An implicit-explicit discontinuous Galerkin scheme using a newton-free Picard iteration for a thin-film model

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An implicit-explicit discontinuous Galerkin scheme using a newton-free Picard iteration for a thin-film model

by

Caleb Logemann

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in partial fulfillment of the requirements for the degree of

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The student author, whose presentation of the scholarship herein was approved by the program of study committee, is solely responsible for the content of this thesis. The Graduate College will ensure this thesis is globally accessible and will not permit alterations after a degree is conferred.

Iowa State University
Ames, Iowa
2019

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This paper provides a high-order numerical scheme for solving thin-film models of the form
\[ q_t + (q^2 - q^3)_x = -(q^3q_{xxx})_x. \]
The second term in this equation is of nonlinear hyperbolic type, while the right-hand side is of nonlinear parabolic type. The nonlinear hyperbolic term is discretized with the standard modal discontinuous Galerkin method, and the nonlinear parabolic term is discretized with the local discontinuous Galerkin method. Propagation in time is done with an implicit-explicit Runge-Kutta scheme so as to allow for larger time steps. The timestep restriction for these methods is determined by the hyperbolic wavespeed restriction, and is not limited by the nonlinear parabolic term. A novel aspect of this method is that the resulting nonlinear algebraic equations are solved with a Newton-free iteration specifically with a Picard iteration. The number of iterations required to converge is less than or equal to the estimated order of the method. We have demonstrated with this method up to third order convergence.
CHAPTER 1. INTRODUCTION

In this paper we look at the model equation,

\[ q_t + (q^2 - q^3)_x = -(q^3 q_{xxx})_x \quad (x, t) \in [a, b] \times [0, T]. \quad (1.1) \]

This equation describes the motion of a thin film of liquid flowing over a one-dimensional domain, \([a, b]\), where \(q(x, t) \geq 0\) is the height of the liquid. This fluid is acted upon by gravity, by forces on the surface, and by surface tension. The surface forces can have many different causes, including wind shear, thermocapillary forces, or molecular forces. In all cases an equivalent model can be derived. This model is useful in many different applications including airplane de-icing [20, 21] and industrial coating. Some experimental study [5, 14, 19] has been done and numerical results have shown good agreement with those experiments in [3].

Previous numerical methods for this type of equation have focused on finite difference approaches. Bertozzi and Brenner [2] used a fully implicit centered finite difference scheme to explore instabilities. Ha et al. [13] explored several different finite difference schemes, some fully implicit and some using the Crank-Nicolson method. In their analysis, they considered several different methods for the hyperbolic terms including WENO, Godunov, and an adapted upwind method. All of these methods were limited to just first or second order, and they required solving a Newton iteration. Finite difference methods also lack provable stability.

We chose to use discontinuous Galerkin methods as they allow for high order convergence. The discontinuous Galerkin methods were first introduced by Reed and Hill [26], and then were formalized by Cockburn and Shu in a series of papers [9, 8, 7, 6, 11]. We use the original modal discontinuous Galerkin method as well as the local discontinuous Galerkin method. The local discontinuous Galerkin method was also formulated by Cockburn and Shu [10] to handle convection-diffusion equations with the discontinuous Galerkin method. We use the modal discontinuous
Galerkin method to discretize the convection term, \((q^2 - q^3)_x\), and the local discontinuous Galerkin method to discretize the parabolic term, \(-\left(q^3 q_{xxx}\right)_x\).

The nonlinear diffusion in the parabolic term is much stiffer, as it has an infinite wavespeed, than the hyperbolic convection, which only has a finite wavespeed. If the parabolic term is handled explicitly in time, then a very strict time step restriction is needed to insure stability. Therefore the nonlinear parabolic term should be solved implicitly. The whole system could be solved implicitly, however we chose to use an implicit-explicit (IMEX) Runge-Kutta scheme for propagating in time. These schemes were first introduced by Ascher et al. [1] and have been expanded on by Kennedy and Carpenter [15] and Pareschi and Russo [24]. The IMEX Runge-Kutta schemes allows us not only to solve the nonlinear parabolic term implicitly, but it also allows for the nonlinear convection to be propagated explicitly. Solving the convection explicitly fully captures the nonlinear behavior without a nonlinear solve.

The parabolic term still requires a nonlinear solve, but this is simpler than solving both convection and parabolic terms nonlinearly. We solve the nonlinear system using a Picard iteration, first introduced by Émile Picard and then formalized by Lindelöf [17]. We find that very minimal number of iterations is required for the iteration to converge. The number of iterations is less than or equal to the overall order of accuracy of the method. With these methods we have demonstrated up to third order accuracy with a manufactured solution example. We also demonstrate the nonlinear traveling wave behavior through several numerical examples introduced by Bertozzi [4]. These examples demonstrate that the steady state traveling wave may not be unique. They also show a double shock structure with an undercompressive shock.
CHAPTER 2. THE THIN-FILM MODEL

In this model we consider a thin film of liquid on a flat surface with a free interface. This liquid is driven by gravity, shear and normal forces on the surface, and surface tension (see Figure 2.1). We begin by considering the two dimensional incompressible Navier-Stokes equations, which have the form,

\[ u_x + w_z = 0, \]  
\[ \rho (u_t + uu_x + uw_z) = -p_x + \mu \Delta u - \phi_x, \]  
\[ \rho (w_t + uw_x + ww_z) = -p_z + \mu \Delta w - \phi_z, \]

where \( \rho \) is the density, \( u \) is the horizontal velocity, \( w \) is the vertical velocity, \( p \) is the pressure, and \( \phi \) is the force of gravity. Equation (2.1) is the incompressibility condition and also represents conservation of mass. Equations (2.2) and (2.3) represent the conservation of momentum in the \( x \) and \( z \) directions respectively. We take a no penetration and no slip boundary condition at the lower boundary and the kinematic boundary condition at the upper boundary. These boundary conditions can be expressed as follows,

\[ w = 0, \quad u = 0, \quad \text{at } z = 0, \]  
\[ w = h_t + uh_x, \quad \text{at } z = h, \]

where \( h \) is the height of the liquid. We can also describe the stress tensor, \( T \), at the free surface, \( z = h \), as

\[ T \cdot n = (-\kappa \sigma + \Pi_0)n + \left( \frac{\partial \sigma}{\partial s} + \tau_0 \right)t, \quad \text{at } z = h, \]

where \( \kappa \) is the mean curvature, \( \sigma \) is the surface tension, and \( \Pi_0 \) and \( \tau_0 \) are the normal and tangential components of the forcing respectively.
2.1 Derivation

These equations completely describe the fluid, but we are now going to make a lubrication approximation, through a scaling argument similar to that given by Oron et al. [23]. For the lubrication approximation we are going to assume that the average height of the liquid, $h_0$, is much smaller than the characteristic wavelength of the liquid, $\lambda$. We will denote the ratio of these two lengths as $\varepsilon$, that is

$$\varepsilon = \frac{h_0}{\lambda} \ll 1. \quad (2.6)$$

Now we nondimensionalize the rest of the variables with respect to this ratio. We denote the nondimensional variables as the uppercase variables. As we stated earlier the characteristic height is $h_0$ and the characteristic length is $\lambda = h_0/\varepsilon$, so the nondimensional length variables are

$$Z = \frac{z}{h_0}, \quad X = \frac{\varepsilon x}{h_0}, \quad H = \frac{h}{h_0}. \quad (2.7)$$

Let $U_0$ be the characteristic horizontal velocity, then

$$U = \frac{u}{U_0}, \quad W = \frac{w}{\varepsilon U_0}, \quad (2.8)$$

where the nondimensional vertical velocity, $W$, follows from the continuity equation, equation (2.1).

It follows that time will be scaled by $\lambda/U_0$, so nondimensional time is

$$T = \frac{\varepsilon U_0 t}{h_0}. \quad (2.9)$$
Finally we assume that the flow is locally parallel or equivalently $p_z \sim \mu u_{zz}$. This gives us the proper scaling for the pressure, gravity, and surface stresses,

$$P = \frac{\varepsilon h_0}{\mu U_0} p, \quad \Phi = \frac{\varepsilon h_0}{\mu U_0} \phi, \quad \Pi = \frac{\varepsilon h_0}{\mu U_0} \Pi_0, \quad \tau = \frac{h_0}{\mu U_0} \tau_0.$$  \hspace{1cm} (2.10)

Lastly we can nondimensionalize the surface tension as

$$\Sigma = \frac{\varepsilon \sigma}{\mu U_0}.$$ \hspace{1cm} (2.11)

Substituting these nondimensional variables into the continuity equation (2.1), gives

$$u_x + w_z = 0,$$

$$\frac{\varepsilon U_0}{h_0} U_X + \frac{\varepsilon U_0}{h_0} W_Z = 0,$$

$$U_X + W_Z = 0.$$

This computation also justifies the scaling of $w$ as the nondimensional variables should also conserve mass.

