-order terms in the implicit treatment allows them to be discretized spatially using centered differencing to maintain stability. First discretize the terms in (8) in time with backward Euler, including the time derivative,

$$h^{n+1} + \Delta t \nabla \cdot \left(\frac{h^3}{\mu(\phi)} \nabla \nabla^2 h + \frac{\rho(\phi)}{\mu(\phi)} h^3 \hat{\mathbf{x}}\right)^{n+1} = h^n.$$
(9)

<sup>148</sup> Write out the operators in (9) fully,

$$h^{n+1} + \Delta t \left[ \partial_x \left( \frac{h^3}{\mu(\phi)} h_{xxx} \right) + \partial_y \left( \frac{h^3}{\mu(\phi)} h_{yyy} \right) + \partial_x \left( \frac{\rho(\phi)}{\mu(\phi)} h^3 \right) \right]^{n+1} + \Delta t \left[ \partial_x \left( \frac{h^3}{\mu(\phi)} h_{yyx} \right) + \partial_y \left( \frac{h^3}{\mu(\phi)} h_{xxy} \right) \right]^{n+1} = h^n.$$
(10)

The idea behind the ADI approach is to reduce the implicit part of (10), with derivatives in both x and y, to a product of two operators, each with only derivatives in either x or y. To achieve this, the terms involving only x-derivatives and only y-derivatives are grouped together. Define the operators

$$D_x = \partial_x \left(\frac{h^3}{\mu(\phi)}\partial_{xxx} + \frac{\rho(\phi)}{\mu(\phi)}h^2I\right)^{n+1}, \ D_y = \partial_y \left(\frac{h^3}{\mu(\phi)}\partial_{yyy}\right)^{n+1}.$$
 (11)

Then replacing the terms in (10) with the definitions in (11), we have

$$h^{n+1} + \Delta t (D_x + D_y) h^{n+1}$$

$$+ \Delta t \left[ \partial_x \left( \frac{h^3}{\mu(\phi)} h_{yyx} \right) + \partial_y \left( \frac{h^3}{\mu(\phi)} h_{xxy} \right) \right]^{n+1} = h^n.$$

$$(12)$$

In order to obtain an ADI scheme from (12), note that  $I + \Delta t D_x + \Delta t D_y = (I + \Delta t D_x)(I + \Delta t D_y) - (\Delta t)^2 D_x D_y$  and so the left-hand side, with the addition of an  $O(\Delta t^2)$  term, can be written as a product of two one-dimensional operators.

$$(I + \Delta t D_x)(I + \Delta t D_y)h^{n+1} - (\Delta t)^2 D_x D_y h^{n+1}$$

$$+ \Delta t \left[ \partial_x \left( \frac{h^3}{\mu(\phi)} h_{yyx} \right) + \partial_y \left( \frac{h^3}{\mu(\phi)} h_{xxy} \right) \right]^{n+1} = h^n.$$

$$(13)$$

<sup>157</sup> To handle the nonlinear terms, which occur in front of derivatives, and mixed-derivative terms in <sup>158</sup> (13), define them as approximate, denoted by a tilde (e.g.,  $\tilde{h}^{n+1}$ ). The approximate terms can be <sup>159</sup> chosen in some reasonable manner, such as time-lagged or extrapolated. This will be discussed <sup>160</sup> in more detail later. Subtract the mixed-derivative terms from and add the  $O(\Delta t^2)$  term to both <sup>161</sup> sides. This leaves a scheme in which all the terms operating on  $h^{n+1}$  are known, as is the entire <sup>162</sup> right-hand side.

$$(I + \Delta t \tilde{D}_x)(I + \Delta t \tilde{D}_y)h^{n+1} = h^n$$

$$+ \left\{ (\Delta t)^2 \tilde{D}_x \tilde{D}_y - \Delta t \left[ \partial_x \left( \frac{\tilde{h^3}}{\mu(\tilde{\phi})} \partial_{yyx} \right) + \partial_y \left( \frac{\tilde{h^3}}{\mu(\tilde{\phi})} \partial_{xxy} \right) \right] \right\}^{n+1} \tilde{h}^{n+1}.$$

$$(14)$$

<sup>163</sup> For simplicity, define the operators in (14) as

$$\tilde{L}_x = I + \Delta t \tilde{D}_x, \ \tilde{L}_y = I + \Delta t \tilde{D}_y.$$

<sup>164</sup> Subtracting  $\tilde{L}_x \tilde{L}_y \tilde{h}^{n+1}$  from both sides of (14), which cancels the  $O(\Delta t^2)$  term, yields

$$\tilde{L}_x \tilde{L}_y \left( h^{n+1} - \tilde{h}^{n+1} \right) = -\left( \tilde{h}^{n+1} - h^n \right) - \Delta t \nabla \cdot \left( \frac{\tilde{h}^3}{\mu(\tilde{\phi})} \nabla \nabla^2 \tilde{h} + \frac{\rho(\tilde{\phi})}{\mu(\tilde{\phi})} \tilde{h}^3 \right)^{n+1}.$$
(15)

At this point, the implicit part of the scheme is complete and the explicit terms can be added back into (15) using forward Euler.

$$\tilde{L}_{x}\tilde{L}_{y}\left(h^{n+1}-\tilde{h}^{n+1}\right) = -\left(\tilde{h}^{n+1}-h^{n}\right) - \Delta t\nabla \cdot \left(\frac{\tilde{h}^{3}}{\mu(\tilde{\phi})}\nabla\nabla^{2}\tilde{h} + \frac{\rho(\tilde{\phi})}{\mu(\tilde{\phi})}\tilde{h}^{3}\right)^{n+1}$$

$$+\Delta t\nabla \cdot \left\{D(\alpha)\left[\frac{h^{3}}{\mu(\phi)}\nabla\left(\rho(\phi)h\right) - \frac{5}{8}\frac{h^{4}}{\mu(\phi)}\nabla\left(\rho(\phi)\right)\right]\right\}^{n}.$$
(16)

167 Define

$$u = h^{n+1} - \tilde{h}^{n+1}.$$

which can be thought of as a correction term to the approximation of  $h^{n+1}$ , and (16) can be written as a three-step process: two one-directional solves (17)-(18) and an update step (19).

$$\tilde{L}_{x}v = -\left(\tilde{h}^{n+1} - h^{n}\right) - \Delta t \nabla \cdot \left(\frac{\tilde{h}^{3}}{\mu(\tilde{\phi})} \nabla \nabla^{2} \tilde{h} + \frac{\rho(\tilde{\phi})}{\mu(\tilde{\phi})} \tilde{h}^{3}\right)^{n+1} + \Delta t \nabla \cdot \left\{D(\alpha) \left[\frac{h^{3}}{\mu(\phi)} \nabla \left(\rho(\phi)h\right) - \frac{5}{8}\frac{h^{4}}{\mu(\phi)} \nabla \left(\rho(\phi)\right)\right]\right\}^{n},$$
(17)

$$L_y u = v, (18)$$

$$h^{n+1} \approx \tilde{h}^{n+1} + u. \tag{19}$$

Since the operators  $\tilde{L}_x$  and  $\tilde{L}_y$  involve at most fourth-order terms, the spatial discretization of them will lead to a five-point stencil in the x- and y-direction, respectively. This discretization is discussed fully in Section 5. Along each row/column of the discretized domain, this results in a pentadiagonal linear algebra problem. This can be solved using a pentadiagonal solver, or a more generic banded matrix solver.

