Computation of Steady and Unsteady Laminar Flames: Theory

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Abstract

In this paper we describe the numerical analysis underlying our efforts to develop an accurate and reliable code for simulating flame propagation using complex physical and chemical models. We discuss our spatial and temporal discretization schemes, which in our current implementations range in order from two to six. In space we use staggered meshes to define discrete divergence and gradient operators, allowing us to approximate complex diffusion operators while maintaining ellipticity. Our temporal discretization is based on the use of preconditioning to produce a highly efficient linearly implicit method with good stability properties. High order for time accurate simulations is obtained through the use of extrapolation or deferred correction procedures. We also discuss our techniques for computing stationary flames. The primary issue here is the automatic generation of initial approximations for the application of Newton’s method. We use a novel time-stepping procedure, which allows the dynamic updating of the flame speed and forces the flame front towards a specified location. Numerical experiments are presented, primarily for the stationary flame problem. These illustrate the reliability of our techniques, and the dependence of the results on various code parameters.

1 Introduction

Comprehensive mathematical models of reacting gases have been known for some time. (See, e.g., [18].) Nonetheless, due to the complexity of the models, it is necessary to use numerical methods to extract detailed predictions and evaluate their ability to faithfully reproduce experimental results. In this note we review our ongoing efforts to design and implement reliable numerical methods for the solution of these complex models. Of course, we are not the first researchers to attempt to solve these equations. Of particular note is the work of groups at the Ballistic Research Laboratory [5], Sandia National Laboratory [11] and the Naval Research Laboratory [10]. Nonetheless, some of our approaches differ from those pursued by other groups and seem worthy of additional investigation. In the design process, we have been motivated by potential efficiency in multidimensional simulations. In this work, however, we will restrict ourselves to examples in one space dimension.

2 Models

Our starting point is the equations for conservation of mass, momentum, energy and species:

\[ \frac{\partial p}{\partial t} + \nabla \cdot (pu) = 0, \]  
\[ \frac{\partial u}{\partial t} + (u \cdot \nabla)u + \frac{1}{\rho} \nabla \cdot \rho \nabla \phi = \frac{1}{\rho} \nabla \cdot J, \]  
\[ \frac{\partial e}{\partial t} + (u \cdot \nabla)e = -\frac{1}{\rho} \nabla \cdot q - \frac{P}{\rho} \nabla \cdot u, \]  
\[ \frac{\partial Y_i}{\partial t} + (u \cdot \nabla)Y_i = -\frac{1}{\rho} \nabla \cdot (\rho Y_i V_i) + r_i. \]

Here, \( \rho \) is the density of the gas mixture, \( u \) is the velocity vector, \( e \) is the internal energy and \( Y_i, i = 1, \ldots, n_s \), is the mass fraction of the \( i \)th species.

We consider a zero Mach number model, akin to the incompressible equations for fluid flow. Precisely, our gas law equation of state takes the form:

\[ \rho RT \sum_{i=1}^{n_s} \frac{Y_i}{M_i} = P(t), \quad e = \sum_{i=1}^{n_s} \frac{e_i(T)Y_i}{M_i}, \]
That is, we assume that the pressure, to leading order, is spatially constant. The quantity \( \phi \) is the spatially varying pressure perturbation. It is only retained in the momentum equation. In a closed domain, an additional equation is needed to specify the temporal evolution of \( P \). In an open domain, however, we may assume that \( P \) is constant. Detailed mathematical analyses of these equations and their derivation are given in [13, 8].

There are both advantages and disadvantages to using the zero Mach number system. The primary advantage is in the design of time stepping strategies, as large imaginary eigenvalues of the spatial operators are suppressed. Disadvantages are its restricted applicability to problems where the Mach number remains small and the necessity to solve additional equations to simulate sound production.

