Development and Application of a Computer Model for Large-Scale Flame Acceleration Experiments

K. D. Marx
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DEVELOPMENT AND APPLICATION OF
A COMPUTER MODEL FOR LARGE-SCALE
FLAME ACCELERATION EXPERIMENTS

K. D. Marx

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ABSTRACT

A new computational model for large-scale premixed flames is developed and applied to the simulation of flame acceleration experiments. The primary objective is to circumvent the necessity for resolving turbulent flame fronts; this is imperative because of the relatively coarse computational grids which must be used in engineering calculations. The essence of the model is to artificially thicken the flame by increasing the appropriate diffusivities and decreasing the combustion rate, but to do this in such a way that the burn velocity varies with pressure, temperature, and turbulence intensity according to prespecified phenomenological characteristics. The model is particularly aimed at implementation in computer codes which simulate compressible flows. To this end, it is applied to the two-dimensional simulation of hydrogen-air flame acceleration experiments in which the flame speeds and gas flow velocities attain or exceed the speed of sound in the gas. It is shown that many of the features of the flame trajectories and pressure histories in the experiments are simulated quite well by the model. Using the comparison of experimental and computational results as a guide, some insight is developed into the processes which occur in such experiments.

* This work was performed at the Combustion Research Facility and supported by the U. S. Nuclear Regulatory Commission.
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Roman:

\( A \)  Coefficient in burn velocity formula
\( A_c \)  Combustion coefficient
\( B \)  Coefficient in burn velocity formula
\( c_p \)  Specific heat at constant pressure
\( c_{p0} \)  Constant in approximation for specific heat
\( c_p' \)  Slope in approximation for specific heat
\( C \)  Progress variable
\( C_1 \)  Coefficient in burn velocity formula
\( C_{2e} \)  Coefficient in approximation for specific heat
\( C_{1e} \)  Slope in approximation for specific heat
\( C_{2e} \)  Progress variable
\( D \)  Diffusion coefficient
\( E \) (App. A only)  Special burn rate function
\( f(T) \)  Special burn rate function
\( f_p(u,k,\epsilon) \) (App. A only)  Special burn rate function
\( h(p, T_u) \)  Special burn rate function
\( h_i \)  Specific Enthalpy of \( i^{th} \) species
\( I \)  Specific energy
\( k \)  Turbulent kinetic energy
\( k_e \)  Effective turbulent kinetic energy
\( k_m \)  Threshold turbulent kinetic energy
\( k_T \)  Thermal conductivity
\( L \)  Turbulent length scale
\( L_e \)  Effective turbulent length scale
\( L_m \)  Threshold turbulent length scale
\( m \)  Pressure exponent in burn velocity formula
\( m_i \)  Molecular weight of \( i^{th} \) species
\( n \)  Temperature exponent in burn velocity formula
\( P \) (App. A only)  Pressure
\( P_r \)  Prandtl number
\( P' \) (App. A only)  \( C_{\mu} f_p \)
\( Q \) (App. A only)  \( \rho k \)
\( Q_i^c \)  Rate of energy release due to chemistry
\( r \)  Thermal expansion ratio
\( R_T \)  Turbulent Reynolds number
\( S_c \)  Schmidt number
\( S_L \)  Laminar burn velocity
\( S_T \)  Turbulent burn velocity
\( S_{TI} \)  Initial turbulent burn velocity
\( S_T^e \) (Table I only)  Expected turbulent burn velocity
\( S_T^i \)  \( dS_T/d\sqrt{k} \)
$t$  Time
$t_t$  Typical transit time
$t_T$  Turbulence time scale
$T$  Temperature
$T_a$  Adiabatic flame temperature
$T_o$  Initial gas temperature
$T_u$  Unburned gas temperature
$t_X$  Thermal diffusion time
$u$  Gas velocity
$u_*$  Wall shear velocity
$u'$  Turbulence intensity
$v_{fo}$  Initial flame speed
$v_T$  Turbulent flame speed
$w$  Channel width
$x$  Spanwise coordinate
$y$  Vertical coordinate
$y$ (App. B only)  Coordinate measuring distance from wall
$Y_i$  Mass fraction of the $i^{th}$ species
$Y_p$ (App. A only)  Mass fraction of product species for completed burn
$z$  Lengthwise coordinate

Greek:

$\beta$  Refers to $\beta$-transformation
$\delta$  Unit tensor
$\delta_c$ (Table I only)  Computational flame thickness
$\delta_c^e$ (Table I only)  Expected computational flame thickness
$\delta_L$  Laminar flame thickness
$\delta_T$  Turbulent flame thickness
$\Delta t$  Time step
$\Delta x$  Grid spacing
$\Delta y$ (App. B only)  Grid spacing
$\epsilon$  Turbulence dissipation rate
$\kappa$  Von Kármán constant
$\mu$  Total viscosity
$\mu_L$  Laminar viscosity
$\mu_T$  Eddy viscosity
$\nu_i$  Stoichiometric coefficient of $i^{th}$ species
$\xi$ (App. A only)  Q/E
$\rho$  Total density
$\rho_i$  Density of $i^{th}$ species
\[ \dot{\rho}_f \] Rate of consumption of fuel
\[ \dot{\rho}_i^e \] Rate of change of density of \( i^{th} \) species due to reactions
\( \sigma \) Stress tensor
\( \sigma_\epsilon \) Coefficient in diffusion of \( \epsilon \)
\( \tau \) Wall shear stress
\( \chi \) Thermal diffusivity

**Subscripts:**

- \( e \): Effective values
- \( f \): Fuel
- \( i \): Species
- \( i \) (App. B only): Grid index
- \( L \): Laminar
- \( p \): Product species
- \( T \): Turbulent
- \( \epsilon \): Refers to turbulent dissipation rate

**Superscripts:**

- \( \text{base} \) (Table I only): Refers to baseline calculation
- \( c \): Chemistry
- \( \epsilon \) (Table I only): Expected values
- \( T \): Transpose of tensor
- \( \prime \) (on \( c_p, S_T \)): Differentiation
- \( \prime \) (on \( \mu, l_T, \mu_T \)): Values of quantities in final form of combustion model

**Abbreviations:**

- CJ: Chapman-Jouguet
- DDT: Deflagration-to-detonation transition
- FLAME: Flame Acceleration Measurements and Experiments facility
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Executive Summary

This report represents a major portion of the work performed at Sandia National Laboratories, Livermore under the Sandia Hydrogen Program for the U.S. Nuclear Regulatory Commission. Computational methods are described which permit techniques previously used for modeling small-scale laboratory experiments to be extended to the simulation of experiments with much larger length scales. The specific configurations considered are the propagation of hydrogen-air burns in the FLAME facility at Sandia National Laboratories, Albuquerque. The largest length scales appearing in these experiments approach in size those which might occur in nuclear reactor accident scenarios.

These length scales exceed the thicknesses of the hydrogen flames by a large factor. Because of this, it was necessary to invent a new procedure to represent the flow variables on a finite-difference grid in which computer limitations require that the grid spacing exceed the physical flame thickness. The new method consists of imposing minimum values of turbulence intensity and turbulent length scale so as to artificially thicken the flame. This is done in such a way that the burn velocity obeys established dependencies on turbulence level, pressure, and unburned gas temperature.

The resulting model is applied to the simulation of one of the experiments performed with obstructions in the channel of the FLAME facility. By using only the initial flame velocity to adjust parameters in the model, a quite successful representation of the experimental flame trajectory and pressure histories is obtained. The calculation does display some inadequacies; it fails to reproduce a sudden acceleration midway in the experiment, and it exhibits a detonation near the end. Nonetheless, it provides a meaningful study of many features of the gas flow and flame propagation.

As one example of this, the computations point out the way in which the acceleration of the flame and its propagation in the choking regime are accompanied by large increases in the amplitude of pressure waves which reflect off the obstacles. This mechanism contributes to the positive feedback which drives the flame acceleration in two ways: (1) it leads to an increased combustion rate via compression and shock heating and (2) it produces increased turbulence through the generation of large flow velocities with high shear, thereby augmenting the thermal mixing. These increasing pressure and turbulence levels are also undoubtedly indicative of a buildup of sensitivity to detonation. This emphasizes the importance of a careful interpretation of experiments with regard to the degree of confinement of the gas, which has been well-established experimentally. The results are consistent with a reduction in flame acceleration in the presence of pressure relief due to venting. (It is noted, however, that there is a potential for increased production of turbulence in the presence of partial venting which remains to be investigated.)