To help with the inaccuracy in the nonlinear and mixed-derivative terms resulting from approximation, an iterative procedure can be used at each timestep to improve the solution and size of the timestep. This was first suggested for the ADI scheme in the context of thin film equations [34]. This procedure amounts to repeating the three-step process associated with solving each equation at each timestep and updating the approximate solution with the most recent solution, until the new and approximate solutions sufficiently converge. This is similar to fixed-point iteration.

For the first equation, when entering the timestep, a choice must be made as to the value of  $\tilde{h}^{n+1}$ 181 and  $(\phi h)^{n+1}$ . Using h as an example, two reasonable choices would be a time-lagged approximation, 182  $h^n$ , which is a first-order accurate approximation in time, or an extrapolated approximation,  $2h^n$  – 183  $h^{n-1}$ , which is second-order in time. For adaptive timestepping, this extrapolation is given by 184  $h^n + (\Delta t/\Delta t_{old})(h^n - h^{n-1})$ , where  $\Delta t$  is the prospective timestep between  $t^n$  and  $t^{n+1}$  and  $\Delta t_{old}$ 185 is the timestep between  $t^{n-1}$  and  $t^n$ . While the second choice of an approximation is second-order, 186 it also requires storing an extra set of data, namely  $h^{n-1}$ . With this choice made, the three-step 187 process can be implemented, obtaining a solution,  $h^{n+1}$ . At this point, the approximation can 188 be redefined,  $\tilde{h}^{n+1} = h^{n+1}$ , and the process run again. This can be continued until convergence 189 between the approximate and new solution, or equivalently when the correction term u is small in 190 a chosen norm. 191

For (2), the ADI method is applied to  $\phi h$  as a whole, since the time derivative is on this term. The applicable terms in the equation are

$$\nabla \cdot \left[ -D(\alpha) \left( \Delta \frac{(\phi h)h^2}{\mu(\phi)} \nabla(\phi h) \right) + \phi h \left( \frac{\rho(\phi)}{\mu(\phi)} h^2 + (1-\phi) V_s f(\phi) w(h) \right) \hat{\mathbf{x}} \right].$$
(20)

As with (1), the time discretization of (20) is based on a backward Euler method

$$(\phi h)^{n+1} + \Delta t \nabla \cdot \left[ -D(\alpha) \left( \Delta \frac{(\phi h)h^2}{\mu(\phi)} \nabla(\phi h) \right) + \phi h \left( \frac{\rho(\phi)}{\mu(\phi)} h^2 + (1-\phi) V_s f(\phi) w(h) \right) \hat{\mathbf{x}} \right]^{n+1} = (\phi h)^n.$$
(21)

<sup>195</sup> Writing out the operators in (21) explicitly,

$$(\phi h)^{n+1} - \Delta t D(\alpha) \Delta \left[ \partial_x \left( \frac{(\phi h)h^2}{\mu(\phi)} \partial_x(\phi h) \right) + \partial_y \left( \frac{(\phi h)h^2}{\mu(\phi)} \partial_y(\phi h) \right) \right]^{n+1} + \Delta t \partial_x \left[ \phi h \left( \frac{\rho(\phi)}{\mu(\phi)} h^2 + (1-\phi) V_s f(\phi) w(h) \right) \right]^{n+1} = (\phi h)^n.$$
(22)

Define the operators in (22) involving only x-derivatives and only y-derivatives as  $\mathcal{D}_x$  and  $\mathcal{D}_y$ , respectively.

$$\mathcal{D}_x = -D(\alpha)\Delta\partial_x \left(\frac{(\phi h)h^2}{\mu(\phi)}\partial_x\right)^{n+1} + \partial_x \left(\left[\frac{\rho(\phi)}{\mu(\phi)}h^2(1-\phi)V_s f(\phi)w(h)\right]I\right)^{n+1}, \qquad (23)$$
$$\mathcal{D}_y = -D(\alpha)\Delta\partial_y \left(\frac{(\phi h)}{\mu(\phi)}h^2\partial_y\right)^{n+1}.$$

<sup>198</sup> Using (23), the equation can be compactly written as

$$(\phi h)^{n+1} + \Delta t \left( \mathcal{D}_x + \mathcal{D}_y \right) (\phi h)^{n+1} = (\phi h)^n.$$
(24)

Note that there are no mixed-derivative terms to handle in (24). The left-hand side can be written as the product of two one-dimensional operators, incurring an  $O(\Delta t^2)$  term in the process.

$$(I + \Delta t \mathcal{D}_x) \left( I + \Delta t \mathcal{D}_y \right) (\phi h)^{n+1} - (\Delta t)^2 \mathcal{D}_x \mathcal{D}_y (\phi h)^{n+1} = (\phi h)^n.$$
(25)

Add the  $O(\Delta t^2)$  term to both sides of (25), and make all terms that occur nonlinearly at time  $t^{n+1}$  approximate, as before.

$$\left(I + \Delta t \tilde{\mathcal{D}}_x\right) \left(I + \Delta t \tilde{\mathcal{D}}_y\right) (\phi h)^{n+1} = (\phi h)^n + (\Delta t)^2 \tilde{\mathcal{D}}_x \tilde{\mathcal{D}}_y (\tilde{\phi} \tilde{h})^{n+1}.$$
(26)