To complete the problem specification, we must describe the computation of the reaction rates, \( r_i \), the diffusion velocities, \( V_i \), the heat flux, \( q \), the enthalpies, \( h_i \), and the viscosity coefficients. Our programs are designed to accommodate any reasonable specifications of these quantities without requiring basic changes in the algorithm. In this work the species production rates were obtained by solving the multicomponent diffusion equation, which is a matrix equation—see equation E18 in [18].

\[ e_i(T) = h_i(T) - RT. \] (2.0.6)

Supplementing (3.0.7) and (3.0.9) by (3.0.8), the burnt state is finally determined. In practice, the chemical equilibrium state is not computed using equations (3.0.7) and (3.0.9). It is instead obtained by minimizing the Gibbs function—for details see [9, 16].

### 3 Plane Flames

A fundamental problem of computational combustion is to determine flame speeds and species profiles, or to establish the nonexistence of the flame, given the mixture composition, temperature and pressure. Mathematically, the plane flame is a traveling wave solution of the governing equations. That is, if \( x \) is the coordinate normal to the flame front and \( v \) is the flame speed we seek a solution of (2.0.1)-(2.0.4) which is a function of \( z = x + vt \). Equivalently, we may make a Galilean transformation and seek a steady solution depending on \( x \) alone with a nonvanishing \( x \)-velocity, \( u = v \), at \( x = -\infty \).

Generally, boundary conditions for traveling wave problems involve specifying limiting rest states at \( \pm \infty \). This description does not strictly apply to the plane flame problem, as there is no rest state associated with the unburnt gas. (This is sometimes called the cold boundary difficulty.) Here we simply impose an unburnt state, that is fix the unburnt fuel mixture, \( Y_{i,0} \), temperature, \( T_0 \), and pressure, \( P_0 \). This raises doubts concerning the strict existence of the unbounded domain solution. However, so long as the unburnt state is an approximate equilibrium (that is the reaction rates are sufficiently small), we expect that our truncated problems do possess solutions which correctly model the physics.

True equilibria, which will define the burnt state, are defined by the \( n_s \) equations:

\[ r_i(Y_{i,1},\ldots,Y_{n_s,1},T_b) = 0. \] (3.0.7)

As the number of unknowns exceeds the number of equations by one, it is reasonable to expect that a one-parameter family of equilibria exist. To uniquely determine the burnt state, we note that under the zero Mach number assumption, an additional conservation law for the enthalpy can be derived:

\[ \frac{\partial}{\partial x} \left( q + \sum_{i=1}^{n_s} \rho(u + v_i)V_ih_i \right) = 0. \] (3.0.8)

(Here \( h_i \) is the enthalpy of species \( i \).) Integrating this equation from the unburnt to the burnt state, imposing the vanishing of the gradients and, hence, of \( q \) and \( V_i \) in these states, and taking into account mass conservation, we arrive at the additional relation:

\[ \sum_{i=1}^{n_s} Y_{i,0}h_{i,0} = \sum_{i=1}^{n_s} Y_{i,1}h_{i,1}. \] (3.0.9)

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An additional property of traveling wave solutions is their translation invariance. That is, given any solution a new solution can be obtained by adding any constant to the independent variable, effectively translating the profile. Mathematically, this implies that the space derivative of the solution is a nullfunction of the linearized equations. To fix our profile in space we impose an additional phase condition. Many such conditions are possible. (For an interesting discussion of the discretization theory of traveling wave problems see Beyn [4].) What is essential is that the additional condition be nonvanishing when applied to the derivative of the solution. We choose a temperature normalization:

\[ T(x_h) = T_h, \quad T_0 < T_h < T_b. \] (3.0.10)
For this to be effective we must have:

\[
\frac{dT}{dx}(x_h) > 0. \tag{3.0.11}
\]

So long as \( T = T_h \) occurs within the flame front, (3.0.11) is expected to be valid, and has been in all our experiments. Later we will discuss a new method for dealing with the phase condition in the context of a time stepping procedure for computing the flame.