In summary, this work contains two major contributions: (1) the development of the numerical model, and (2) the use of the computer calculations to provide insight into the processes accompanying flame acceleration in experiments in the FLAME facility. It defines the attributes and limitations of current computational capabilities in this research area, and suggests additional steps that could be taken to improve them.
I. INTRODUCTION

In this paper, we describe a computational model for simulating the propagation of premixed turbulent flames which extend over spatial length scales which are large compared to those which characterize the turbulent flame zone. In these situations, the geometric complexity is often such that it is impossible to provide accurate resolution of the flame front on a computational grid which encompasses the entire domain of interest. The specific configurations to which this work has been applied are experiments performed in the FLAME facility\textsuperscript{1−3} at Sandia National Laboratories in Albuquerque, New Mexico (see Figure 1). The FLAME facility consists of a reinforced concrete channel 2.44 m high by 1.83 m wide by 30.5 m long (8 ft by 6 ft by 100 ft). The particular experiments considered here involve the ignition of a hydrogen-air mixture at one (closed) end and the propagation of the flame past periodically spaced obstacles toward the other (open) end (see Figure 2).

Turbulence generated ahead of the flame as the unburned gas flows past the obstacles results in an increase in the combustion rate. This, in turn, causes the flow velocity of the unburned gas to increase, with a concurrent increase in the turbulence level. Hence, there is a positive feedback mechanism which leads to acceleration of the flame as it propagates down the channel.

This mechanism for flame acceleration has been described previously for experiments on a smaller scale.\textsuperscript{4−6} Some work has also been done on large-scale experiments.\textsuperscript{7} The primary purpose of the FLAME facility is to provide relatively large-scale results for hydrogen-air burns. The ultimate goal of the research is to determine scaling laws which will permit assessment of potential hazards to nuclear reactors in case of hydrogen production in reactor accidents (see, e.g., Reference 8). The length scales existing in the FLAME facility are much larger than those typically occurring in the small-scale experiments described in References 4-6, but are somewhat smaller than some which might be encountered in a reactor containment.

The purposes of this paper are to describe our method for simulating flame acceleration in such large-scale configurations, to compare computational results with a limited set of experimental data, and to provide some analysis of the processes occurring in the experiments.

The computer modeling of reacting flows as complex as those existing in the present experiments is extremely difficult. There are, in principle, a large number of chemical reactions to be accounted for. In all but the simplest flow configurations, one must approximate the true chemical kinetics with a reduced model. Furthermore, as noted above, turbulence plays a crucial role in determining the rate of mixing of burned and unburned gas, and therefore has a great influence over the rate of combustion. A complete solution to the Navier-Stokes equations for turbulent flow is manifestly impossible in practical engineering calculations such as these. It is therefore necessary to provide a turbulence model which permits the computation of turbulence intensities and time scales. In this work, we have used the $k-\epsilon$ model.\textsuperscript{9−10} Given such a method for the determination of turbulence parameters, one must couple it to the combustion model in order to have a realistic determination of the rate at which the unburned gas is consumed. There have been various proposals for reasonable ways to do this. Most notable for our purposes are the Magnussen-Hjertager model\textsuperscript{11} and the eddy breakup model.\textsuperscript{12−13}
Figure 1. Sketch of the FLAME facility. (From Reference 3.) The vertical coordinate is assumed ignorable in the two-dimensional computer calculations. (Note: FLAME is an acronym for "FLame Acceleration Measurements and Experiments.")
Figure 2. Schematic of the FLAME facility with obstacles installed. (From Reference 34.) The obstacles used in the experiments described in this work are shown to scale; the blockage ratio is $\frac{1}{3}$. The enclosed area in the top view is the two-dimensional domain of the computer calculations.
The Magnussen-Hjertager model is useful in situations where the flame propagation is
determined by the rate of thermal mixing. It has been previously employed in studies of flame
acceleration in small-scale experiments.\textsuperscript{14} However, when it was used in conjunction with the $k-\epsilon$
model in attempted simulations of flame acceleration experiments, the results were unfavorable.
Strong spatial oscillations of the flow variables and turbulence parameters appeared in the
solution. Although the oscillations were of short wavelength, i.e., alternating on adjacent grid
points, no temporally increasing numerical instability appeared. It appeared that the difficulty was
introduced by trying to resolve phenomena on a length scale incompatible with the grid spacing.
As noted above, this is anticipated when features in the flow configuration are large compared
to the flame thickness. In such cases, in order to achieve a numerically viable description of
the fluid, it is necessary to find some way to define the flame front on a numerical grid which
extends over a large spatial domain. Although there has been some success in the application
of adaptive gridding techniques to this problem,\textsuperscript{15-16} we know of no attempts to apply them to
calculations where a turbulence model must be used to define combustion rates. The reason for
this is that the resulting turbulent flame profile may not be particularly well-justified theoretically,
so there is no point in devoting a great deal of computational effort to its definition. It is also true
that the geometric complexity of the present experiments would make application of adaptive
gridding very difficult.