We can nondimensionalize the conservation of momentum equations as follows,

$$\rho(u_t + uu_x + wu_z) = -p_x + \mu \Delta u - \phi_x,$$

$$\rho \left( \frac{\varepsilon U_0^2}{h_0} W_T + \frac{\varepsilon U_0^2}{h_0} U W_X + \frac{\varepsilon U_0^2}{h_0} W W_Z \right) = -\frac{\mu U_0}{h_0^2} P_X + \mu \left( \frac{\varepsilon^2 U_0}{h_0} U_X + \frac{U_0}{h_0^2} U ZZ \right) - \frac{\mu U_0}{h_0^2} \Phi_X,$$

$$\frac{\varepsilon U_0}{h_0} U_T + U U_X + W W_Z = \frac{\mu U_0}{h_0^2} (-P_X + (\varepsilon^2 U_{XX} + U_{ZZ}) - \Phi_X),$$

$$\frac{\varepsilon U_0 h_0}{\mu} (U_T + U U_X + W W_Z) = (-P_X + (\varepsilon^2 U_{XX} + U_{ZZ}) - \Phi_X),$$

and

$$\rho(w_t + wu_x + ww_z) = -p_z + \mu \Delta w - \phi_z,$$

$$\rho \left( \frac{\varepsilon^2 U_0^2}{h_0} W_T + \frac{\varepsilon^2 U_0^2}{h_0} U W_X + \frac{\varepsilon^2 U_0^2}{h_0} W W_Z \right) = -\frac{\mu U_0}{h_0^2} P_Z + \mu \left( \frac{\varepsilon^3 U_0}{h_0^2} W_{XX} + \frac{U_0}{h_0^2} W ZZ \right) - \frac{\mu U_0}{h_0^2} \Phi_Z,$$

$$\frac{\varepsilon^2 U_0 h_0}{h_0} (W_T + U W_X + W W_Z) = \frac{\mu U_0}{h_0^2} (-P_Z + (\varepsilon^4 W_{XX} + \varepsilon^2 W_{ZZ}) - \Phi_Z),$$

$$\varepsilon^3 \rho U_0 h_0 \mu (W_T + U W_X + W W_Z) = (-P_Z + \varepsilon^2 (\varepsilon^2 W_{XX} + W_{ZZ}) - \Phi_Z).$$
The boundary conditions are nondimensionalized as below,

\[ \begin{align*}
  w &= 0, \quad u = 0, \quad \text{at } z = 0, \\
  \varepsilon U_0 W &= 0, \quad U_0 U = 0, \quad \text{at } Z = 0, \\
  W &= 0, \quad U = 0, \quad \text{at } Z = 0,
\end{align*} \]

with

\[ \begin{align*}
  w &= h_t + u h_x, \quad \text{at } z = h, \\
  \varepsilon U_0 W &= \varepsilon U_0 H_T + \varepsilon U_0 H x, \quad \text{at } Z = H, \\
  W &= H_T + U H_x, \quad \text{at } Z = H.
\end{align*} \]

In order to nondimensionalize the stress tensor at the free surface, we consider the normal and tangential components of \( T \cdot n \) separately. The normal and tangential vector at the surface can be expressed in terms of the free surface as

\[ \begin{align*}
  n &= \frac{\langle -h_x, 1 \rangle}{(1 + h_x^2)^{1/2}}, \quad t = \frac{\langle 1, h_x \rangle}{(1 + h_x^2)^{1/2}},
\end{align*} \] (2.12)

and the mean curvature, \( \kappa \), can be expressed in terms of \( h \) as

\[ \kappa = -\frac{h_{xx}}{(1 + h_x^2)^{3/2}}. \] (2.13)

We would now like to write the following vector equation in terms of its normal and tangential components.

\[ T \cdot n = (-\kappa \sigma + \Pi_0)n + \left( \frac{\partial \sigma}{\partial s} + \tau_0 \right) t, \quad \text{at } z = h. \]

Note that we are using a Newtonian stress tensor which is given by

\[ T = \begin{bmatrix} -p & 0 \\ 0 & -p \end{bmatrix} + \mu \begin{bmatrix} 2u_x & u_z + w_x \\ u_z + w_x & 2w_z \end{bmatrix} \] (2.14)

The normal component of this vector equation is given by

\[ \langle T \cdot n, n \rangle = (-\kappa \sigma + \Pi_0). \] (2.15)
First we will simplify the left hand side.

\[
a = (1 + h_x^2)^{1/2},
\]

\[
\langle T \cdot n, n \rangle = \frac{1}{a^2} \left[ -h_x \quad 1 \right] \left[ \begin{array}{cc} -p + 2 \mu u_x & \mu(u_z + w_x) \\ \mu(u_z + w_x) & -p + 2 \mu w_x \end{array} \right] \left[ \begin{array}{c} -h_x \\ 1 \end{array} \right]
\]

\[
= \frac{1}{a^2} \left[ -h_x \quad 1 \right] \left[ \begin{array}{c} -h_x(-p + 2 \mu u_x + \mu(u_z + w_x)) \\ -h_x \mu(u_z + w_x) + -p + 2 \mu w_x \end{array} \right]
\]

\[
= \frac{1}{a^2} \left( h_x^2(-p + 2 \mu u_x) - \mu h_x(u_z + w_x) - \mu h_x(u_z + w_x) + -p + 2 \mu w_x) \right),
\]

\[
= \frac{1}{a^2} \left( (1 + h_x^2)(-p) + 2 \mu(h_x^2 u_x + w_z) - 2 \mu h_x(u_z + w_x) \right),
\]

\[
= -p + 2\mu \left( (h_x^2 u_x + w_z) - h_x(u_z + w_x) \right).
\]

Next simplify the right hand side,

\[
(-\kappa \sigma + \Pi_0) = \frac{h_{xx}}{(1 + h_x^2)^{3/2}} \sigma + \Pi_0.
\]

This gives the following scalar equation,

\[
-p - \Pi_0 + \frac{2\mu}{1 + h_x^2} \left( (h_x^2 u_x + w_z) - h_x(u_z + w_x) \right) = \frac{h_{xx}}{(1 + h_x^2)^{3/2}} \sigma.
\] (2.16)

The tangential component is given by

\[
\langle T \cdot n, t \rangle = \left( \frac{\partial \sigma}{\partial s} + \tau_0 \right).
\] (2.17)
First simplify the left hand side,

\[
\left\langle \mathbf{T} \cdot \mathbf{n} , \mathbf{n} \right\rangle = \frac{1}{a^2} \begin{bmatrix} 1 & h_x \\ \mu (u_z + w_x) & -p + 2 \mu u_x \end{bmatrix} \begin{bmatrix} -h_x \\ 1 \end{bmatrix},
\]

\[
= \frac{1}{a^2} \begin{bmatrix} 1 & h_x \\ -h_x(-p + 2 \mu u_x) + \mu (u_z + w_x) & -h_x \mu (u_z + w_x) + -p + 2 \mu w_z \end{bmatrix},
\]

\[
= \frac{1}{a^2} (-h_x(-p + 2 \mu u_x) + \mu (u_z + w_x) - h_x^2 \mu (u_z + w_x) + h_x(-p + 2 \mu w_z)),
\]

\[
= \frac{1}{a^2} (-h_x(2 \mu u_x) + \mu (u_z + w_x) - h_x^2 \mu (u_z + w_x) + h_x(2 \mu w_z)),
\]

\[
= \frac{\mu}{a^2} (2h_x(w_z - u_x) + (1 - h_x^2)(u_z + w_x)).
\]

Next simplify the right hand side

\[
\left( \frac{\partial \sigma}{\partial s} + \tau_0 \right) = \frac{\partial \sigma}{\partial x} \frac{\partial x}{\partial s} + \tau_0,
\]

\[
= \frac{\partial \sigma}{\partial x} \frac{1}{(1 + h_x^2)^{1/2}} + \tau_0.
\]

This gives the following scalar equation,

\[
\mu (2h_x(w_z - u_x) + (1 - h_x^2)(u_z + w_x)) = \frac{\partial \sigma}{\partial x} (1 + h_x^2)^{1/2} + \tau_0 (1 + h_x^2). \tag{2.18}
\]
Lastly we will nondimensionalize these two equations,

\[-p - \pi + \frac{2\mu}{1 + h_x^2} \left( (h_x^2 u_x + w_z) - h_x(u_z + w_x) \right) = \frac{h_{xx}}{(1 + h_x^2)^{3/2}} \sigma,\]

\[-\frac{\mu U_0}{\varepsilon h_0} P - \frac{\mu U_0}{\varepsilon h_0} \Pi + \frac{2\mu}{1 + \varepsilon^2 H_X^2} \left( \left( \varepsilon^2 H_X^2 \frac{\varepsilon U_0}{h_0} U_X + \frac{\varepsilon U_0}{h_0} W_Z \right) - \varepsilon H_X \left( \frac{U_0}{h_0} U_Z + \frac{\varepsilon^2 U_0}{h_0} W_X \right) \right) = \frac{\varepsilon^2}{h_0} \left( 1 + \varepsilon^2 H_X^2 \right)^{3/2} \Sigma,\]

\[\left( -\frac{\mu U_0}{\varepsilon h_0} P - \frac{\mu U_0}{\varepsilon h_0} \Pi + \frac{\varepsilon \mu U_0}{h_0} \left( \frac{2}{1 + \varepsilon^2 H_X^2} \left( (\varepsilon^2 H_X^2 U_X + W_Z) - H_X (U_Z + \varepsilon^2 W_X) \right) \right) = \frac{\varepsilon \mu U_0}{h_0} \left( 1 + \varepsilon^2 H_X^2 \right)^{3/2} \Sigma,\]