4 Spatial Discretization and Adaptivity

We approximate spatial derivatives using second to sixth order central differences, switching to lopsided schemes where necessary near the boundaries. We expect that higher order methods will prove to be more efficient, particularly when the physical models are complex. In these cases, the overhead associated with the use of higher order differences is small compared with the cost of evaluating various physical quantities at each point. Hence, any reduction in the number of mesh points results in a reduction of computing time.

Two distinct spatial difference formulas are employed. The first, which is denoted \( D_g \) and used to approximate first order terms in the equation, takes a function defined on a uniform grid and produces approximate derivatives on that grid. The second operator, \( D_h \), approximates derivatives midway between gridpoints. These operators are composed to approximate second order derivatives on the grid. For example, we approximate the species conservation equations (2.0.4) by:

\[
uD_gY_i = -\frac{1}{\rho}D_h((I_h\rho)(I_hY_i)V_i) + r_i, \tag{4.0.12}
\]

where \( I_h \) is an interpolation operator of appropriate order producing function values midway between gridpoints from values on the grid. The diffusion velocities are computed, again midway between gridpoints, by:

\[V_i = V_i(D_hY_j, D_hT, I_hY_j, I_hT). \tag{4.0.13}\]

The reason for staggering the grids in this way is to guarantee the ellipticity of the resulting discrete diffusion operators. The only grid functions annihilated by \( D_g \) are constant, whereas \( D_h \) also annihilates the function \((-1)^j\) where \( j \) is a grid index. Therefore, the Fourier transforms of second order operators formed by composing \( D_h \) with itself are always large for large wavenumbers while those obtained by composing \( D_g \) with itself decrease to zero as the wavenumber approaches \( \pi/\Delta x \), that is as we approach the largest wavenumber representable on the grid.

For the one-dimensional problems considered here, the essential length scales are the computational domain length, \( L \), which we cannot make too small without using far more sophisticated boundary treatments, and the width of the flame zone. In practice the ratio of these is sufficiently small to merit an adaptive mesh strategy. Currently, we use a parametrized family of coordinate maps. This strategy has been successfully employed and analyzed by Bayliss and coworkers \cite{1, 2} in the context of spectral simulations of simpler combustion models. The maps we use are slightly different. Precisely, we write:

\[x = x_l + \frac{1}{2} \gamma y \left( \epsilon + \frac{(1 - \epsilon)(y/\beta)^p}{1 + (y/\beta)^p} \right), \tag{4.0.14}\]

and employ a uniform grid in \( y \). For \( \epsilon < 1 \) there is a clustering of gridpoints near \( x = x_l \). Using the temperature as our monitor function, we choose \( x_l \) to coincide with the location of the maximum derivative of \( T \) and choose \( \epsilon \) according to the ratio:

\[\epsilon = \frac{T_h - T_0}{L(T_{max})}. \tag{4.0.15}\]

Throughout our simulations we have chosen \( p = 8 \). The parameters \( \gamma \) and \( \beta \) are chosen to scale the length of the \( y \)-interval and to guarantee that sufficiently many points are in the refinement layer. Our exact formulas are:

\[\beta = (1 + 3\epsilon)^{-1}, \quad \gamma = \frac{1 + 3\epsilon}{4\epsilon} \cdot L. \tag{4.0.16}\]

A defect of the very simple mapping described above is its inability to handle multiple fronts. An alternative method we have used which is capable of handling multiple fronts is given by the mapping:

\[y = A \int_0^x \left(1 + \xi |T_x| + (1 - \xi)|T_{xx}|^{1/2}\right)^{1/2} dz. \tag{4.0.17}\]

where \( A \) is a normalization parameter and \( 0 \leq \xi \leq 1 \).

In all cases we use static rezoning, that is update the mesh after some fixed number of time steps and interpolate the solution onto the new mesh. Each of these methods would require extensive modification for use in multiple dimensions. Therefore, a more flexible approach such as the AMR techniques of Berger and coworkers (e.g. \cite{3}) will be considered.