203 Define

$$\tilde{\mathcal{L}}_x = I + \Delta t \tilde{\mathcal{D}}_x, \ \tilde{\mathcal{L}}_y = I + \Delta t \tilde{\mathcal{D}}_y$$

and subtract  $\tilde{\mathcal{L}}_x \tilde{\mathcal{L}}_y (\tilde{\phi} \tilde{h})^{n+1}$  from both sides of (26) to obtain

$$\tilde{\mathcal{L}}_{x}\tilde{\mathcal{L}}_{y}\left((\phi h)^{n+1} - (\tilde{\phi}\tilde{h})^{n+1}\right) = -\left((\tilde{\phi}\tilde{h})^{n+1} - (\phi h)^{n}\right)$$

$$-\Delta t\nabla \cdot \left[-D(\alpha)\left(\Delta \frac{(\tilde{\phi}\tilde{h})\tilde{h}^{2}}{\mu(\tilde{\phi})}\nabla(\tilde{\phi}\tilde{h})\right) + \tilde{\phi}\tilde{h}\left(\frac{\rho(\tilde{\phi})}{\mu(\tilde{\phi})}\tilde{h}^{2} + (1-\tilde{\phi})V_{s}f(\tilde{\phi})w(\tilde{h})\right)\hat{\mathbf{x}}\right]^{n+1}.$$

$$(27)$$

The remaining terms can be incorporated into (27) via forward Euler.

$$\tilde{\mathcal{L}}_{x}\tilde{\mathcal{L}}_{y}\left((\phi h)^{n+1} - (\tilde{\phi}\tilde{h})^{n+1}\right) = -\left((\tilde{\phi}\tilde{h})^{n+1} - (\phi h)^{n}\right) \\
-\Delta t\nabla \cdot \left[-D(\alpha)\left(\Delta \frac{(\tilde{\phi}\tilde{h})\tilde{h}^{2}}{\mu(\tilde{\phi})}\nabla(\tilde{\phi}\tilde{h})\right) + \tilde{\phi}\tilde{h}\left(\frac{\rho(\tilde{\phi})}{\mu(\tilde{\phi})}\tilde{h}^{2} + (1-\tilde{\phi})V_{s}f(\tilde{\phi})w(\tilde{h})\right)\hat{\mathbf{x}}\right]^{n+1} \\
-\Delta t\nabla \cdot \left[\phi h\left(\frac{h^{2}}{\mu(\phi)}\nabla\nabla^{2}h - D(\alpha)\left(\frac{h^{2}}{\mu(\phi)}\nabla h - \frac{5}{8}\frac{h^{3}}{\mu(\phi)}\nabla(\rho(\phi))\right)\right) - F_{diff}\right]^{n}.$$
(28)

206 Define

$$w = (\phi h)^{n+1} - (\tilde{\phi}\tilde{h})^{n+1}.$$

Then (28) can be written out as the three-step process (29)-(31):

$$(\phi h)^{n+1} \approx (\tilde{\phi}\tilde{h})^{n+1} + w.$$
(31)

The spatial operators in the  $\hat{\mathcal{L}}_x$  and  $\hat{\mathcal{L}}_y$  terms are at most second-order, and spatial discretization leads to a three-point stencil in each direction. Similar to with the first equation, a tridiagonal solver or banded matrix solver can be used to solve along each row/column.

Solving the system, as a whole, at each timestep can be then done by solving the first equation for  $h^{n+1}$ , solving the second equation for  $(\phi h)^{n+1}$ , then recovering the particle concentration as  $\phi^{n+1} = (\phi h)^{n+1}/h^{n+1}$ .

#### **4.** Adaptive Timestepping

We use an adaptive timestepping scheme to advance the solution. The scheme utilizes the 215 solution at consecutive timesteps  $t^{n-1}, t^n, t^{n+1}$ . Based on a measure of error, it decides whether or 216 not to accept the new solution, and if it is reasonable to increase the size of the timestep. This 217 is a modification of the scheme used in Bertozzi et al. [4], in which it serves as an estimate of a 218 dimensionless local truncation error in time. Consider the solution of the film thickness, h, at times 219  $t^{n-1}, t^n$ , and  $t^{n+1}$ . Calculate  $e^{n+1} = (h^{n+1} - h^n)/h^n$  and  $e^n = (h^n - h^{n-1})/h^n$ . The modification 220 from the original method is to divide by the value  $h^n$  at each point rather than  $h_{max}^n = max_{i,j}\{h_{i,j}^n\}$ , 221 since it produces a better-working adaptive scheme for this problem. Denote the timestep going 222 from time  $t^n$  to  $t^{n+1}$  as  $\Delta t$  and from  $t^{n-1}$  to  $t^n$  as  $\Delta t_{old}$ . Then define 223

$$Error = \left\| \left| e^{n+1} - \frac{\Delta t}{\Delta t_{old}} e^n \right\| \right|.$$
(32)

This provides a dimensionless estimate of the local truncation error in time, accumulated over the grid. The solution will be accepted if this error is less than some tolerance, denoted  $Tol_1$ . If the error is less than a smaller tolerance,  $Tol_2 < Tol_1$ , for a fixed number of steps, the timestep is increased by a scale factor. If the error is larger than  $Tol_1$ , the maximum number of iterations within a timestep is surpassed, or the solution becomes negative, the timestep is reduced by a factor of 2.

Since (32) only takes into account one of the two variables, this error can be computed for  $\phi h$  as well. These two errors can be combined into an overall measure of the error by taking the maximum of the two, or by some other reasonable combination.

## 233 5. Spatial Discretization

We use centered finite differences for all spatial discretizations. Using the notation,  $h_{i+1/2,j} \approx (h_{i,j} + h_{i+1,j})/2$ , the fourth-order term in (1) is

$$\nabla \cdot \left(\frac{h^{3}}{\mu(\phi)} \nabla \nabla^{2} h\right)_{i,j}$$

$$\approx \left(\frac{h^{3}_{i+1/2,j}}{\mu(\phi_{i+1/2,j})} h_{xxx,i+1/2,j} - \frac{h^{3}_{i-1/2,j}}{\mu(\phi_{i-1/2,j})} h_{xxx,i-1/2,j}\right) / \Delta x$$

$$+ \left(\frac{h^{3}_{i+1/2,j}}{\mu(\phi_{i+1/2,j})} h_{yyx,i+1/2,j} - \frac{h^{3}_{i-1/2,j}}{\mu(\phi_{i-1/2,j})} h_{yyx,i-1/2,j}\right) / \Delta x$$

$$+ \left(\frac{h^{3}_{i,j+1/2}}{\mu(\phi_{i,j+1/2})} h_{xxy,i,j+1/2} - \frac{h^{3}_{i,j-1/2}}{\mu(\phi_{i,j-1/2})} h_{xxy,i,j-1/2}\right) / \Delta y$$

$$+ \left(\frac{h^{3}_{i,j+1/2}}{\mu(\phi_{i,j+1/2})} h_{yyy,i,j+1/2} - \frac{h^{3}_{i,j-1/2}}{\mu(\phi_{i,j-1/2})} h_{yyy,i,j-1/2}\right) / \Delta y.$$
(33)