\[\frac{\mu U_0}{\varepsilon h_0} \left( -P - \Pi + \frac{2\varepsilon^2}{1 + \varepsilon^2 H_X^2} \left( (\varepsilon^2 H_X^2 U_X + W_Z) - H_X (U_Z + \varepsilon^2 W_X) \right) \right) = \frac{\mu U_0}{\varepsilon h_0} \frac{\varepsilon^2 H_X^2}{\left( 1 + \varepsilon^2 H_X^2 \right)^{3/2}} \Sigma,\]

and

\[\mu (2h_x(w_z - u_x) + (1 - h_x^2)(u_z + w_x)) = \frac{\partial \sigma}{\partial x} (1 + h_x^2)^{1/2} + \tau_0 (1 + h_x^2),\]

\[\mu \left( 2\varepsilon H_X \left( \frac{\varepsilon U_0}{h_0} W_Z - \frac{\varepsilon U_0}{h_0} U_X \right) + (1 - \varepsilon^2 H_X^2) \left( \frac{U_0}{h_0} U_Z + \frac{\varepsilon^2 U_0}{h_0} W_X \right) \right) = \frac{\mu U_0}{h_0} \Sigma X \left( 1 + \varepsilon^2 H_X^2 \right)^{1/2} + \frac{\mu U_0}{h_0} \tau (1 + \varepsilon^2 H_X^2),\]

\[\frac{\mu U_0}{h_0} \left( 2\varepsilon^2 H_X (W_Z - U_X) + (1 - \varepsilon^2 H_X^2) (U_Z + \varepsilon^2 W_X) \right) = \frac{\mu U_0}{h_0} \left( \Sigma X \left( 1 + \varepsilon^2 H_X^2 \right)^{1/2} + \tau (1 + \varepsilon^2 H_X^2) \right),\]

\[2\varepsilon^2 H_X (W_Z - U_X) + (1 - \varepsilon^2 H_X^2) (U_Z + \varepsilon^2 W_X) = \Sigma X \left( 1 + \varepsilon^2 H_X^2 \right)^{1/2} + \tau (1 + \varepsilon^2 H_X^2).\]
The full nondimensional equations are thus

\[ U_X + W_Z = 0, \quad (2.19) \]
\[ \frac{\varepsilon U_0 \rho h_0}{\mu} (U_T + UU_X + WU_Z) = -P_X + (\varepsilon^2 U_{XX} + U_{ZZ}) - \Phi_X, \quad (2.20) \]
\[ \varepsilon^2 \rho U_0 h_0 \mu (W_T + UW_X + WW_Z) = -P_Z + \varepsilon^2 (U_{XX} + W_{ZZ}) - \Phi_Z, \quad (2.21) \]

at \( Z = 0 \)

\[ W = 0, \quad U = 0, \quad (2.22) \]

and at \( Z = H \),

\[ W = H_T + U_H, \quad (2.23) \]
\[ -P - \Pi + \frac{2\varepsilon^2}{1 + \varepsilon^2 H_X^2} \left( (\varepsilon^2 H_X^2 U_X + W_Z) - H_X (U_Z + \varepsilon^2 W_X) \right) = \frac{\varepsilon^2 H_X^2}{(1 + \varepsilon^2 H_X^2)^{3/2}} \Sigma, \quad (2.24) \]
\[ 2\varepsilon^2 H_X (W_Z - U_X) + (1 - \varepsilon^2 H_X^2) (U_Z + \varepsilon^2 W_X) = \Sigma X (1 + \varepsilon^2 H_X^2)^{1/2} + \tau (1 + \varepsilon^2 H_X^2). \quad (2.25) \]

We can now let \( \varepsilon \to 0 \) which results in

\[ U_X + W_Z = 0, \quad (2.26) \]
\[ P_X + \Phi_X = U_{ZZ}, \quad (2.27) \]
\[ P_Z + \Phi_Z = 0, \quad (2.28) \]

at \( Z = 0 \),

\[ W = 0, \quad U = 0, \quad (2.29) \]

\[ \quad (2.30) \]

and at \( Z = H \)

\[ W = H_T + U_H, \quad (2.31) \]
\[ -P - \Pi = \overline{\Sigma} H_{XX}, \quad (2.32) \]
\[ U_Z = \Sigma X + \tau. \quad (2.33) \]
Note that we assume that the surface tension is large, so that \( \bar{\Sigma} = \varepsilon^2 \Sigma = O(1) \). This is important in order to keep surface tension effects in the final equation.

Next we integrate the continuity equation over \( Z \),

\[
\int_0^H U_X + W_Z \, dZ = 0, \\
\int_0^H U_X \, dZ + W_{\mid Z=0} = 0, \\
\int_0^H U_X \, dZ + H_T + U H_X = 0, \\
H_T + \int_0^H U_X \, dZ + U H_X = 0, \\
H_T + \frac{\partial}{\partial X} \left( \int_0^H U \, dZ \right) = 0.
\]

Using the boundary conditions we can solve for an expression of \( U \) as follows

\[
\int_Z^H U_{ZZ} \, dZ = \int_Z^H P_X + \Phi_X \, dZ, \\
U_{\mid Z}^H = (P_X + \Phi_X)(H - Z), \\
(\tau + \Sigma_X) - U_Z = (P_X + \Phi_X)(H - Z), \\
\int_0^Z (\tau + \Sigma_X) - U_Z \, dZ = \int_0^Z (P_X + \Phi_X)(H - Z) \, dZ, \\
(\tau + \Sigma_X)Z - U_{\mid Z=0} = (P_X + \Phi_X) \left( HZ - \frac{1}{2} Z^2 \right), \\
(\tau + \Sigma_X)Z - U = (P_X + \Phi_X) \left( HZ - \frac{1}{2} Z^2 \right), \\
U = \left( \tau + \Sigma_X \right)Z + (P_X + \Phi_X) \left( \frac{1}{2} Z^2 - HZ \right).
\]

The boundary conditions also give an expression for \( P + \Phi \),

\[
P_Z + \Phi_Z = 0, \\
\int_Z^H P_Z + \Phi_Z \, dZ = 0, \\
P_{\mid Z=H} - P + \Phi_{\mid Z=H} - \Phi = 0, \\
-\Pi - \bar{\Sigma} H_{XX} - P + \Phi_{\mid Z=H} - \Phi = 0, \\
P + \Phi = \Phi_{\mid Z=H} - \Pi - \bar{\Sigma} H_{XX}.
\]
Plugging both of these into the integrated continuity equation gives,

\[ H_T + \frac{\partial}{\partial X} \left( \int_0^H U \, dZ \right) = 0, \]
\[ H_T + \frac{\partial}{\partial X} \left( \int_0^H (\tau + \Sigma X)Z + (P_X + \Phi_X) \left( \frac{1}{2}Z^2 - HZ \right) \, dZ \right) = 0, \]
\[ H_T + \left( \frac{1}{2}(\tau + \Sigma X)H^2 + (P_X + \Phi_X) \left( \frac{1}{6}H^3 - \frac{1}{2}H^3 \right) \right)_X = 0, \]
\[ H_T + \left( \frac{1}{2}(\tau + \Sigma X)H^2 - \frac{1}{3}(P + \Phi)_X H^3 \right)_X = 0, \]
\[ H_T + \left( \frac{1}{2}(\tau + \Sigma X)H^2 - \frac{1}{3}(\Phi|_{Z=H} - \Pi - \Sigma H_{XX})_X H^3 \right)_X = 0, \]
\[ H_T + \left( \frac{1}{2}(\tau + \Sigma X)H^2 - \frac{1}{3}(\Phi|_{Z=H} - \Pi)_X H^3 \right)_X = -(\Sigma H^3 H_{XXX})_X. \]

This is our final thin-film equation and taking all of the constants to be one gives

\[ H_T + (H^2 - H^3)_X = -(H^3 H_{XXX})_X. \] (2.34)

For the rest of the paper we will relabel the variables and use variables originally shown in equation 1.1,

\[ q_t + (q^2 - q^3)_x = -(q^3 q_{xxx})_x. \] (2.35)

2.2 Hyperbolic Convection Dynamics

Consider the thin-film model with only the hyperbolic convection term, that is

\[ q_t + (q^2 - q^3)_x = 0. \] (2.36)

Equations of the type \( q_t + f(q)_x = 0 \) are known as conservation laws, where \( f \) is called the flux function. The dynamics of this equation are determined by the flux function. In the system case, the conservation law is called hyperbolic at \( q \) if \( f'(q) \) is diagonalizable with real eigenvalues. In the scalar case, this reduces to having a real valued flux function, so this term is indeed hyperbolic. We also need to consider if our flux function is convex or nonconvex. It is known that \( f \) is nonconvex if \( f''(q) = 0 \) for some \( q \). In our case \( f''(q) = 2 - 6q \), so \( f''(q) = 0 \) at \( q = \frac{1}{3} \). We can also see in Figure 2.2 that there is an inflection point at \( q = \frac{1}{3} \). Nonconvex flux functions can exhibit more
complex behavior than convex flux functions. Nonconvex flux functions can involve compound waves including rarefaction-shocks. See Leveque [16] for more details.