For these reasons, we have developed a combustion model which combines some of the
features of the Magnussen-Hjertager and eddy breakup models with some artificial definitions
of turbulence parameters in the flame zone for flame propagation at low turbulence levels. This
allows us to resolve the flame thickness over a number of grid points sufficient for good numerical
behavior, and to specify the burn velocity as a realistic function of turbulence level, pressure, and
temperature of the unburned gas. It is important to note that, while we thereby do not compute
the burn velocity from first principles, we are able to achieve fairly good agreement with many
aspects of the experiment with only minimal adjustment of a small number of parameters.

In the next section, we give a brief description of the Conchas-Spray computer code\textsuperscript{17}
which we have used for the calculations. (The simulations described here have been performed
in a two-dimensional domain, but the model itself is not restricted to two dimensions.) We also
discuss some features of the version of the $k-\epsilon$ turbulence model that we have implemented
into the computer code. In Section III, we derive the new features of our combustion model and
discuss its capabilities when integrated into the code.

Results of the simulation of an actual experiment in the FLAME facility are presented
in Section IV. The reason for the concentration of effort on only one experiment is that only
three experiments have been performed within the particular configuration to which our model
currently applies, and they do not cover a large range of hydrogen concentrations. Both the
experiments and the computer calculations are expensive; this unfortunately limits the data base
and the number of simulations permissible.

A particularly noteworthy result is that there is a period of time during which the computed
flame velocities and pressures agree quite well with the experimental values. During that time
period, the flow velocities are approximately sonic, and it appears likely that these velocities
are determined primarily by a choking mechanism. The pressure signals are characterized by
oscillations determined by the burnout of the individual chambers formed by the obstacles.

Some conclusions and recommendations are offered in the final section of the paper.
II. COMPUTER CODE AND TURBULENCE MODEL

Flow Variables and Chemistry

The Conchas-Spray code (modified to include the $k$-$\epsilon$ turbulence model) solves the following system of equations for the flow variables:

\begin{align}
\frac{\partial \rho_i}{\partial t} + \nabla \cdot (\rho_i \mathbf{u}) &= \nabla \cdot \left[ D \nabla (\rho_i \rho) \right] + \dot{\rho}_i^c \quad (1) \\
\frac{\partial (\rho \mathbf{u})}{\partial t} + \nabla \cdot (\rho \mathbf{u} \mathbf{u}) &= -\nabla p + \nabla \cdot \sigma \quad (2) \\
\frac{\partial (\rho I)}{\partial t} + \nabla \cdot (\rho I \mathbf{u}) &= -p \nabla \cdot \mathbf{u} + \nabla \cdot \left[ k_T \nabla T + \rho D \sum_i h_i \nabla (\rho_i / \rho) \right] + \dot{Q}_c + \rho \epsilon \quad (3)
\end{align}

where $\rho_i$ is species density, $\mathbf{u}$ is velocity, $I$ is specific internal energy, $\dot{\rho}_i^c$ and $\dot{Q}_c$ are the rates of change of $\rho_i$ and energy per unit volume due to chemical reactions, $k$ is the turbulent kinetic energy, and $\epsilon$ is the rate of dissipation of $k$. The index $i$ denotes the individual species. In this paper four species are considered: $H_2$, $O_2$, $N_2$, and $H_2O$. In the interest of conciseness, the identification and method of evaluation of many of the rest of the terms in these equations will only be briefly outlined. For further details, see References 17-19. The total mass density $\rho$ is obtained by summing the $\rho_i$ over all species. The temperature $T$ is computed from the energy $I$ by assuming that the species enthalpies $h_i$ are functions only of $T$. The pressure $p$ is then obtained from the ideal gas law. Diffusion of mass and heat are accounted for by the terms involving the mean diffusion coefficient $D$ and the thermal conductivity $k_T$. The stress tensor $\sigma$ is given by