Here, the third derivatives are calculated at half-grid points by differencing consecutive standard
 second-order approximations. Two representative examples are

$$h_{xxx,i+1/2,j} \approx \left(h_{i+2,j} - 3h_{i+1,j} + 3h_{i,j} - h_{i-1,j}\right) / \Delta x^3, \tag{34}$$

$$h_{xxy,i,j+1/2} \approx \left( (h_{i+1,j+1} - 2h_i, j+1 + h_{i-1,j+1}) / \Delta x^2 - (h_{i+1,j} - 2h_i, j+h_{i-1,j}) / \Delta x^2 \right) / \Delta y.$$
(35)

<sup>238</sup> The two second-order terms are discretized as

$$\nabla \cdot \left(\frac{h^3}{\mu(\phi)} \nabla(\rho(\phi)h)\right)_{i,j} \approx \left(\frac{h^3_{i+1/2,j}}{\mu(\phi_{i+1/2,j})} (\rho(\phi_{i+1,j})h_{i+1,j} - \rho(\phi_{i,j})h_{i,j}) - \frac{h^3_{i-1/2,j}}{\mu(\phi_{i-1/2,j})} (\rho(\phi_{i,j})h_{i,j} - \rho(\phi_{i-1,j})h_{i-1,j})\right) / \Delta x^2 \quad (36)$$
$$+ \left(\frac{h^3_{i,j+1/2}}{\mu(\phi_{i,j+1/2})} (\rho(\phi_{i,j+1})h_{i,j+1} - \rho(\phi_{i,j})h_{i,j}) - \frac{h^3_{i,j-1/2}}{\mu(\phi_{i,j-1/2})} (\rho(\phi_{i,j})h_{i,j} - \rho(\phi_{i,j-1})h_{i,j-1})\right) / \Delta y^2,$$

$$\nabla \cdot \left(\frac{h^4}{\mu(\phi)} \nabla(\rho(\phi))\right)_{i,j} \approx \left(\frac{h^4_{i+1/2,j}}{\mu(\phi_{i+1/2,j})} \left(\rho(\phi_{i+1,j}) - \rho(\phi_{i,j})\right) - \frac{h^4_{i-1/2,j}}{\mu(\phi_{i-1/2,j})} \left(\rho(\phi_{i,j}) - \rho(\phi_{i-1,j})\right)\right) / \Delta x^2$$

$$+ \left(\frac{h^4_{i,j+1/2}}{\mu(\phi_{i,j+1/2})} \left(\rho(\phi_{i,j+1}) - \rho(\phi_{i,j})\right) - \frac{h^4_{i,j-1/2}}{\mu(\phi_{i,j-1/2})} \left(\rho(\phi_{i,j}) - \rho(\phi_{i,j-1})\right)\right) / \Delta y^2.$$
(37)

<sup>239</sup> The advective term is discretized using a standard centered-differencing scheme.

The terms in (2) are discretized in the same manner since many of them are similar to those in (1). The fourth- and second-order terms that come from  $v_{av}$  are discretized as in (33)-(37), with hreplaced by  $\phi h$ . Both advective terms are done via standard centered differencing.

The shear-induced diffusion term is discretized the same way as (36)-(37).

$$\nabla \cdot \left( \hat{D}(\phi) \frac{h^2 \rho(\phi)}{\mu(\phi)} \nabla \phi \right)_{i,j} \approx \left( \hat{D}(\phi_{i+1/2,j}) \frac{h_{i+1/2,j}^2 \rho(\phi_{i+1/2,j})}{\mu(\phi_{i+1/2,j})} (\phi_{i+1,j} - \phi_{i,j}) - \hat{D}(\phi_{i-1/2,j}) \frac{h_{i-1/2,j}^2 \rho(\phi_{i-1/2,j})}{\mu(\phi_{i-1/2,j})} (\phi_{i,j} - \phi_{i-1,j}) \right) / \Delta x^2 + \left( \hat{D}(\phi_{i,j+1/2}) \frac{h_{i,j+1/2}^2 \rho(\phi_{i,j+1/2})}{\mu(\phi_{i,j+1/2})} (\phi_{i,j+1} - \phi_{i,j}) - \hat{D}(\phi_{i,j-1/2}) \frac{h_{i,j-1/2}^2 \rho(\phi_{i,j-1/2})}{\mu(\phi_{i,j-1/2})} (\phi_{i,j} - \phi_{i,j-1}) \right) / \Delta y^2.$$

Centered differencing is not used for the moving reference frame, if one is employed. Instead, a second-order upwind differencing scheme is used, which will be discussed in the next section.

#### 246 6. Reference Frame

The area of interest in the simulations is near the front of the flow, where effects like the capillary 247 and particle-rich ridge occur. With a fixed reference frame, the spatial domain would need to be 248 taken as the entire area over which the flow would evolve, leading to large portions of the domain 249 where no change is occurring. This issue can be easily addressed by using a moving reference frame. 250 To implement a moving reference frame, we add an extra term to each equation,  $-sh_x$  on the 251 left-hand side of (1) and  $-s(\phi h)_x$  on (2). Here, s is the speed at which the moving reference frame 252 travels. Zhou et al. [35] approximate the front speed by removing all terms from the equations 253 which are higher than first order, leaving only the advective terms. They observe that these terms 254 capture the large scale dynamics, including the speed of the shocks, and the ridges that develop in 255 h and  $\phi$ . This leaves a 2 × 2 system of scalar conservation laws of the form 256

$$h_t + [F(h,\phi h)]_x = 0, (\phi h)_t + [G(h,\phi h)]_x = 0,$$
(38)  
$$F(h,\phi h) = \frac{\rho(\phi)}{\mu(\phi)} h^3, G(h,\phi h) = \frac{\rho(\phi)}{\mu(\phi)} (\phi h) h^2 + (\phi h) (1-\phi) V_s f(\phi) w(h).$$

<sup>257</sup> The initial conditions for (38) are

$$h(x,0) = \begin{cases} h_l, & x \le 0, \\ h_r, & x > 0, \end{cases}$$
(39)

$$(\phi h)(x,0) = \begin{cases} \phi_0 h_l, & x \le 0, \\ \phi_0 h_r, & x > 0. \end{cases}$$
(40)

where  $h_l$  and  $h_r$  in (39) and (40) are the initial film thickness and the height of the precursor b, respectively, and  $\phi_0$  in (40) is the initial particle concentration of the fluid. These initial conditions specify a Riemann problem [18]. From the initial shock in both equations, an intermediate state emerges,  $(h_i, (\phi h)_i)$ . The weak form of this system produces two Rankine-Hugoniot jump conditions, which define the shock speeds, ahead and behind the intermediate states. For  $s_1$ , the speed of the shock behind the intermediate state, and  $s_2$ , the speed ahead, these conditions are given by

$$s_{1} = \frac{F(h_{i},(\phi h)_{i}) - F(h_{l},(\phi h)_{l})}{h_{i} - h_{l}} = \frac{G(h_{i},(\phi h)_{i}) - G(h_{l},(\phi h)_{l})}{(\phi h)_{i} - (\phi h)_{l}},$$

$$s_{2} = \frac{F(h_{r},(\phi h)_{r}) - F(h_{i},(\phi h)_{i})}{h_{r} - h_{i}} = \frac{G(h_{r},(\phi h)_{r}) - G(h_{i},(\phi h)_{i})}{(\phi h)_{r} - (\phi h)_{i}}.$$
(41)

This nonlinear system (41) of four equations and four unknowns,  $h_i$ ,  $\phi_i h_i$ ,  $s_1$ , and  $s_2$ , can be solved via Newton's method. For the simulations shown in Section 7, our reference frame speed is an average of the two speeds,  $s = (s_1 + s_2)/2$ .