Another interesting feature of hyperbolic equations is that they have a finite wavespeed. The wavespeed of the conservation law is determined by $f'(q)$. In our case, $f'(q) = 2q - 3q^2$, so the wavespeed can be as large as $\frac{1}{3}$. Having a finite wavespeed can be very useful for numerical methods, as this limits the speed of propagation of information. Each point in the domain has a finite radius of influence for each future time.

### 2.3 Parabolic Dynamics

Now consider the thin-film model with only the parabolic term, that is

\[ q_t = -(q^3 q_{xxx})_x. \]  

(2.37)

Using the product rule of differentiation this is equivalent to

\[ q_t = -(3q^2 q_x) q_{xxx} - q^3 q_{xxxx}. \]  

(2.38)

The dynamics of the third and fourth derivative terms can be analyzed with the Fourier transform.

Consider, $q_t = -q_{xxx}$, and take the Fourier transform of both sides, which results in

\[ \mathcal{F}(q)_t = -i\xi^3 \mathcal{F}(q). \]  

(2.39)
Solving this ordinary differential equation (ODE) gives

\[ \mathcal{F}(q) = e^{-i\xi q}. \]  \hspace{1cm} (2.40)

We know that this type of solution has a dispersive dynamics.

Similarly consider, \( q_t = -q_{xxxx} \), and take the Fourier transform of both sides. This gives the following ordinary differential equation

\[ \mathcal{F}(q)_t = -\xi^4 \mathcal{F}(q). \]  \hspace{1cm} (2.41)

The solution to this ODE is

\[ \mathcal{F}(q) = e^{-\xi^4 t}. \]  \hspace{1cm} (2.42)

This is a function that decays rapidly in time, and the higher the wavenumber, \( \xi \), the faster the diffusion.

Therefore we expect this parabolic term to exhibit dispersive and diffusive dynamics. The dynamics are also nonlinearly coupled together making the behavior even more difficult to predict. The one thing that is clear is that the overall model is very stiff. The vast time scale difference between the finite wavespeed of the hyperbolic convection and the rapid decay of the nonlinear diffusion, make this model very stiff. Our numerical method must be able to handle this stiffness in a way that is not too computationally intensive.
CHAPTER 3. NUMERICAL METHODS

3.1 Implicit-Explicit Runge-Kutta Scheme

The thin-film model we are trying to solve is a very stiff equation due to the high order derivatives. In order to take reasonable timesteps despite this stiffness we use implicit-explicit (IMEX) Runge-Kutta schemes to propagate our solution through time. The IMEX Runge-Kutta scheme propagates time for ordinary differential equations or systems of equations of the form

\[ q_t = F(t, q) + G(t, q), \]  \hspace{1cm} (3.1)

where \( F \) is solely handled explicitly, but \( G \) needs to be solved implicitly. A single step of the Runge-Kutta IMEX scheme is computed as follows,

\[ q^{n+1} = q^n + \Delta t \sum_{i=1}^{s} (b'_i F(t_i, u_i)) + \Delta t \sum_{i=1}^{s} (b_i G(t_i, u_i)), \]  \hspace{1cm} (3.2)

\[ u_i = q^n + \Delta t \sum_{j=1}^{i-1} (a'_{ij} F(t_j, u_j)) + \Delta t \sum_{j=1}^{i} (a_{ij} G(t_j, u_j)) \]  \hspace{1cm} (3.3)

\[ t_i = t^n + c_i \Delta t \]  \hspace{1cm} (3.4)

where the coefficients \( a_{ij}, b_i, c_i, a'_{ij}, b'_i, \) and \( c'_i \) are set in the double Butcher tableau

\[
\begin{array}{c|c|c}
  c' & a' & c \\
  \hline
  (b')^T & a & b^T
\end{array}
\]

Specifically we used the IMEX schemes first introduced by Pareschi and Russo [24, 25]. These schemes are strong stability preserving (SSP) in the sense of Gottlieb et al. [12] and they are designed for stiff systems. The double Butcher tableaus are shown below for the one stage, L-stable, 1st order method:

\[
\begin{array}{c|c|c}
  0 & 0 & 1 \\
  \hline
  1 & 1 & \end{array}
\]
the three stage, 2nd order method:

\[
\begin{array}{cccc|ccc}
0 & 0 & 0 & \frac{1}{2} & \frac{1}{2} & 0 & 0 \\
0 & 0 & 0 & 0 & -\frac{1}{2} & \frac{1}{2} & 0 \\
1 & 0 & 1 & 0 & 1 & \frac{1}{2} & \frac{1}{2} \\
\hline
0 & \frac{1}{2} & \frac{1}{2} & 0 & 1 & \frac{1}{2} & \frac{1}{2}
\end{array}
\]

and the four stage, L-stable, third order method:

\[
\begin{array}{cccc|cccc}
0 & 0 & 0 & 0 & 0 & \alpha & \alpha & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & -\alpha & \alpha & 0 & 0 \\
1 & 0 & 1 & 0 & 0 & 1 & 0 & 1-\alpha & \alpha & 0 \\
\frac{1}{2} & 0 & \frac{1}{4} & \frac{1}{4} & 0 & \frac{1}{2} & \beta & \eta & \zeta & \alpha \\
\hline
0 & \frac{1}{6} & \frac{1}{6} & \frac{2}{3} & 0 & 1 & \frac{1}{6} & \frac{1}{6} & \frac{2}{3}
\end{array}
\]

where

\[
\alpha = 0.24169426078821, \\
\beta = 0.06042356519705, \\
\eta = 0.1291528696059, \\
\zeta = \frac{1}{2} - \beta - \eta - \alpha.
\]

For our thin-film model \(F\) and \(G\) will be the spatial discretizations of the hyperbolic convection and parabolic terms respectively, that is

\[
F(t, q) = -(q^2 - q^3)_x \quad (3.5)
\]

\[
G(t, q) = -(q^3 q_{xxx})_x. \quad (3.6)
\]

Since \(G\) is the stiffest part of our model, we would like to handle this term implicitly so that our time step is less restricted. Also handling \(F\) explicitly allows us to capture the nonlinear behavior without a nonlinear solve.
3.2 Space Discretization

We chose to use the discontinuous Galerkin Method to discretize our equation in space. First let \( \{x_{j+1/2}\}_{0}^{N} \) partition the domain, \([a,b]\), and denote each interval as \( I_{j} = [x_{j-1/2},x_{j+1/2}] \) with \( \Delta x_{j} = x_{j+1/2} - x_{j-1/2} \). The discontinuous Galerkin solution then exists in the following finite dimensional space,

\[
V_{h}^{k} = \left\{ v \in L^{1}([a,b]) : v|_{I_{j}} \in P_{k}(I_{j}), j = 1, \ldots, N \right\}, \tag{3.7}
\]

where \( P_{k}(I_{j}) \) denotes the set of polynomials of degree \( k \) or less on \( I_{j} \).

3.2.1 Modal Discontinuous Galerkin for Hyperbolic Convection

First we consider the continuous operator \( F(t,q) = -(q^{2} - q^{3})_{x} \). Note that \( F : [0,T] \times C^{1}(a,b) \to C^{0}(a,b) \), and we would like to form an approximation, \( F_{h} \) to this continuous operator, such that \( F_{h} : [0,T] \times V_{h}^{k} \to V_{h}^{k} \). In order to do this, consider the weak formulation of the continuous operator, \( F \). The weak form requires finding \( F(t,q) \in C^{0}(a,b) \) such that

\[
\int_{a}^{b} F(t,q)v(x) \, dx = -\int_{a}^{b} (q^{2} - q^{3})_{x} v(x) \, dx \tag{3.8}
\]

for all smooth functions, \( v \in C^{\infty}(a,b) \). The discontinuous Galerkin approximation is formed by replacing these function spaces with the DG space, \( V_{h}^{k} \). This gives that the weak formulation of the approximation \( F_{h} \) is to find \( F_{h}(t,q_{h}) \in V_{h}^{k} \) such that

\[
\int_{a}^{b} F_{h}(t,q_{h})v_{h}(x) \, dx = -\int_{a}^{b} (q_{h}^{2} - q_{h}^{3})_{x} v_{h}(x) \, dx \tag{3.9}
\]

for all \( v_{h} \in V_{h}^{k} \). Using integration by parts this is equivalent to finding \( F_{h}(t,q_{h}) \in V_{h}^{k} \) such that

\[
\int_{I_{j}} F_{h}(t,q_{h})v_{h}(x) \, dx = \int_{I_{j}} (q_{h}^{2} - q_{h}^{3})_{x} v_{h}(x) \, dx \tag{3.10}
\]

\[
- \hat{f}(q_{h})_{j+1/2}v_{h}(x_{j+1/2}) + \hat{f}(q_{h})_{j-1/2}v_{h}(x_{j-1/2})
\]

for \( j = 1, \ldots, N \) and for all \( v_{h} \in V_{h}^{k} \). Since \( q_{h} \) is discontinuous at the cell interfaces \( x_{j\pm1/2} \), some numerical flux \( \hat{f} \) must be chosen.
We chose to use the Local Lax-Friedrich’s numerical flux, that is
\[
\hat{f}(q_l, q_r) = \frac{1}{2} (f(q_l) + f(q_r) - \lambda_{\text{max}}(q_r - q_l))
\]
(3.11)
where \(f\) is the flux function, \(q_l\) and \(q_r\) and the left and right states at the interface, and \(\lambda_{\text{max}}\) is the locally maximum wavespeed, that is
\[
\lambda_{\text{max}} = \max_{q \in [\nu, \mu]} \{f'(q)\},
\]
(3.12)
where \(\nu = \min\{q_l, q_r\}\) and \(\mu = \max\{q_l, q_r\}\). In our case \(f(q) = q^2 - q^3\). At the interface \(x_{j+1/2}\) we have \(q_l = q^{-}_h(x_{j+1/2})\) and \(q_r = q^{+}_h(x_{j+1/2})\).