5 Temporal Discretization

It is well known that the equations of reacting flows may be stiff when realistic reaction mechanisms are
Therefore, if an explicit temporal discretization were employed, the time step could be severely restricted by stability requirements. Fully implicit discretizations, on the other hand, require the solution of a discrete approximation to a nonlinear elliptic system at each time step. Solving such a problem is potentially quite expensive in multiple dimensions. Here we take an intermediate approach, based on the concept of preconditioning.

Consider, for simplicity, a system of ordinary differential equations:

\[ \frac{du}{dt} = F(u). \quad (5.0.18) \]

Introduce a preconditioning matrix, \( G(u) \). A first order approximation to (5.0.18) is given by:

\[ (I - \Delta t G(u(t)))w = \Delta t F(u(t)), \quad (5.0.19) \]
\[ u(t + \Delta t) = u(t) + w. \quad (5.0.20) \]

For reasons of efficiency, we want to choose \( G \) so that (5.0.19) may be easily solved. To study the stability of the approximation, we consider the special case \( F = Ju \). Then we have:

\[ u(t + \Delta t) = (I - \Delta t G)^{-1}(I + \Delta t(J - G))u(t) \quad (5.0.21) \]

so that the method is stable if and only if for some subordinate matrix norm:

\[ \|(I - \Delta t G)^{-1}(I + \Delta t(J - G))\| \leq 1. \quad (5.0.22) \]

As a simple example, suppose \( J \) is a discrete approximation to a variable coefficient multidimensional diffusion operator:

\[ J = \sum_j D_{h,j}(\sigma_j(x) D_{h,j}), \quad (5.0.23) \]

where \( D_h \) are as discussed above. Then it is possible to take:

\[ (I - \Delta t G) = \prod_j (I - \Delta t \bar{\sigma}_j D_{+,j} D_{-,j}) \quad (5.0.24) \]

for suitably chosen \( \bar{\sigma}_j \). Then we only need to solve small tridiagonal systems, no matter what stencil width our actual difference approximations employ.

The time stepping scheme, as outlined above, is only first order. This is sufficient for the computation of stationary flames, where time-stepping is simply used to generate an initial approximation for Newton's method. For time accurate simulations, on the other hand, it is more efficient to use a higher order method. There are two distinct approaches we have considered for automatically producing higher order time advancement procedures based on the preconditioning strategy: extrapolation, as in the ode solver SEULIM (e.g. [6]), and iterated deferred correction, as discussed in [7]. As the second approach is less familiar, and is the one employed in the example given below, we describe it briefly.

Suppose \( u(t_n) \) is our approximate solution at \( t = t_n \) and that we wish to compute \( u(t_{n+1}) \). Recall the formula obtained by integrating (5.0.18) from \( t_n \) to an arbitrary time, \( t \):

\[ u(t) = u(t_n) + \int_{t_n}^t F(u(s))ds. \quad (5.0.25) \]

Choose nodes (typically of Gauss, Radau or Lobatto type),

\[ t_n \leq s_1 < s_2 < \ldots < s_m \leq t_{n+1}. \quad (5.0.26) \]

Let the \( p \)th approximate solution on the nodes be given by:

\[ v^p_j \approx u(s_j), \quad (5.0.27) \]

where the initial approximation is computed by (5.0.19),(5.0.20) with \( \Delta t \) replaced by \( s_{j+1} - s_j \). To compute corrections, define the interpolant, \( Q^p(s) \), to \( (s_j, F(v^p_j)) \) and the residual:

\[ \epsilon^p_j = u(t_n) + \int_{t_n}^{s_j} Q^p(s)ds - v^p_j. \quad (5.0.28) \]