The discretization of the terms for the moving reference frame is done explicitly using forward Euler combined with second-order upwind-differencing,

$$-sh_x \approx -s\frac{-h_{i+2,j}+4h_{i+1,j}-3h_{i,j}}{2\Delta x}$$

This was chosen for simplicity and that it visually produced better results than the other discretizations that were tried. The effects of this choice appear to be some minor dissipation and dispersion in some cases behind the particle-rich ridge in the concentration.

The moving reference frame can be used for both the 1-D and 2-D cases (see Figures 4 and 5). To demonstrate this, simulations were run under the same conditions as those in Section 7. The theory-based solution for the problem without higher-order terms (38)-(40) aligns well with the 1-D numerical solution for the full problem. The 2-D solution for the full problem with a perturbation to the initial film thickness leads to a finger that moves faster than the 1-D case and the troughs, to the sides of the finger, move slower.

This can be viewed more succinctly in Figures 6 and 9, where the contours of the 1-D and 278 perturbed 2-D cases are overlayed. The position of the finger runs ahead of the 1-D front while 279 the troughs lag behind. Figure 7 shows the average front position of the flow for both cases. The 280 averaging of the front position was first done by Huppert [13] for experiments involving clear fluids. 281 Both simulations start with the same volume and, after an initial transient, the average front 282 positions of the film for the 1-D and perturbed 2-D cases (measured at h = 0.5) remains constant 283 and close to each other. Figure 8 shows the position of the finger and the trough in the 2-D case 284 over time. 285

### 286 7. Simulations

A rectangular domain is used with the x-direction oriented down the inclined plane and the y-direction across the inclined plane. In all cases, the particle concentration is initially taken to be



Figure 4: Comparison of theory and simulations at time t = 100 for the film thickness, h: theory without higher-order terms (solid line), 1-D solution to the full problem (dashed line), perturbed 2-D finger (dotted line), and perturbed 2-D trough (dot-dashed line).



Figure 5: Comparison of theory and simulations at time t = 100 for the particle concentration,  $\phi$ . The labels are the same as in Figure 4.

 $\phi(x, y, 0) = \phi_0$ , where  $0 \le \phi_0 \le \phi_{max}$ . This corresponds to having a well-mixed initial fluid. The film thickness far behind the contact line is set at h(x, y, 0) = 1 and ahead of the flow, a precursor of height h(x, y, 0) = b is assumed. At the contact line, a perturbation to a linear front can be applied to induce behavior such as a fingering instability. The parameters in the model are taken to be:  $a = 0.1, \Delta = 1.7, Ca = 10^{-3}, \alpha = \pi/4$ . The constant  $\phi_{max}$  is taken to be 0.67, in line with the simulations in Cook et al. [7]. The initial timestep is set to  $\Delta t = 10^{-6}$ .

<sup>295</sup> For the model, two sources contribute to the height of the film thickness and particle concentra-



Figure 6: A contour plot of the simulation at times t = 0 and t = 100 for the film thickness, h, in the 1-D and perturbed 2-D cases. The perturbation in 2-D leads to a fingering instability not seen in the 1-D case.



Figure 7: The average front position of the film thickness, h, of the 1-D and perturbed 2-D cases up to time t = 100. After an initial transient, the speeds stay close to constant and to each other.

tion near the front of the flow. The first is the higher-order terms, such as surface tension, which produce smooth ridges in both h and  $\phi$ . Second, even without these terms, an intermediate state at the front emerges for both variables, higher than either of their respective left or right states. These heights are dependent on the precursor b.

The height of the precursor in the following simulations is chosen out of convenience, to keep the size of  $\Delta x$  close to the precursor height. In general, choosing a different precursor has a small effect on the speed of the flow, but a large effect on both the film thickness and particle concentration. To illustrate this, Table 1 shows the height of the intermediate states for both h and  $\phi$  as well as the speeds of the trailing and leading shocks obtained from the theory-based solution to the system of scalar conservation laws (38)-(40) (see Section 6 for a more in-depth discussion). The



Figure 8: The front position of the film thickness, h, of the perturbed 2-D case up to time t = 100 along the finger and trough.



Figure 9: A contour plot of the simulation data at time t = 100 for the particle concentration,  $\phi$ , in the 1-D and perturbed 2-D cases. The perturbation leads to a particle-rich ridge that outlines and begins to fill in the finger.

intermediate film thickness  $h_i$  and particle concentration  $\phi_i$  increase as the height of the precursor 306 b decreases. For the shock speeds, a smaller precursor leads to the trailing shock speed  $s_1$  staying 307 relatively the same, but the leading shock speed  $s_2$  slows down and approaches  $s_1$ . These results 308 agree with the previous ones related to solving the system of scalar conservation laws [6, 35]. For 309 this model, the smallest precursor for which a solution exists is  $b \approx 9 \times 10^{-4}$  [6]. A precursor 310 close to this case, b = 0.001, produces shocks speeds which are close together and an intermediate 311 particle concentration near the maximum packing fraction. An alternative settling function that 312 permits solutions with smaller precursors,  $f_B(\phi) = (1 - \phi/\phi_{max})^5$ , is examined in Cook et al. [6]. 313

| b       | $h_i$   | $\phi_i$ | $s_1$    | $s_2$    |
|---------|---------|----------|----------|----------|
| 0.1     | 1.01653 | 0.307566 | 0.459323 | 0.510221 |
| 0.05    | 1.03478 | 0.315538 | 0.459314 | 0.483782 |
| 0.025   | 1.07107 | 0.330331 | 0.459301 | 0.471418 |
| 0.0125  | 1.1427  | 0.356006 | 0.459289 | 0.465441 |
| 0.00625 | 1.28276 | 0.396078 | 0.459294 | 0.462488 |
| :       | :       | :        | :        | :        |
| 0.001   | 9.14247 | 0.635545 | 0.459788 | 0.459916 |

Table 1: The intermediate states and shock speed solutions from equation (41) based on the precursor thickness b. As the precursor decreases, both  $h_i$  and  $\phi_i$  increase and the shock speeds converge.