### 3.2.2 Local Discontinuous Galerkin Scheme for Parabolic Term

Next we look at the continuous operator \(G(t, q) = -\left(q^3q_{xxx}\right)_x\). In discretizing this operator we follow the local discontinuous Galerkin (LDG) method first introduced by Cockburn and Shu [10] for convection-diffusion systems. We first introduce three auxilliary variables, \(r, s, u\), and rewrite the equation as the following system,
\[
\begin{align*}
    r &= q_x, \\
    s &= r_x, \\
    u &= s_x, \\
    G(t, q) &= -\left(q^3u\right)_x.
\end{align*}
\]
(3.13, 3.14, 3.15, 3.16)
The weak form of this system is solved by finding functions \(r, s, u, G\) such that
\[
\begin{align*}
    \int_a^b r(x)w(x)\,dx &= \int_a^b q_x(x)w(x)\,dx, \\
    \int_a^b s(x)y(x)\,dx &= \int_a^b r_x(x)y(x)\,dx, \\
    \int_a^b u(x)z(x)\,dx &= \int_a^b s_x(x)z(x)\,dx, \\
    \int_a^b G(t, q)v(x)\,dx &= -\int_a^b \left(q^3(x)u(x)\right)_x v(x)\,dx,
\end{align*}
\]
(3.17, 3.18, 3.19, 3.20)
for all smooth functions \( w, y, z, v \in C^\infty(a, b) \). The LDG method arrives from applying the standard DG method to each of these equations. That is we replace the continuous function spaces with the discontinuous finite dimensional DG space, \( V_k^h \). The approximate operator \( G_h \) becomes the process of finding \( r_h, s_h, u_h, G_h(t, q_h) \in V_h \) such that for all test functions \( v_h, w_h, y_h, z_h \in V_h \) the following equations are satisfied

\[
\int_a^b r_h w_h \, dx = \int_a^b (q_h)_x w_h \, dx, \tag{3.21}
\]
\[
\int_a^b s_h y_h \, dx = \int_a^b (r_h)_x y_h \, dx, \tag{3.22}
\]
\[
\int_a^b u_h z_h \, dx = \int_a^b (s_h)_x z_h \, dx, \tag{3.23}
\]
\[
\int_a^b G_h(t, q_h) v_h \, dx = -\int_a^b (q_h^3 u_h)_x v_h \, dx, \tag{3.24}
\]
given \( t \) and \( q_h \in V_k^h \). This is equivalent to the following equations for all \( j \), if we use integration by parts,

\[
\int_{I_j} r_h w_h \, dx = \left( (\hat{q}_h w_h^+)_{j+1/2} - (\hat{q}_h w_h^-)_{j-1/2} \right) - \int_{I_j} q_h(w_h)_x \, dx, \tag{3.25}
\]
\[
\int_{I_j} s_h y_h \, dx = \left( (\hat{r}_h y_h^-)_{j+1/2} - (\hat{r}_h y_h^+)_{j-1/2} \right) - \int_{I_j} r_h(y_h)_x \, dx, \tag{3.26}
\]
\[
\int_{I_j} u_h z_h \, dx = \left( (\hat{s}_h z_h^-)_{j+1/2} - (\hat{s}_h z_h^+)_{j-1/2} \right) - \int_{I_j} s_h(z_h)_x \, dx, \tag{3.27}
\]
\[
\int_{I_j} G_h(t, q_h) v_h \, dx = -\left( (\hat{q}_h^3 u_h v_h^-)_{j+1/2} - (\hat{q}_h^3 u_h v_h^+)_{j-1/2} \right) + \int_{I_j} q_h^3 u_h(v_h)_x \, dx, \tag{3.28}
\]

where \( \hat{q}, \hat{r}, \hat{s}, \hat{q}_h^3 u \) are suitably chosen numerical fluxes. A common choice of numerical fluxes are the so-called alternating fluxes, shown below

\[
\hat{q}_h = q_h^-, \tag{3.29}
\]
\[
\hat{r}_h = r_h^+, \tag{3.30}
\]
\[
\hat{s}_h = s_h^-, \tag{3.31}
\]
\[
\hat{q}_h^3 u_h = (q^3 u)_h^+. \tag{3.32}
\]
These numerical fluxes are one-sided fluxes that alternate sides for each derivative. They are chosen to make this method stable, and they also allow for the auxilliary variables to be locally solved in terms of $q_h$, hence where the local discontinuous Galerkin method gets its name.

### 3.3 Nonlinear Solver

Using these discretizations in the Runge-Kutta IMEX scheme, requires solving the nonlinear system,

$$ u_i - \Delta t a_{ij} G_h(t_j, u_j) = q^n + \Delta t \sum_{j=1}^{i-1} (a'_{ij} F_h(t_j, u_j)) + \Delta t \sum_{j=1}^{i-1} (a_{ij} G_h(t_j, u_j)), \quad (3.33) $$

for $u_i$. The standard approach to solving this system would be to use a Newton iteration, however that would require the Jacobian of the operator $I - \Delta t a_{ij} G_h$, which would be relatively intractable. Therefore we chose to linearize this operator and use a Picard iteration instead, which does not require the Jacobian.

Suppose we are trying to solve the nonlinear equation $L(u) = b$, where $L'(v, u)$ is the nonlinear operator linearized about $v$ acting on $u$. The Picard iteration for solving this nonlinear equation starts with some initial guess $u_0$. The next solution is found by solving the following linear problem,

$$ L'(u_i, u_{i+1}) = b. \quad (3.34) $$

In other words the next iteration is found by solving the problem linearized about the the previous iteration.

We linearize the operator $G_h(t, q_h)$ by first linearizing $G(t, q)$. The continuous operator linearized about $v$ is given by

$$ G'(v, t, q) = -(v^3 q_{xxx})_x. \quad (3.35) $$

The linearized discrete operator is now just the LDG method applied to this linearized continuous operator.

We find that the Picard iteration approach provides good results with relatively few iterations. In fact in Section 4.1 we show that we can achieve first, second, and third order accuracy with
only one, two, and three iterations respectively per stage. This approach is more tractable than a
Newton iteration and it converges quickly to the nonlinear solution.

These methods were implemented in a python code called PyPacked [18] developed by the
authors. PyPacked uses the packages NumPy [22] and SciPy [27] extensively. In particular
they are used to solve linear systems of equations and to integrate numerically. All of the examples
and tests shown in Chapter 4 were done using this code. Key parts of the code are shown in the
Appendix.
CHAPTER 4. RESULTS

4.1 Manufactured Solution

In order to confirm the order of accuracy of our method, we use the method of manufactured solutions. The method of manufactured solutions picks an exact solution \( \hat{q} \) and then adds a source term to the initial partial differential equation to make \( \hat{q} \) the true solution to the new differential equation. So we actually solve

\[
q_t + (q^2 - q^3)_x = -(q^3 q_{xxx})_x + s,
\]

where

\[
s = q_t + (\hat{q}^2 - \hat{q}^3)_x + (\hat{q}^3 \hat{q}_{xxx})_x.
\]

This new PDE’s exact solution is now \( \hat{q} \). We chose

\[
\hat{q} = 0.1 \times \sin(2\pi/20.0 \times (x - t)) + 0.15,
\]

on \( x \in [0, 40] \) with periodic boundary conditions to be our manufactured solution. This manufactured solution is chosen so that our flux \( q^2 - q^3 \) is in its convex region.

We solve this problem until \( t = 5.0 \) with constant, linear, and quadratic basis polynomials, that is with \( k = 0, 1, \) and \( 2 \). The time step size for these simulations is determined by the Courant–Friedrichs–Lewy (CFL) condition. The CFL condition states that

\[
\Delta t = \nu \frac{\Delta x}{\lambda},
\]

where \( \lambda \) is the wavespeed and \( \nu \) is the CFL number. For hyperbolic problems solved with explicit Runge-Kutta time-stepping, \( \nu \), typically follows the pattern \( \frac{1}{2n-1} \) where \( n \) is the order of the method. For this example the wavespeed is 1, and the timestep is chosen just smaller than the typical CFL number. Specifically the timesteps were \( \Delta t = 0.9\Delta x \), \( \Delta t = 0.2\Delta x \), and \( \Delta t = 0.1\Delta x \) for first, second and third order respectively.
Table 4.1 shows the error for each simulation as the mesh is refined. The table also shows the rate of convergence to the true solution, and in each case the rate of convergence approaches the expected order. Note that the first, second, and third order methods only require one, two, and three Picard iterations respectively.