Then solve:

\[ (I - k_j G(v^p_j))(\delta^p_{j+1} - \delta^p_j) = (5.0.29) \]
\[ k_j (F(v^p_{j+1}) - F(v^p_j)) + \epsilon^p_{j+1} - \epsilon^p_j, \quad (5.0.30) \]

\[ k_j = s_{j+1} - s_j, \quad (5.0.31) \]

The overall method order increases by one at each iteration, up to a maximum determined by the integration scheme. Assuming this order is not less than \( l \), an \( l \)th order approximation to \( u(t_{n+1}) \) is given by:

\[ u(t_{n+1}) = u(t_n) + \int_{t_n}^{t_{n+1}} Q^{l-1}(s)ds. \quad (5.0.32) \]

Note that our semidiscretized system is in fact a differential-algebraic rather than a differential equation. Formally, the index of this system is three. In particular, no equation for the time derivative of the pressure variation, \( \phi \), is given. To obtain such an equation, one must differentiate the constraint, namely the equation of state (2.0.5), three times. (The first differentiation produces \( \rho_t \); substituting for \( \rho_t \) using (2.0.1), the second differentiation
produces \( u_t \); finally, substituting for \( u_t \) using (2.0.2) yields an equation for \( \phi \) which can be differentiated and solved for \( \phi_t \). In one space dimension, however, (2.0.2) and, hence, \( \phi \) may be eliminated. Then we have no equation for \( u_t \), and the index, following the same reasoning as above, is reduced to two.

In the current code, we only apply our time-stepping procedure to the spatially discretized approximations to (2.0.3) and (2.0.4). The density is computed using (2.0.5) and the velocity is updated using (2.0.1). For stationary flames this reduces to:

\[
  u = \frac{\rho_0}{\rho} \cdot v, \tag{5.0.33}
\]

where \( v \) is the velocity at \( x = -\infty \). For time accurate approximations we found it difficult to use (2.0.1) directly. Instead, we replace \( \rho_t \) by first expressing it in terms of \( e_t \) and \( Y_{1,t} \) and then using (2.0.3) and (2.0.4) to eliminate all time derivatives. This results in a somewhat complicated expression, which is nevertheless easily integrated for \( u \).

In our code, \( G \) is chosen by replacing the heat conduction and species diffusion terms in equations (2.0.3) and (2.0.4) by simplified physical approximations and by using second order differences. It also includes the exact Jacobian of the reaction rates.

Finally, we note that the time steps are automatically adjusted, within preset upper and lower bounds, according to the relative change in the solution.

### 6 Stationary Flames

As discussed in Section 3, the plane flame problem is to compute stationary solutions to equations (2.0.1)-(2.0.4) along with the normalization condition (3.0.10) and the boundary conditions:

\[
  (e, Y_i) \to (e_0, Y_{i,0}), \quad x \to -\infty, \tag{6.0.34}
\]
\[
  (e, Y_i) \to (e_b, Y_{i,b}), \quad x \to \infty, \tag{6.0.35}
\]
\[
  u \to v, \quad x \to -\infty. \tag{6.0.36}
\]

Recall that the state \((e_0, Y_{i,0})\) is specified, the state \((e_b, Y_{i,b})\) is computed given \((e_0, Y_{i,0})\) by minimizing the Gibbs function, and the flame speed, \( v \), is to be determined as part of the calculation.

A direct approach to the computation is to truncate the infinite domain by a finite domain, replacing (6.0.34)-(6.0.36) by approximate boundary conditions at the artificial boundaries, then to approximate the spatial derivatives by differences on a (generally nonuniform) mesh, and finally to use some variant of Newton’s method to solve the resulting algebraic system. The problem here is in the last step, as the convergence of Newton’s method is only guaranteed for a sufficiently accurate initial guess, which seems difficult to automatically generate for this problem. A more reliable alternative is to use a time-stepping procedure in place of Newton iterations. In this approach one makes use of the apparently large dynamical basin of attraction of the plane flame solution. Disadvantages associated with standard time-stepping procedures include slow convergence to steady state, an inability to fix the flame front, and difficulties in determining a precise speed.