The boundary conditions for h are Dirichlet in front and behind the flow and Neumann on the sides. The same is done for  $\phi$ . In addition, all third derivatives in h, normal to the boundary, are set to 0. More specifically, for a rectangular domain with length  $X_0$  and width  $Y_0$ , the boundary conditions are

$$h(0, y) = 1, \ h_{xxx}(0, y) = 0, \ h(X_0, y) = b, \ h(X_0, y) = 0,$$
  
$$h_y(x, 0) = 0, \ h_{yyy}(x, 0) = 0, \ h_y(x, Y_0) = 0, \ h_{yyy}(x, Y_0) = 0,$$
  
$$\phi(0, y) = \phi_0, \ \phi(X_0, y) = \phi_0, \ \phi_y(x, 0) = 0, \ \phi_y(x, Y_0) = 0.$$

The simulations are all run using moving reference frames, with the speed of the frame determined as in Section 6.



Figure 10: The speed-up gained by going from 1 to N processors using OpenMP. The line y = N is shown as a point of reference.

The code is written in parallel using the C++ OpenMP package. This choice of parallelization was made since the majority of calculations are done via *for* loops and OpenMP works well with <sup>322</sup> loop-heavy code. This includes the calculation of all finite differences and the solves along rows and <sup>323</sup> columns associated with the ADI part of the scheme. This is especially useful since rows/columns <sup>324</sup> can be solved independently of each other for each equation. In addition, writing special solvers <sup>325</sup> for linear systems of equations across multiple processors [24, 27] is avoided by this approach. The <sup>326</sup> speed-up attained using N processors is calculated by dividing the runtime for one processor by <sup>327</sup> the runtime for N processors (Speed –  $Up = Time(1 \ Processor)/Time(N \ Processors))$ .

Based on Figure 10, the scaling seems close to linear up to 4 processors, with a small drop-off 328 in performance as the number increases. This almost-linear behavior is a result of all of the code, 329 outside of a few minor calculations and the recording of the data, being amenable to parallelization. 330 To test some preferences that need be chosen a priori in the simulation, we conducted short-331 time tests to gauge the effectiveness of each approach. The ones considered here are (a) whether to 332 time-lag or extrapolate the approximate terms and (b) whether or not to perform iterations past 333 a single solve to improve the approximate terms, and therefore the solution at each timestep (see 334 Table 2). 335

| (a) Approximate Terms | Time-Lag      | Extrapolate         |
|-----------------------|---------------|---------------------|
| (b) Iterations        | One Iteration | Multiple Iterations |

Table 2: The two choices to be made when implementing the numerical scheme. One must choose whether to (a) time-lag or extrapolate the approximate terms and (b) whether to perform additional iterations past the initial solve.

Consider an initial condition of  $\phi_0 = 0.3$  and a front perturbed from Riemann initial data, h(x, y, 0) = 1 behind far behind the front, h(x, y, 0) = 0.05 far ahead of the front. At the jump from fluid to precursor, the shape of the front given as  $x_{front} = X_0/2 - \cos(2\pi y/Y_0)$ . This initial data is then smoothed using hyperbolic tangent and matched to the boundary condition (see Figure 13). This has the effect that the initial timestep can be taken more leniently.

We ran this initial simulation for each of the four combinations in Table 2 to time t = 1 and the maximum timestep allowed, average number of iterations per timestep, and the total runtime, in seconds, are listed in the table below (Table 3). This choice was made as the timestep changes dramatically over this time interval and can provide insight as to what methods seem practical for long-time runs. Since adaptive timestepping is utilized here, the tolerances are tuned so as to ensure that the simulation stays stable, not only to time t = 1 but for some time afterwards as well (it is taken up to t = 100 in this case, which is the length of the long-run simulations).

|                                 | $\Delta t_{max}$         | Avg. Iter. | Runtime |
|---------------------------------|--------------------------|------------|---------|
| Time-Lagged and One Iteration   | 0.000568341              | 1.0        | 518.2   |
| Time-Lagged and Iterations      | 0.00183296               | 2.20997    | 601.468 |
| Extrapolation and One Iteration | $4.07743 \times 10^{-5}$ | 1.0        | 19596.1 |
| Extrapolation and Iterations    | 0.00486338               | 1.29668    | 376.603 |

Table 3: Results for time t = 1 based on various choices for implementation.

<sup>348</sup> Using *Iterations* does well for both choices of approximate terms in that the total runtimes are <sup>349</sup> low, the maximum timesteps are large, and the number of iterations stays close to 1. Between these two, *Extrapolation and Iterations* does best, with nearly one fewer iteration required per timestep, on average, and a runtime that is 37% shorter. Performing *One Iteration*, the runtime for *Time-Lagged* is in between the two cases with *Iterations*, but for *Extrapolation*, it performs poorly, producing a runtime that is 33 to 52 times worse than the other three options. This is due to the small maximum timestep that is associated with this approach, which is 14 to 119 times smaller than the other three. At this point, it makes sense to discard the *Extrapolation and One Iteration* approach due to its excessive runtime and explore the remaining ones.

Under the same conditions, we ran a longer simulation, this time to t = 100. Using the best remaining options, we can glean some idea as to which one(s) will work best for a longer simulation.

|                               | $\Delta t_{max}$ | Avg. Iter. | Runtime |
|-------------------------------|------------------|------------|---------|
| Time-Lagged and One Iteration | 0.00107169       | 1.0        | 17811.3 |
| Time-Lagged and Iterations    | 0.00329173       | 2.95498    | 13153.8 |
| Extrapolation and Iterations  | 0.0106161        | 2.01204    | 3364.93 |

Table 4: Results for time t = 100.