\begin{equation}
\sigma = -\frac{2}{3} \rho k \delta + \mu \left[ \nabla \mathbf{u} + \nabla \mathbf{u}^T - \frac{2}{3} (\nabla \cdot \mathbf{u}) \delta \right] \quad (4)
\end{equation}

where $\delta$ is the unit tensor, the superscript $T$ denotes the transpose operator, and the viscosity is

\begin{equation}
\mu = \mu_L + \mu_T \quad (5)
\end{equation}

where $\mu_L$ is laminar viscosity and $\mu_T$ is turbulent eddy viscosity. The eddy viscosity is obtained from the $k$-$\epsilon$ turbulence model, and will be discussed later. The laminar viscosity is obtained from Sutherland's formula with appropriate coefficients, but is actually negligible in our calculations.

The term $\epsilon$ in (3) is the rate of dissipation of turbulent kinetic energy per unit mass. (The dissipation term describes the conversion of turbulent kinetic energy into heat; hence it is an
additive term in the energy equation (3). Equations (1)–(3) are in Favre-averaged form, assuming that the turbulence model can appropriately account for the Reynolds stresses via Equations (4) and (5), and that \( \epsilon \) can be correctly computed. We are neglecting differential species diffusion by using an average value of diffusion coefficient \( D \). This is not well-justified for a hydrogen-air mixture, but in view of the fact that we are using an artificial model to define the flame zone, this discrepancy does not detract from the results.

The thermal conductivity \( k_T \) and diffusion coefficient \( D \) are related to viscosity \( \mu \) by

\[
k_T = \frac{c_p \mu}{P_r} \tag{6}
\]
\[
D = \frac{\mu}{\rho S_c} \tag{7}
\]

where \( c_p \) is the specific heat at constant pressure and \( P_r \) and \( S_c \) are turbulent Prandtl and Schmidt numbers. We have used \( P_r = S_c = 0.52 \). However, because of the aforementioned flame front modeling, our results are not sensitive to this choice.

The following quantities appearing in the equations remain to be determined: The chemical contribution to the species conservation equation \( \dot{\rho}_i^c \), and the chemical heat release \( \dot{Q}_c \). These terms describe processes that are too complex to be evaluated in detail numerically for the present experimental configuration. Our model for them will be derived in Section III.

In this work, the chemistry is assumed to consist of the global one-step reaction

\[
2H_2 + O_2 \rightarrow 2H_2O \tag{8}
\]

In this case, given the \( \rho_i \)'s and \( h_i \)'s, all the chemical rates \( \dot{\rho}_i^c \) and \( \dot{Q}_c \) are completely defined by specifying only one of the species production rates. (We choose to use \( \dot{\rho}_{H_2}^c \) for this purpose.)

At this point, it is convenient to define one more variable for later use. Models which assume that combustion is limited by the rate of thermal mixing usually make use of a progress variable, or something related to such a quantity. Our progress variable is defined as

\[
C = \frac{Y_p}{Y_p^\infty}
\]

where \( Y_p \) is the mass fraction of the product species and \( Y_p^\infty \) is the mass fraction of product that would result if the locally extant mixture was allowed to react to completion. (Subscripts \( f \) and \( p \) will be used for fuel and product henceforth. In our case these correspond to \( H_2 \) and \( H_2O \).) Then

\[
Y_p^\infty = \frac{\rho_p + \frac{\nu_{f,m_p}}{\nu_{i,m_f}} \rho_f}{\rho}
\]

where the \( \nu_i \) are stoichiometric coefficients (both equal to 2 in our one-step \( H_2 \)-air reaction–see Equation (8)), and the \( m_i \) are the corresponding molecular weights. Hence,
\[ C = \frac{\rho_p}{\rho_p + \frac{\nu_p m_p}{\nu_f m_f} \rho_f} = \frac{n_p}{n_p + \frac{\nu_p n_f}{\nu_f n_f}} \]  

where \( n_f \) and \( n_p \) are the number densities of the fuel and product species. The quantity \( C \) always satisfies \( 0 \leq C \leq 1 \). (Note that the denominators in Equation (9) are the densities of product species instantaneously available under conditions of complete reaction of a lean mixture (at constant volume), and \( C = 1 \) when the fuel is completely consumed. This equation can be used for a fuel-rich mixture; if the subscript \( f \) is replaced by \( o \) (denoting the oxidizer), \( C = 1 \) when the oxidizer is consumed.)