In this work we use a hybrid approach. We begin with a novel time-stepping procedure, which allows us to dynamically compute a flame speed and to modify the dynamics to push the front towards its normalized location. Similar ideas have been proposed by Sermange [17], but ours differ in the details. Second, either after the time derivatives become sufficiently small or after a maximum number of time steps are carried out, we switch to a damped Newton iterator to achieve final convergence.

The time-stepping procedure is most easily explained in a general setting. Consider the nonlinear, algebraic eigenvalue problem:

\[
  F(w) - vSw = 0. \tag{6.0.37}
\]

Here \( v \) is the eigenvalue and \( S \) is a square matrix. Discrete approximations to the plane flame problem take this form when the velocity, \( u \), is replaced using (5.0.33) and \( \rho \) is eliminated using (2.0.5). Equation (6.0.37) then represents the discrete approximation to (2.0.3)-(2.0.4) multiplied by \( \rho \). The matrix \( S \) is the discrete derivative operator.

As mentioned above, solutions to the plane flame problem are only unique up to translation. This implies that the equations linearized about a plane flame solution have a nullfunction which is the derivative of the solution itself. Moreover, we determine a unique solution by imposing an additional normalization condition (3.0.10). We represent our discrete approximation to (3.0.10) by

\[
  N(w) = 0, \tag{6.0.38}
\]

where \( N \) is scalar valued. Corresponding to the translation invariance, we assume that if \( w^* \) is the solution of (6.0.37)-(6.0.38), then the matrix:

\[
  A \equiv DF(w^*) - vS, \tag{6.0.39}
\]

has 0 as a simple eigenvalue with normalized eigenvector \( q_0 = S w^* / |S w^*| \). (This assumption is generally false after domain truncation. However, dropping it complicates the analysis beyond the scope of...
this note.) We further assume that $w^*$ is dynamically stable, that is, that all other eigenvalues, $\lambda_j$, of $A$, have negative real part. Moreover, we impose the nondegeneracy condition:

$$DN(w^*)Sw^* > 0. \quad (6.0.40)$$

Note that if the left-hand side of (6.0.40) were zero, the normalization would fail to guarantee a unique solution. In our case, (6.0.40) is equivalent to (3.0.11).

Our time-stepping procedure is to replace (6.0.37) by the dynamical problem:

$$w_t = G(w), \quad (6.0.41)$$

with $G$ defined by:

$$G(w) = F(w) - (\Phi(w) + \mu N(w))Sw, \quad (6.0.42)$$

where $\Phi$ is a scalar function and $\mu > 0$ is a damping parameter. In order for $w^*$ to be a rest point of the new system, it is necessary to guarantee that:

$$\Phi(w^*) = v. \quad (6.0.43)$$

This we accomplish by choosing:

$$\Phi(w) = \frac{(Sw)^T R F(w)}{(Sw)^T R (Sw)}, \quad (6.0.44)$$

for some symmetric semidefinite matrix $R$. In our current code, $R$ is simply the projection onto the internal energy.

The effectiveness of our approach depends on the spectrum of $DG(w^*)$, which must lie in the left half-plane. By direct computation we find:

$$DG(w^*) = \left( I - \frac{(Sw^*)(Sw^*)^T R}{(Sw)^T R (Sw)} \right) A \quad (6.0.45)$$

$$-\mu (Sw^*)DN(w^*).$$

Let $Q$ be an orthogonal matrix whose first column is $q_0$. Then we have:

$$Q^T AQ = \begin{pmatrix} 0 & \gamma^T \\ 0 & A^{(1)} \end{pmatrix}, \quad (6.0.46)$$

and

$$Q^T DG(w^*)Q = \begin{pmatrix} \lambda_0 & \gamma^T \\ 0 & A^{(1)} \end{pmatrix}, \quad (6.0.47)$$

$$\lambda_0 = -\mu DN(w^*)Sw^* < 0. \quad (6.0.48)$$

Therefore, the spectrum of $DG(w^*)$ is given by the eigenvalues of $A^{(1)}$, which by (6.0.46) are also the nonzero eigenvalues of $A$, and $\lambda_0$. Hence, the local dynamic stability of $w^*$ is assured.