Comparing Tables 3 and 4, the maximum timestep for each approach has increased. Using 359 Iterations, the average number has gone up in for both *Time-Lagged* and *Extrapolations*. However, 360 the average number of iterations per timestep for *Extrapolation* is approximately one iterations fewer 361 than for *Time-Lagged*. Also the runtime takes about 2.9 times longer for *Time-Lagged* compared to 362 *Extrapolation*. One can see the benefit of performing iterations instead of using a smaller timestep in 363 comparing the results for *Time-Lagged and One Iteration* and *Time-Lagged and Iterations*. *Time-*364 Lagged and One Iteration advances the solution approximately the same time forward with three 365 timesteps as *Time-Lagged and Iterations* does with one timestep and three iterations. However, 366 doing two extra timesteps costs more than two extra iterations, as seen in their respective runtimes. 367 This is because the explicit terms do not need to be re-calculated for each iteration while they do 368 for each timestep. Therefore, the only two options which make sense to use are the ones involving 369 *Iterations.* Of these, *Extrapolation* is the clear favorite. 370

In Figure 11, we see that by time t = 8, all three approaches have settled into a respective timestep. The timestep for *Extrapolations and Iterations* does best, followed by *Time-Lagged and Iterations* and *Time-Lagged and One Iteration*. The timestep for *Extrapolation and Iterations* is 3.2 times better than *Time-Lagged and Iterations* and 9.9 times better than *Time-Lagged and One Iteration*. The benefit of the larger timestep for both approaches with *Iterations* is partially offset by the need for extra calculations related to the iterations.

Figure 12 shows the number of iterations required throughout the simulation. For *Extrapolation* 377 and Iterations, the increase in iterations approximately between times t = 20 and t = 30 corresponds 378 to the finger forming and stretching out ahead of the flow in the film thickness and the particle-rich 379 ridge growing higher and outlining the finger. While the number of iterations jumps once to 3 and 380 then back down to 2 for Extrapolation and Iterations, it remains constant at 3 for Time-Lagged 381 and Iterations. The cost of storing extra data and performing a small computation to find the 382 extrapolated approximations seems a small price to pay to save one iteration per timestep, which 383 includes recalculating values involving the approximate terms and performing the ADI solves. 384

Using the simulation data up to t = 100, we can examine the effects of the initial perturbation graphically. For the film thickness, a small capillary ridge forms in the center of the perturbation



Figure 11: The adaptive timestep up to time t = 20. The timestep,  $\Delta t$ , is recorded in intervals of 0.25 for the three cases. Extrapolation and Iterations has a significantly larger timestep than either Time-Lagged and One Iteration or Time-Lagged and Iterations.



Figure 12: The number of iterations up to time t = 100. The iterations are recorded in intervals of 0.25 for the two cases. Using *Extrapolation and Iterations* does better than *Time-Lagged and Iterations* in terms of fewest number of iterations.

(Figure 14) and begins to stretch out ahead of the bulk flow (Figures 15 and 16). This is the wellknown fingering instability present in thin-film flows. For the particle concentration, a particlerich ridge initially forms at the contact line (Figure 17) and, as the fingering instability evolves, outlines the shape of the finger (Figures 18 and 19). Directly behind the ridge, a pocket of lower concentration forms. The interior of the finger is slowly encroached upon by the particles that have accumulated near the back and sides of the finger. This can be seen in Figure 19 as an interior



Figure 13: The initial film thickness. It is perturbed by a cosine wave along y and smoothed along x by hyperbolic tangent.



Figure 14: Film thickness at time t = 25. A small capillary ridge forms in the center of the flow.

<sup>393</sup> layer along the inside of the particle-rich ridge. It is possible that this phenomenon is not physical, <sup>394</sup> and may be a result of the current model not containing all of the necessary physics.

#### <sup>395</sup> 8. Comparison to Experiments

Experiments for particle-laden thin film flows have been compared in one dimension to the solution, both analytically and numerically, for clear-fluid flows. The average front position for clear fluids is given by a power law, where the location of the front scales like  $t^{1/3}$  [13]. Ward et al. [32] compare the average front position of the flow to this scaling. Grunewald et al. [11] compare



Figure 15: Film thickness at time t = 50. A fingering instability begins to develop from the ridge.



Figure 16: Film thickness at time t = 100. The finger stretches out ahead of the bulk flow.

the average front position to a re-derived 1-D model, based on results from Huppert [13] with a precursor, and to numerical solutions of the 1-D problem. Both find some agreement between the experiments and the scaling for the 1-D front position for clear fluids. We seek to compare the numerical solution in two dimensions to images of experiments, taking into account that variations occur across the front of the flow.

We use 1000 cSt Polydimethylsiloxane (PDMS), a silicone oil, for the liquid component of the fluid. For the particles, glass beads with diameters in the range of  $250 - 425 \ \mu m$  are used. The two components are then well-mixed and released down an inclined plane from a reservoir. This corresponds to a constant-volume experiment, whereas our simulations are constant-flux.



Figure 17: Particle concentration at time t = 25. A small particle-rich ridge forms at the front of the flow with a slightly higher concentration in the center.



Figure 18: Particle concentration at time t = 50. The particle-rich ridge increases in concentration and has a higher concentration in and around the fingering instability.

The experiment, which we will compare to simulation, is a fluid of approximately 90  $cm^3$ 409 containing a volume which is 35.9% particles. The plane is inclined at a 32-degree angle. The fluid is 410 allowed to flow down the plane, which is 14 cm across and 90 cm down. In the experiments, the flow 411 starts out close to uniform across the front, away from the edges, and over time develops instabilities, 412 in the form of fingers stretching out ahead of the bulk flow. Since, for simulations, starting with a 413 uniform front along the y-direction leads to a uniform solution, we start the simulation some time 414 after the start-time to add a perturbation to the initial data, which induces the type of behavior 415 seen in the latter stages of the experiments. 416



Figure 19: Particle concentration at time t = 100. The particle-rich ridge is composed of two parts: one outlining the finger and one entering the finger from the troughs.

In order to avoid simulating the problem over the entire domain, we truncate the solution near the front and treat the problem locally as being constant-flux. We are interested in the dynamics of finger formation during which time the film thickness only changes by at most 20%, so a local approximation by constant-flux is reasonable.



Figure 20: The initial condition of the experiment, used for comparing with the simulation. At this point, the front of the flow has begun to develop perturbations, which will lead to fingering instabilities.

We use two images, taken three minutes apart, to compare with the simulation. The first image is taken when the front of the flow has reached approximately 53 cm down the plane. The shape of the front is parabolic-like with two large perturbations at either end of the front. In between, smaller perturbations exist which lead to fingering instabilities. The two outer perturbations lead
to longer and thicker fingers than the smaller inner perturbations. We take a front similar to this
in our simulation.

The scales for a constant-flux problem can be taken from Cook et al. [6], which are the same as for the clear-fluid case. The height scale is taken to be  $h_0 = 1 \ mm$ . The length scale is  $x_0 = (l^2 h_0)^{1/3}$ , where the capillary length, l, is  $l = \sqrt{\gamma/\rho_l g_{\parallel}}$ . The constants are  $\gamma$ , the coefficient of surface tension;  $\rho_l$ , the liquid density; and  $g_{\parallel}$ , the component of gravity parallel to the inclined plane. The time scale is  $t_0 = (3\mu_l/\gamma)x_0l^2/h_0^2$ , where  $\mu_l$  is the dynamic liquid viscosity. The capillary number is given by  $Ca = \mu_l x_0/\gamma t_0 = h_0^2/3l^2$ .