The relative $L^2$ error is defined as

$$E = \|c(x)\|_{L^2(a,b)} = \sqrt{\frac{\int_a^b (\hat{q} - q_h)^2 \, dx}{\int_a^b \hat{q}^2 \, dx}}$$

(4.5)

Since we are using an orthonormal basis of $V_h^k$, there is an easier way to compute this error numerically. The basis consists of functions $\phi_j^l$, that are polynomials of degree $l$ on $I_j$ and are zero everywhere else for $j = 1, \ldots, N$ and $l = 0, \ldots, k$. By definition of orthonormality we have that

$$\int_a^b \phi_k^i \phi_j^l \, dx = \delta_{kl} \delta_{ij}$$

(4.6)

where $\delta$ is the Kronecker delta. In other words on cell $I_j$, $\phi_j^l$ is a Legendre polynomial. Given this basis we can express our numerical solution, $q_h \in V_h^k$, as

$$q_h|_{I_j} = \sum_{l=0}^k (Q_j^l \phi_j^l),$$

(4.7)

for all $j$, where $Q_j^l$ are constant coefficients.

In order to maintain leading order accuracy we project our exact solution, $\hat{q}$, onto the space $V_h^{k+1}$, that is with one degree higher polynomials. Let $\hat{q}_h$ denote this projection. We use the same orthonormal basis with the addition of $\phi_j^{k+1}$ on each cell. In this case we have the following representations

$$\hat{q}_h|_{I_j} = \sum_{l=0}^{k+1} \left( \hat{Q}_j^l \phi_j^l(x) \right) \text{ and } q_h|_{I_j} = \sum_{l=0}^{k+1} \left( Q_j^l \phi_j^l(x) \right),$$

(4.8)

where $Q_j^{k+1}$ will be zero. Substituting these representations into the $L^2$ error, equation (4.5), and using the orthonormality property gives the following expression for the error,

$$E_N = \sqrt{\frac{\sum_{j=1}^N \sum_{l=0}^{k+1} \left( \hat{Q}_i - Q_i \right)^2}{\sum_{j=1}^N \sum_{l=0}^{k+1} \hat{Q}_i^2}}.$$

(4.9)
Table 4.1 Convergence table with a constant, linear, quadratic polynomial bases. Nonlinear solve uses one, two, or three Picard iterations respectively. CFL = 0.9, 0.2, 0.1 respectively.

<table>
<thead>
<tr>
<th>n</th>
<th>1st Order error</th>
<th>order</th>
<th>2nd Order error</th>
<th>order</th>
<th>3rd Order error</th>
<th>order</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>0.136</td>
<td>—</td>
<td>7.34 x 10^{-3}</td>
<td>—</td>
<td>5.29 x 10^{-4}</td>
<td>—</td>
</tr>
<tr>
<td>40</td>
<td>0.0719</td>
<td>0.91</td>
<td>1.99 x 10^{-3}</td>
<td>1.89</td>
<td>5.38 x 10^{-5}</td>
<td>3.30</td>
</tr>
<tr>
<td>80</td>
<td>0.0378</td>
<td>0.93</td>
<td>5.60 x 10^{-4}</td>
<td>1.83</td>
<td>7.47 x 10^{-6}</td>
<td>2.85</td>
</tr>
<tr>
<td>160</td>
<td>0.0191</td>
<td>0.99</td>
<td>1.56 x 10^{-4}</td>
<td>1.85</td>
<td>9.97 x 10^{-7}</td>
<td>2.91</td>
</tr>
<tr>
<td>320</td>
<td>0.00961</td>
<td>0.99</td>
<td>3.98 x 10^{-5}</td>
<td>1.97</td>
<td>1.26 x 10^{-7}</td>
<td>2.98</td>
</tr>
<tr>
<td>640</td>
<td>0.00483</td>
<td>0.99</td>
<td>1.00 x 10^{-5}</td>
<td>1.99</td>
<td>1.58 x 10^{-8}</td>
<td>3.00</td>
</tr>
<tr>
<td>1280</td>
<td>0.00242</td>
<td>1.00</td>
<td>2.50 x 10^{-6}</td>
<td>2.00</td>
<td>1.98 x 10^{-9}</td>
<td>3.00</td>
</tr>
</tbody>
</table>

This expression is equivalent to equation (4.5) to leading order. The order of convergence is then estimated from these errors as

\[
\text{order} \approx \log_2 \left( \frac{E_N}{E_{2N}} \right). \tag{4.10}
\]

### 4.2 Traveling Waves

In this section we showcase several numerical examples that demonstrate the traveling wave profiles of equation (1.1). The traveling wave profiles differ from the standard hyperbolic wave profile in several ways. They differ in that they may not be unique and may include undercompressive shocks. These examples were first shown in [4] with first order accuracy.

In these examples, we use a moving reference frame to keep the mesh size reasonable. This is done by actually simulating the following equation,

\[
q_t + (q^2 - q^3 - sq)_x = -(q^3 q_{xxx})_x \tag{4.11}
\]

where \( s \) is the Rankine-Hugoniot wavespeed of the original numerical flux, \( f \)

\[
s = \frac{f(q_t) - f(q_r)}{q_t - q_r} = q_t + q_r - (q^2_t q_r + q^2_r). \tag{4.12}
\]

This modified equation will have zero wavespeed in most cases and simulates the original PDE in a moving reference frame.
These examples consider Riemann Problems with different left and right states. Depending on the left and right states they are several different wave profiles. For these examples we will fix the right state and vary the left state. The wave profiles would be qualitatively equivalent for different values for the right state, however the values of the left state would also need to change accordingly.

4.2.1 Case 1: Unique Weak Lax Shock

Consider a Riemann Problem with left state, \( q_l = 0.3 \), and right state, \( q_r = 0.1 \). We will use the following smoothed out profile as an initial condition for this problem

\[
q_0(x) = (\tanh(-x) + 1) \frac{q_l - q_r}{2} + q_r.
\]

For this initial condition the numerical solution approaches a steady wave profile with a unique Lax type shock. Figure 4.1 shows a plot of the solution with this initial condition after enough time for the solution to hit its steady state. Note that this behavior persists for all \( q_l \) up to some bound which depends on \( q_r \).

4.2.2 Case 2: Multiple Lax Shocks

If the left state is increased to \( q_l = 0.3323 \), then the wave profile is no longer unique. In this case the steady traveling wave depends on the initial conditions.
For the following initial conditions,

\[ q_0(x) = (\tanh(-x) + 1) \frac{q_l - q_r}{2} + q_r, \quad (4.14) \]

the behavior is the same as in case 1. The numerical solution for this initial condition is shown in Figure 4.2.

Now consider the following initial conditions,

\[ q_0(x) = \begin{cases} 
((0.6 - q_l)/2) \tanh(x) + ((0.6 + q_l)/2) & x < 5 \\
-((0.6 - q_r)/2) \tanh(x - 10) + ((0.6 + q_r)/2) & x > 5
\end{cases}. \quad (4.15) \]

The reader might expect this initial condition to approach the traveling wave shown earlier as it has the same far field boundary values, however this is not the case. The traveling wave profile that is approaches is shown in Figure 4.2.

These are not the only possible wave profiles possible with these left and right states. Figure 4.2 part (c) shows the result for the following initial condition,

\[ q_0(x) = \begin{cases} 
((0.6 - q_l)/2) \tanh(x) + ((0.6 + q_l)/2) & x < 10 \\
-((0.6 - q_r)/2) \tanh(x - 20) + ((0.6 + q_r)/2) & x > 10
\end{cases}. \quad (4.16) \]

This initial condition has a larger hump then equation (4.15), and the traveling wave reflects this aspect. In this case the traveling wave is not a steady wave, and there are in fact two shocks. The right shock is an undercompressive shock and the left shock is a traditional compressive shock. Both shocks travel slower than the moving reference frame. They also travel at different speeds from one another so the wave profile changes over time.

4.2.3 Case 3: Undercompressive Double Shock

For the left state in the next regime, there is again a unique shock profile for all initial conditions. However this shock profile is not the single Lax shock seen in Case 1. With the following initial conditions,

\[ q_0(x) = (\tanh(-x) + 1) \frac{q_l - q_r}{2} + q_r, \quad (4.17) \]
Figure 4.2 Case 2: Distinct Lax Shocks. Panels (a) and (b) show two distinct steady travelling waves formed from different initial conditions. Panel (c) shows an unsteady wave with a double shock structure from a third initial condition.
where $q_l = 0.4$ and $q_r = 0.1$, Figure 4.3 shows a double shock structure. Similar to Figure 4.2 part (c), we see a undercompressive shock on the right and a Lax shock on the left.