Finally, we consider the boundary conditions imposed at the artificial boundaries introduced to truncate the infinite domain. At the cold boundary we simply impose the Dirichlet conditions:

$$(e, Y_i) = (e_0, Y_{i,0}). \quad (6.0.49)$$

Our justification for this is the rapid exponential decay to this state which we observe in the computations. At the hot boundary, where the observed decay rates are much slower, we impose the conditions analyzed by Lohéac [12] for advection-diffusion equations:

$$e_{xx} = Y_{i,xx} = 0. \quad (6.0.50)$$

Although our experiments indicate that these conditions do provide sufficient accuracy, more sophisticated conditions could be used and might lead to improvements. See [4] for a detailed discussion.

7 Results

In this section we present the results of some numerical experiments with our code. The vast majority of these involve computations of a stationary, stoichiometric hydrogen-air flame at a temperature of 300 degrees Kelvin and atmospheric pressure. Our mechanism includes 13 species.

This problem is rather standard, and was included in a series of benchmark problems in laminar flame computations sixteen years ago [15]. We note that our mechanism does differ from the one used there. Our complete set of results is summarized in Table 1. Our computed flame speeds vary between 235.77 cms and 239.08 cms. If the somewhat less reliable sixth order results are ignored, this variance is reduced from 236.86 to 237.92 - i.e. about one-half of one percent. It is interesting to note that these results show far better agreement with the experimental results of Milton and Keck [14] than do any presented in [15], though this no doubt says more about the accuracy of our models than it does about the merits of our code. Plots of the temperature, velocity, and various mole fractions for a typical case (fourth order, 149 points, domain width $L = 1$) are presented in Figures 1 and 2.

In all cases we tried except one, the code successfully ran to completion. (The one failure was with the sixth order differencing and our coarsest mesh, $n = 49$.) The maximum number of time steps was set to 7000 and the code automatically switched to damped Newton iterations whenever:

$$\frac{||dw/dt||}{||w||} \leq 5 \text{sec}^{-1}. \quad (7.0.51)$$
(Here $w$ is the complete discrete solution.) The convergence criterion for Newton’s method was a relative change less than $10^{-5}$. This criterion was not always achieved, in which case the final estimated relative error is listed in the tolerance column. Such failures to fully converge are most likely due to a highly ill-conditioned Jacobian. Note that Newton’s method fails to converge at all if the initial conditions for the time-stepping are used as its initial guess.

**Mesh Refinement**: Results are shown for varying numbers of mesh points and various orders. Fixing all other parameters, the results clearly converge under refinement.

**Varying Order**: Both the second and fourth order codes show consistent convergence behavior. However, fixing all other parameters, the flame speeds computed using the second order code are slightly smaller. One would, of course, hope for even better agreement here. Again this may be attributable to conditioning problems. The results for the sixth order code are a little worse, at least as judged by its convergence under mesh refinement. It is possible that the sixth order method is more susceptible to stability problems due to the large one-sided stencils it uses near the boundaries. It may be necessary to use some additional mesh clustering at the boundary to improve its performance.

**Varying Domain Length, $L$**: We have computed solutions on both a small domain, $L = 1$cm, and a large domain, $L = 10$cm. Again, for the second and fourth order codes there is excellent agreement (about $1/20$ of one percent) between the results on different domains. This indicates that our artificial boundary conditions are adequate for the domain lengths considered.