The scales, given these parameters, are  $h_0 = 0.001 \ m$ ,  $x_0 = 0.00161396 \ m$ ,  $l = 0.00205041 \ m$ , 433  $t_0 = 0.93235 \ s$ , and Ca = 0.0792863. Using this, we can construct an initial condition which 434 resembles the experiment and will produce similar results. This is done by measuring the features 435 of the initial image and creating a similar condition. While the flow in the experiment is asymmetric, 436 we take a symmetric initial condition in the simulation which has features that are approximately, 437 in both location and size, the same as in the experiment. The track is taken to be 86.75 units 438 wide (rounded up to the nearest 0.05 increment), which corresponds to the 14 cm wide track. The 439 precursor in the simulation is set to b = 0.05, as in the previous simulations. 440

A moving reference frame is used since this is taken to be a constant-flux problem locally. The speed of the moving reference frame is approximately s = 0.343198, calculated as in Section 6. Running a simulation over the course of three minutes leads to a distance traveled for the frame of approximately 10.69 cm, where the actual displacement, based on experiments, is around 12 cm, so using the constant-flux assumption seems to produce a decent approximation of the distance the fluid will flow.



Figure 21: The initial condition for the film thickness, h, used in the simulation. This is an artificially-created starting condition to be representative of the state shown for the experiment. The height is in mm.

The initial data is generated using a sine wave to form the two large perturbations and the space away from the edges. The three fingers that develop between these two perturbations are simulated with a cosine wave of small amplitude, 0.25 in dimensionless units. The simulation is 450 run to t = 193.06, the equivalent of three minutes of real-time.



Figure 22: The evolution of the experiment after three minutes. The fingering instability starts to form at the front.



Figure 23: The evolution of the film thickness, h, in the simulation after three minutes. Both the experiment and simulation exhibit a fingering instability, but the instability in the simulation is less pronounced. The height is in mm.

Over the course of the three minutes, the exterior of the outer fingers in the experiments go from 4 cm and 6.5 cm on the left and right, respectively, to 7.5 cm and 12 cm. The interior of these fingers go from less than 1 cm on each side to about 3 cm. The interior fingers are not discernable in the initial image. The flow as a whole, measured from where the fluid touches the walls, has moved about 11 cm down the plane. The interior fingers in the experiment, extend about 0.5 cm ahead of the flow.



Figure 24: Particle concentration,  $\phi$ , for the film thickness in Figure 23.

In the simulation, the moving reference frame accounts for 10.69 cm of movement, so the position 457 where the fluid touches the edges has moved approximately 7.5 cm. The evolution of the fingers 458 in the simulation is slightly less pronounced than in the experiments. This is likely due to the 459 simulation initially undergoing a transient state where the fluid travels slower than at later times, 460 while the transient in the experiment has occurred prior to this three-minute interval. The exterior 461 of the outer fingers is approximately 4.2 cm and interior 1.2 cm. The interior fingers extend ahead 462 of the flow about 0.8 cm. The tip of the longest finger in the experiments has moved 15 cm while in 463 the simulations, it has advanced approximately 11.4 cm. The tips of the fingers, in the z-direction, 464 reach up to 1.37 mm. 465

The particle concentration cannot be determined accurately at the particle-rich ridge in the experiment, but the increased opacity at the leading edge of the flow indicates an increase in the concentration, relative to the ambient concentration. This change in shade is approximately 1 cmlong in the direction of the flow. In the simulations, the thickness of the ridge ranges from 0.6 to 1.1 cm, which is consistent with the experiments.

## 471 9. Discussion

Schemes originally derived for numerically solving high-order parabolic problems have recently been extended to high-order systems, such as the case of surfactants and particle-laden thin films. Handling the higher-order terms in a practical way is necessary for fast and efficient computation. The scheme we have presented here for particle-laden thin film flow provides an easy-to-program and effective way to solve this high-order coupled system. This scheme can provide a blueprint for approaches to solving similar problems.

The numerical scheme developed for particle-laden thin film flow has several nice attributes. The timestep required for this scheme is in the range of  $O(\Delta x^2)$ , which is much better than the  $O(\Delta x^4)$  for a fully explicit scheme. The structure of the scheme allows for the possibility of solving each equation with its own unique timestep for better efficiency, as the particle concentration is typically the equation that fails the timestep restriction criteria. The linear algebra problem that results from an implicit time discretization along with the nonlinearity is reduced to a series of triand pentadiagonal solves, which can be done in parallel along the rows/columns of the grid.

The parallelization of the code is straightforward using OpenMP. The loops for computing the explicit and approximate terms as well as the solves along rows and columns can be done in parallel, leading to a code that scales close to linearly for up to 8 processors, getting close to 8 times speedup. Adding OpenMP implementation to C++ code on any multicore machine is easy to implement, as it only requires adding a few lines of code to existing *for* loops and needs no managing of the movement of data on the programmer's part. Since the code is predominantly such loops, it is easy to parallelize and is highly effective in getting better runtimes.

Implementing *Iterations* within each timestep, which is first presented in Witelski and Bowen [34], but not used in Warner et al. [33], seems to work best for this problem, in terms of allowing for a larger timestep and producing an accurate solution. Among the choices for the approximate terms when performing *Iterations*, *Extrapolation* seems to produce the best runtime and fewest iterations. Implementation requires only storing an extra set of data used in extrapolating the approximate terms but, using the adaptive timestepping discussed here, this data is stored anyway.

The choice of *Extrapolation and Iterations* may work best for this problem, but for other problems or initial conditions, another choice may fare better. It is recommended, as in this case, that a short-term simulation be performed for the different choices of approximate terms and whether or not to perform extra iterations. The small cost of these short runs may allow for a more efficient run for actual simulations. It is also recommended that one examines the results to make sure that the scheme is not only fast with the choice, but sufficiently accurate.

The numerical solution agrees reasonably well with the behavior seen in experiments. This is in part because the model was derived for the case when a particle-rich ridge forms. This is seen in the experiments for high angles of inclination and high concentrations, but will occur in model for all concentrations and angles. The particle-rich ridge in the simulations is two thin layers of particles, one which originates at the front of the flow and the other from the troughs of the emerging fingers, which may not be physical.

The current model assumes a constant, or average, particle concentration throughout the fluid layer in the z-direction. The same is true for the velocity, which is averaged in the z-direction. Theory exists for the vertical movement of the particles [5], whether they will settle to the inclined plane or form a ridge, and incorporating this behavior into a new model is the current research of the authors. It is hoped that the current numerical scheme will be adaptable to this new model.

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