The computer code used was an adaptation of the Conchas-Spray code developed by Los Alamos National Laboratory. It provides a solution to Equations (1)–(3) in two dimensions; the present calculations are performed in rectangular coordinates. The numerical algorithm employed in the code is a finite-difference approximation derived from an integral formulation of the conservation laws. For details see References 17 and 20.

All the computations employ an Eulerian method on a grid with square cells (see Figure 3). The thickness of the obstacles is taken to be two grid spacings for numerical simplicity. (The experimental obstacles were constructed of 13 mm (1/2 inch) plywood, and were therefore considerably thinner. However, the difference has negligible effect on our results.) The \( z \)-axis lies in the midplane of the facility, which is assumed to be a symmetry plane. Hence, the grid on which the computations were carried out encompassed only half the area of the facility. Except where noted otherwise, the grid consists of 13 points in the (transverse) \( x \)-direction by 397 grid points in the (axial) \( z \)-direction. This encompasses sixteen and one-half chambers formed in the channel by the obstacles used in the experiments. This is a very coarse grid; it is not adequate to resolve recirculating flows behind the obstacles with precision. Furthermore, errors will be incurred in the computation of the shock waves which form when the gas flows past obstacles at supersonic speeds. We have investigated the numerical accuracy to the extent possible by refining the grid spacing by a factor of 2/3. The results (discussed in Section IV) show some differences; nonetheless, the coarse grid suffices for calculations which provide insight into the processes involved.

The \( k-\epsilon \) Turbulence Model

The turbulence model which we have used is the \( k-\epsilon \) model as described in References 9 and 10. Although Reference 9 specifically addresses the question of compressible flows, it should be noted that turbulence models for compressible flows are not well-developed. Hence, the choice of the standard \( k-\epsilon \) model in this situation cannot be regarded as definitive.

The model consists of the following transport equations for the evaluation of turbulent kinetic energy \( k \) and dissipation rate \( \epsilon \).

\[
\frac{\partial}{\partial t} (\rho k) + \nabla \cdot (\rho ku) = \sigma \nabla u + \nabla \cdot (\mu \nabla k) - \rho \epsilon \tag{10}
\]

\[
\frac{\partial}{\partial t} (\rho \epsilon) + \nabla \cdot (\rho \epsilon u) = C_{1\epsilon} (\sigma \nabla u) \frac{\epsilon}{k} + \nabla \cdot (\frac{\mu}{\sigma \epsilon} \nabla \epsilon) - \frac{C_{2\epsilon} \rho \epsilon^2}{k} \tag{11}
\]
Figure 3. Problem domain and finite-difference grid used in most of the calculations. The full $16 \frac{1}{2}$ chambers into which the facility is divided is shown on the left. The grid is too fine to be resolved on the scale used there. An expanded view of 3 chambers is given on the right.
The terms on the right sides of these equations represent (from left to right) the production, diffusion, and dissipation of the quantities $k$ and $\epsilon$. Strictly speaking, Favre averaging requires the inclusion of a term in (11) involving the correlation between velocity fluctuations and the pressure. Following a practice often used, we omit this term to avoid undue complexity in a turbulence model which is still under development for compressible flows. The coefficients $C_{1\epsilon}$ and $C_{2\epsilon}$ are modeling constants, and $\sigma_\epsilon$ is a Schmidt number. These constants have been tuned for agreement with certain experiments, with optimum values $C_{1\epsilon} = 1.92$, $C_{2\epsilon} = 1.44$, and $\sigma_\epsilon = 1.3$. (The experiments used for tuning are much simpler than those addressed here. Furthermore, they involve incompressible flows. Hence, the use of these constants should not be regarded as definitive.)

The turbulent part of the viscosity (the eddy viscosity) is given by

$$\mu_T = C_\mu \frac{k^2}{\epsilon}$$

(12)

The coefficient $C_\mu$ is set equal to the usual value of 0.09 (see, e.g., Reference 10).