**Varying $x_h$**: Fixing our domain to be $x \in [0, 1]$, we used the fourth order method with 149 points to compute solutions with $x_h$ varying from $1/4$ (our standard value) to $3/4$. The flame speed varied between 236.86 and 237.90. Results for $x_h \in [1/4, 1/2]$ showed far less variance. We expected this, as the decay rate of the solution to the unburnt state is far greater than its decay to the burnt state.

**Varying $\mu$**: We also varied the damping parameter, $\mu$, with the fourth order method, $x_h = 1/4$, $L = 1$ and $n = 149$. There were significant changes in the number of time steps required, with the best results obtained with $\mu = 1000$. Surprisingly, the flame speeds also varied somewhat. Note that varying $\mu$ only changes the initial guess from the point of view of the Newton iterator. The discrepancies in the results are another indication of ill-conditioning.

**Time Accurate Simulations**: We have, at this time, far less experience running our code for time-dependent problems. Therefore, we include only one preliminary result. It involves using the stationary flame profile as initial data and setting the inlet velocity to zero. The flame then propagates to the right. From the graphs it is clear that the profile and speed are correctly maintained. The data shown were obtained using the fourth order method in space and time with 99 mesh points.

### 8 Conclusions

Our fundamental conclusion, based not only on the results shown here, but on numerous others to be published at a later date, is that our code is both accurate and robust for stationary plane flame calculations. We must, of course, carry out many more experiments to assess its behavior for time-dependent and multidimensional problems. The results here do point to some difficulties in achieving high accuracy, which are most likely due to poor conditioning. This issue should be investigated further.

### References


Appendix

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Table 1: Summary of Results
Figure 1. Computed Solution, Order= 4, n = 149, L = 1.
Figure 2. Computed Solution, Order=4, n = 149, L = 1.
at time $t=0, 2d-4, 4d-4, 6d-4, 8d-4, 1d-3$

Figure 3. Unsteady Sol. $n = 99, L = 1$. 
at time $t=0$, $2d-4$, $4d-4$, $6d-4$, $8d-4$, $1d-3$

**Figure 4.** Unsteady Sol. $n = 99$, $L = 1$. 
Mole fraction of species H2

at time t=0, 2d-4, 4d-4, 6d-4, 8d-4, 1d-3

Figure 5. Unsteady Sol. n = 99, L = 1.
Mole fraction of species O2

at time t=0, 2d⁻⁴, 4d⁻⁴, 6d⁻⁴, 8d⁻⁴, 1d⁻³

Figure 6. Unsteady Sol. n = 99, L = 1.
Figure 7. Unsteady Sol. $n = 99, L = 1$. 

Mole fraction of species H2O
Mole fraction of species $\text{H}_2\text{O}_2$

at time $t=0, 2d^{-4}, 4d^{-4}, 6d^{-4}, 8d^{-4}, 1d^{-3}$

Figure 8. Unsteady Sol. $n = 99, L = 1.$
### Title and Subtitle

Computation of Steady and Unsteady Laminar Flames: Theory

### Authors

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### SUPPLEMENTARY NOTES


### ABSTRACT (Maximum 200 words)

In this paper we describe the numerical analysis underlying our efforts to develop an accurate and reliable code for simulating flame propagation using complex physical and chemical models. We discuss our spatial and temporal discretization schemes, which in our current implementations range in order from two to six. In space we use staggered meshes to define discrete divergence and gradient operators, allowing us to approximate complex diffusion operators while maintaining ellipticity. Our temporal discretization is based on the use of preconditioning to produce a highly efficient linearly implicit method with good stability properties. High order for time accurate simulations is obtained through the use of extrapolation or deferred correction procedures. We also discuss our techniques for computing stationary flames. The primary issue here is the automatic generation of initial approximations for the application of Newton’s method. We use a novel time-stepping procedure, which allows the dynamic updating of the flame speed and forces the flame front towards a specified location. Numerical experiments are presented, primarily for the stationary flame problem. These illustrate the reliability of our techniques, and the dependence of the results on various code parameters.