### 4.2.4 Case 4: Rarefaction-Undercompressive Shock

The final traveling wave structure appears when the left state is greater than the undercompressive shock height. In this case we see a rarefaction wave along with the undercompressive shock. Figure 4.4 shows the numerical solution for initial condition

$$q_0(x) = (\tanh(-x + 110) + 1)\frac{q_l - q_r}{2} + q_r,$$

where $q_l = 0.8$ and $q_r = 0.1$. Note that the rarefaction wave and undercompressive shock are traveling at different speeds so they separate from each other.
Figure 4.4 Case 4: Rarefaction-undercompressive shock.
CHAPTER 5. CONCLUSION

This paper has given a discontinuous Galerkin method for solving a Thin-Film model. The nonlinear convection term, \((q^2 - q^3)_x\), is discretized with the modal discontinuous Galerkin method and the nonlinear parabolic term, \(-(q^3 q_{xxx})_x\), is discretized with the local discontinuous Galerkin method. The nonlinear parabolic term is very stiff and would force a very tight time restriction, therefore implicit-explicit Runge-Kutta methods are used for solving the semi-discretized system. The implicit-explicit time-stepping allows for reasonable time steps to be taken given the stiffness of the problem. As the parabolic term is nonlinear, this still requires solving a nonlinear system, however we are able to solve this nonlinear problem without a Newton iteration. In fact a Picard Iteration was used, and it was demonstrated that only a minimal number of iterations were required to achieve high accuracy. The number of Picard Iterations needed is less than or equal to the order of the LDG discretization. Using this method we have demonstrated up to third order accuracy with the method of manufactured solutions. Also we have showcased several numerical examples first given by Bertozzi. These examples show that our method preserves the nonlinear behavior of the Thin-Film model that was first shown in those examples.
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[19] V. Ludviksson and E. Lightfoot. The dynamics of thin liquid films in the presence of surface-


APPENDIX. PYDOGPACK CODE

The code for the implicit-explicit Runge-Kutta scheme is shown below.

```python
class IMEXRungeKutta:
    # IMEX Runge Kutta scheme for solving \( q_{n+1} = F(t, q) + G(t, q) \)
    # where \( F \) is solved explicitly and \( G \) is solved implicitly
    # these schemes are represented by two Butcher Tableaus
    # \( a, b, c \) for operator \( F \)
    # \( a_p, b_p, c_p \) for operator \( G \)
    # \( y_{n+1} = y_n + \Delta t \sum_{i=1}^{s} \{ b_p_i F(t^n + c_i \Delta t, u_i) \} + \Delta t \sum_{i=1}^{s} \{ b_i G(t^n + c_i \Delta t, u_i) \} \)
    # \( u_i = y_n + \Delta t \sum_{j=1}^{i-1} \{ a_p_{ij} F(t^n + c_j \Delta t, u_j) \} + \Delta t \sum_{j=1}^{i-1} \{ a_{ij} G(t^n + c_j \Delta t, u_j) \} \)
    # means solving
    # \( u_i - \Delta t a_{ii} G(t^n + c_i \Delta t, u_i) = rhs \)
    # \( rhs = y_n + \Delta t \sum_{j=1}^{i-1} \{ a_p_{ij} F(t^n + c_j \Delta t, u_j) \} + \Delta t \sum_{j=1}^{i-1} \{ a_{ij} G(t^n + c_j \Delta t, u_j) \} \)
    def __init__(self, a, b, c, ap, bp, cp):
        self.a = a
        self.b = b
        self.c = c
        self.ap = ap
        self.bp = bp
        self.cp = cp
        self.num_stages = self.c.size

def time_step(self, q_old, t_old, delta_t, explicit_operator, implicit_operator, solve_operator):
    stages = []
    for i in range(self.num_stages):
        stages.append(
            self.stage(
                q_old, t_old, delta_t, explicit_operator, implicit_operator, solve_operator, stages, i,
            )
        )
```
q_new = q.old.copy()
for i in range(self.num_stages):
    if self.b[i] != 0.0:
        time = t.old + self.c[i] * delta_t
        q_new += delta_t * self.b[i] * implicit_operator(time, stages[i])
    if self.bp[i] != 0.0:
        time = t.old + self.cp[i] * delta_t
        q_new += delta_t * self.bp[i] * explicit_operator(time, stages[i])
return q_new

def stage(
    self,
    q.old,
    t.old,
    delta_t,
    explicit_operator,
    implicit_operator,
    solve_operator,
    stages,
    stage_num,
):
    # rhs = y' + \sum_{j = 1}^{i-1} \{ a_{ij} F(t' + c_{j} \ delta_t, u_{j}) \} + \delta t \ \sum_{j = 1}^{i-1} \{ a_{ij} G(t' + c_{j} \ delta_t, u_{j}) \}
    stage_rhs = q.old.copy()
    for j in range(stage_num):
        if self.a[stage_num, j] != 0.0:
            time = t.old + self.c[j] * delta_t
            stage_rhs += (delta_t * self.a[stage_num, j] * implicit_operator(time, stages[j]))
        if self.ap[stage_num, j] != 0.0:
            time = t.old + self.cp[j] * delta_t
            stage_rhs += (delta_t * self.ap[stage_num, j] * explicit_operator(time, stages[j]))
    # d q + c G(t, f q) = rhs
    # solve_function(q, c, f, t, rhs)
    # u_{i} = \delta t a_{ii} G(t' + c_{i} \ delta_t, u_{i}) = rhs
    time = t.old + self.c[stage_num] * delta_t
    e = -1.0 * delta_t * self.a[stage_num, stage_num]
    return solve_operator(1.0,
        e,
        time,
        stage_rhs,
        q.old,
        t.old,
        delta_t,
        implicit_operator,
        stages,
class IMEX1(IMEXRungeKutta):
    def __init__(self):
        ap = np.array([[0.0]])
bp = np.array([1.0])
cp = np.array([0.0])
a = np.array([[1.0]])
b = np.array([1.0])
c = np.array([1.0])
        IMEXRungeKutta.__init__(self, a, b, c, ap, bp, cp)

class IMEX2(IMEXRungeKutta):
    def __init__(self):
        ap = np.array([np.array([[0.0, 0.0, 0.0], [0.0, 0.0, 0.0], [0.0, 1.0, 0.0]])])
bp = np.array([0.0, 0.5, 0.5])
cp = np.array([0.0, 0.0, 1.0])
a = np.array([np.array([[0.5, 0.0, 0.0], [-0.5, 0.5, 0.0], [0.0, 0.5, 0.5]])])
b = np.array([0.0, 0.5, 0.5])
c = np.array([0.5, 0.0, 1.0])
        IMEXRungeKutta.__init__(self, a, b, c, ap, bp, cp)

class IMEX3(IMEXRungeKutta):
    def __init__(self):
        alpha = 0.24169426078821
    beta = 0.06042356519705
    eta = 0.1291528696059
    zeta = 0.5 - beta - eta - alpha
        ap = np.array([np.array([[0.0, 0.0, 0.0, 0.0], [0.0, 0.0, 0.0, 0.0],
                                  [0.0, 1.0, 0.0, 0.0], [0.0, 0.25, 0.25, 0.0]],
                        np.array([[alpha, 0.0, 0.0, 0.0], [-alpha, alpha, 0.0, 0.0],
                                   [0.0, 1.0 - alpha, alpha, 0.0], [alpha, 0.0, 0.0, 0.0]]),
                           [0.0, 1.0 / 6.0, 1.0 / 6.0, 2.0 / 3.0])])
bp = np.array([0.0, 0.0, 1.0])
cp = np.array([0.0, 0.0, 1.0, 0.5])
a = np.array([0.0, 0.0, 1.0])
The discontinuous Galerkin method in operator and matrix form is shown below.

```python
def evaluate_weak_form(
    dg_solution,
    numerical_fluxes,
    quadrature_function,
    vector_left,
    vector_right,
    source_quadrature_function=None,
):
    mesh_ = dg_solution.mesh
    basis_ = dg_solution.basis
    num elems = mesh_.num elems
    num_basis_cpts = basis_.num basis_cpts
    transformed_solution = solution.DGSolution(
        np.zeros((num elems, num_basis_cpts)), basis_, mesh_
    )
    for i in range(num elems):
        left_face_index = mesh_.elems_to_faces[i, 0]
        right_face_index = mesh_.elems_to_faces[i, 1]
        transformed_solution[i, :] = (1.0 / mesh_.elem_metrics[i] * (quadrature_function(i) - (numerical_fluxes[right_face_index] * vector_right - numerical_fluxes[left_face_index] * vector_left)))
    if source_quadrature_function is not None:
        transformed_solution[i, :] += (1.0 / mesh_.elem_metrics[i] * source_quadrature_function(i))
    return transformed_solution

def dg_weak_form_matrix(
    basis_,
    mesh_,
    t,
    boundary_condition,
    numerical_flux,
    quadrature_matrix_function,
    source_quadrature_function=None,
)
num_basis_cpts = basis_.num_basis_cpts
um elems = mesh_.num_elems

# matrix size
n = num_basis_cpts * num elems
L = np.zeros((n, n))
S = np.zeros(n)

phi1 = basis_.evaluate(1.0)
phim1 = basis_.evaluate(-1.0)

C11 = np.matmul(basis_.mass_matrix_inverse, np.outer(phi1, phi1))
C1m1 = np.matmul(basis_.mass_matrix_inverse, np.outer(phi1, phim1))
Cm11 = np.matmul(basis_.mass_matrix_inverse, np.outer(phim1, phi1))
Cm1m1 = np.matmul(basis_.mass_matrix_inverse, np.outer(phim1, phim1))

# iterate over all the rows of the matrix
for i in range(num elems):
    left_elem_index = mesh_.get_left_elem_index(i)
    right_elem_index = mesh_.get_right_elem_index(i)
    left_face_index = mesh_.elems_to_faces[i, 0]
    right_face_index = mesh_.elems_to_faces[i, 1]

    m_i = mesh_.elem_metrics[i]
    x_left = mesh_.get_face_position(left_face_index)
    tuple_ = numerical_flux.linear_constants(x_left, t)
    c_l_imh = tuple_[0]
    c_r_imh = tuple_[1]

    x_right = mesh_.get_face_position(right_face_index)
    tuple_ = numerical_flux.linear_constants(x_right, t)
    c_l_iph = tuple_[0]
    c_r_iph = tuple_[1]

    indices_i = solution.vector_indices(i, num_basis_cpts)

    # 1/m_i (B_i - C11_i + Cm1m1) Q_i
    L[indices_i, indices_i] = (1.0 / m_i) * (quadrate_matrix_function(i) - c_l_iph * C11 + c_r_imh * Cm1m1)

    # S = S_i
    if source_quadrature_function is not None:
        S[indices_i] = (1.0 / m_i) * source_quadrature_function(i)

    # check boundary
    if left_elem_index != -1:
        indices_l = solution.vector_indices(left_elem_index, num_basis_cpts)
        # 1/m_i Cm11_i Q_{i-1}
        L[indices_l, indices_l] = (1.0 / m_i) * c_l_imh * Cm1

    if right_elem_index != -1:
        indices_r = solution.vector_indices(right_elem_index, num_basis_cpts)
The local discontinuous Galerkin method in operator and matrix form in shown below.

```python
# Do boundary conditions
for i in mesh.boundary_faces:
    tuple_ = boundary_condition.evaluate_boundary_matrix(
        mesh_, basis_, i, numerical_flux, t, L, S
    )
    L = tuple_[0]
    S = tuple_[1]

return (L, S)
```

```python
def operator(
    dg_solution,
    t,
    diffusion_function=None,
    source_function=None,
    q_boundary_condition=None,
    r_boundary_condition=None,
    s_boundary_condition=None,
    u_boundary_condition=None,
    q_numerical_flux=None,
    r_numerical_flux=None,
    s_numerical_flux=None,
    u_numerical_flux=None,
    quadrature_matrix_function=None,
):
    (diffusion_function,
     source_function,
     q_boundary_condition,
     r_boundary_condition,
     s_boundary_condition,
     u_boundary_condition,
     q_numerical_flux,
     r_numerical_flux,
     s_numerical_flux,
     u_numerical_flux,
     quadrature_matrix_function,
     ) = get_defaults(
        dg_solution,
        t,
        diffusion_function,
        source_function,
        q_boundary_condition,
        r_boundary_condition,
        s_boundary_condition,
        u_boundary_condition,
        q_numerical_flux,
        r_numerical_flux,
        s_numerical_flux,
        u_numerical_flux,
        quadrature_matrix_function,
    )

```
# Frequently used constants

\[
M^{-1} S \text{quad} = -1.0 \times \text{basis}\_\text{mass}^{-1}\text{stiffness}\_\text{trans} \\
\]

\[
\Phi (1.0) \text{quad} = \text{np multiply}(\text{basis}\_\text{mass}^{-1}\text{stiffness}\_\text{trans}, \text{basis}) \\
\]

\[
\Phi (-1.0) \text{quad} = \text{np multiply}(\text{basis}\_\text{mass}^{-1}\text{stiffness}\_\text{trans}, \text{basis}) \\
\]

\[
\text{quadrature function} = \text{dg-utils get quadrature matrix}(Q, \text{quad matrix}) \\
\]

\[
\text{fluxes}(Q, t, \text{boundary condition}, \text{numerical flux}) \\
\]

\[
\text{weak form}(Q, \text{fluxes}, \text{quad function}, \text{vector left}, \text{vector right}) \\
\]

\[
\text{source quadrature function} = \text{dg-utils get source quadrature}(U, \text{quad matrix}) \\
\]

\[
\text{fluxes}(U, t, \text{boundary condition}, \text{numerical flux}) \\
\]

\[
\text{weak form}(U, \text{fluxes}, \text{quad function}, \text{vector left}, \text{vector right}) \\
\]

\[
\text{source quadrature function} = \text{dg-utils get source quadrature}(U, \text{quad matrix}) \\
\]

\[
\text{fluxes}(U, t, \text{boundary condition}, \text{numerical flux}) \\
\]

\[
\text{weak form}(U, \text{fluxes}, \text{quad function}, \text{vector left}, \text{vector right}) \\
\]

\[
\text{source quadrature function} = \text{dg-utils get source quadrature}(U, \text{quad matrix}) \\
\]

\[
\text{fluxes}(U, t, \text{boundary condition}, \text{numerical flux}) \\
\]

\[
\text{weak form}(U, \text{fluxes}, \text{quad function}, \text{vector left}, \text{vector right}) \\
\]
source_quadrature_function,
)

return L

def matrix(
    dg_solution,
    t,
    diffusion_function=None,
    source_function=None,
    q_boundary_condition=None,
    r_boundary_condition=None,
    s_boundary_condition=None,
    u_boundary_condition=None,
    q_numerical_flux=None,
    r_numerical_flux=None,
    s_numerical_flux=None,
    u_numerical_flux=None,
    quadrature_matrix_function=None,
):

    (diffusion_function,
    source_function,
    q_boundary_condition,
    r_boundary_condition,
    s_boundary_condition,
    u_boundary_condition,
    q_numerical_flux,
    r_numerical_flux,
    s_numerical_flux,
    u_numerical_flux,
    quadrature_matrix_function,
) = get_defaults(
    dg_solution,
    t,
    diffusion_function,
    source_function,
    q_boundary_condition,
    r_boundary_condition,
    s_boundary_condition,
    u_boundary_condition,
    q_numerical_flux,
    r_numerical_flux,
    s_numerical_flux,
    u_numerical_flux,
    quadrature_matrix_function,
)

    basis_ = dg_solution.basis
    mesh_ = dg_solution.mesh

    # quadrature_matrix_function, \quad \mathbf{B}_i = M'(x_i) \mathbf{a}(x_i) \Phi_{x_i} \Phi^T \{x_i
}
# for these problems $a(x_i) = -1.0$ for $r$, $s$, and $u$ equations

```
quadrature_matrix = -1.0 * basis_.mass_inverse_stiffness_transpose
cnst_quadrature_matrix_function = dg_utils.
    get_quadrature_matrix_function_matrix(
        quadrature_matrix
    )

# r - q.x = 0
# $R = A_r Q + V_r$
tuple_ = dg_utils.dg_weak_form_matrix(
    basis_,
    mesh_,
    t,
    q_boundary_condition,
    q_numerical_flux,
    const_quadrature_matrix_function,
)
    r_matrix = tuple_[0]
r_vector = tuple_[1]

# s - r.x = 0
# $S = A_s R + V_s$
tuple_ = dg_utils.dg_weak_form_matrix(
    basis_,
    mesh_,
    t,
    r_boundary_condition,
    r_numerical_flux,
    const_quadrature_matrix_function,
)
s_matrix = tuple_[0]
s_vector = tuple_[1]

# u - s.x = 0
# $U = A_u S + V_u$
tuple_ = dg_utils.dg_weak_form_matrix(
    basis_,
    mesh_,
    t,
    s_boundary_condition,
    s_numerical_flux,
    const_quadrature_matrix_function,
)
    u_matrix = tuple_[0]
u_vector = tuple_[1]
```

# source_quadrature_function

```python
if isinstance(source_function, flux_functions.Zero):
    source_quadrature_function = None
else:
    source_quadrature_function = dg_utils.get_source_quadrature_function(
        source_function, basis_, mesh_, t
    )
```

# $l + (q^3 u).x = 0
# L = A \_l U + V \_l 

tuple\_ = dg\_utils\_. dg\_weak\_form\_matrix(
    basis\_,
    mesh\_,
    t,
    u\_boundary\_condition,
    u\_numerical\_flux,
    quadrature\_matrix\_function,
    source\_quadrature\_function
)

l\_matrix = tuple\_[0]
l\_vector = tuple\_[1]

# R = A \_r Q + V \_r 
# S = A \_s R + V \_s = A \_s ( A \_r Q + V \_r ) + V \_s = A \_s A \_r Q + A \_s V \_r + V \_s 
# U = A \_u S + V \_u = A \_u ( A \_s A \_r Q + A \_s V \_r + V \_s ) + V \_u 
# = A \_u A \_s A \_r Q + A \_u ( A \_s V \_r + V \_s ) + V \_u 
# L = A \_l U + V \_l = A \_l ( A \_u A \_s A \_r Q + A \_u ( A \_s V \_r + V \_s ) + V \_u ) + V \_l 

matrix = np\_. matmul(l\_matrix , np\_. matmul(u\_matrix , np\_. matmul(s\_matrix , r\_matrix )))

vector = (
    np\_. matmul(
        l\_matrix ,
        np\_. matmul(u\_matrix , np\_. matmul(s\_matrix , r\_vector) + s\_vector) + u\_vector
    ),
    + l\_vector
)

return (matrix , vector)