Since $k$ and $\epsilon$ are defined to be nonnegative, it is important that the numerical scheme preserve this property. The production and dissipation terms in the $k$ and $\epsilon$ equations require special consideration in this regard. Since turbulent regimes are sometimes characterized by the fact that production approximately equals dissipation, care must be taken to ensure that numerical errors in either term do not result in negative $k$ or $\epsilon$. A special numerical scheme was devised to alleviate such difficulties. It is given in Appendix A.

In the Conchas-Spray code, the boundary conditions on tangential velocity are obtained by implementing the law of the wall for turbulent boundary layers. To this end, the tangential velocity at walls is not zero, but is allowed to vary to account for fluid momentum in the wall grid cells, which are in the boundary layer. We have essentially retained the numerical boundary condition as in the original code except that we restrict the velocity in a cell at a wall to be less in magnitude than the tangential velocity in the adjacent cell one grid point in from the wall. This prevents occasional nonphysical profiles from occurring.

The boundary conditions on $k$ and $\epsilon$ are likewise obtained from consideration of the law-of-the-wall shear layer. The wall shear stress $\tau$ is obtained by assuming that the tangential velocities at grid points one cell away from the wall are in the law-of-the-wall boundary layer. This allows one to solve a simple equation for the shear stress. Assuming that the Reynolds stresses are nearly constant, it can be shown that the turbulent kinetic energy in the wall cell may be approximated by

$$k = \frac{u_*^2}{\sqrt{C_\mu}}$$

where $u_*$ is the shear velocity

$$u_* = \sqrt{\tau/\rho}$$
This provides the boundary condition on $k$. The associated boundary condition on $\epsilon$ is

$$\epsilon = \frac{v_*^3}{\kappa y}$$

where $y$ is the distance to the wall and $\kappa = 0.4$ is the von Kármán constant. This $1/y$ behavior of $\epsilon$ can give rise to some numerical inaccuracy, especially in the diffusive flux of $\epsilon$. This can be alleviated by implementing a special differentiation formula near the wall. Details are given in Appendix B.

III. THE COMBUSTION MODEL

Properties of Turbulent Flames and Numerical Flames

As noted in the Introduction, simulation of combustion processes in large experiments places extreme demands on the capabilities of even the largest and fastest computers. In the present case, it is necessary to give up the idea of resolving the flame thickness on the computational grid. A possible exception arises in situations in which the turbulent length scale is very large and the turbulent flame thickness, defined as the thickness of a region where burned and unburned mixture both exist, extends over several grid points. But for satisfactory numerical behavior we must usually thicken the flame artificially. To illustrate this point, a plot of the laminar, turbulent, and computational flame profiles are given in Figure 4. Turbulence results in rapidly fluctuating wrinkles and spirals of interpenetrating burned and unburned gas, as shown in the upper sketch in the figure. The turbulent flame thickness $\delta_T$ is defined by the region where the composition is intermittent due to the fluctuations, and the average value of $C$ is greater than zero, but less than unity. For good numerical behavior, the computational flame thickness $\delta_c$ must be equal to a few grid spacings. (The size of one grid interval is indicated by $\Delta x$. The scale in the figure is not necessarily accurate; the various lengths actually satisfy

$$\frac{\delta_c}{4} \approx \Delta x > \delta_T > \delta_L$$

Artificial flame thickening has been done previously by a method known as the $\beta$-transformation.\textsuperscript{23,24} This involves changing the length scales in the problem by artificially increasing the thermal diffusivity and diffusion coefficient and decreasing the combustion rate. We have retained this essential idea. However, the original $\beta$-transformation established the scaling through the use of the local temperature gradient in the problem. Numerical experimentation indicated that this did not work well with mixing-limited combustion models such as the Magnussen-Hjertager model. Hence, it was decided to achieve the flame thickening in a different way.

Given that the flame will be somewhat artificial, it is necessary to first determine what properties it should have. In particular, it is desirable that the burn velocity be realistic. The
Figure 4. Schematic illustration of laminar flame thickness, turbulent flame thickness, and computational flame thickness. Above: A slice through a turbulent flame, plotted in \((x,y)\) coordinates. The thickness of the dark lines is the laminar flame thickness \(\delta_L\). Below: A plot of the progress variable \(C\) (defined in Equation (9)), for \(y\) equal to the constant value defined by the dashed line in the upper figure.
following properties of turbulent flames have been extracted from a list given in Reference 13 ($S_L$ and $S_T$ refer to laminar and turbulent burn velocities, respectively):

1. $S_T > S_L$ (almost always)
2. $S_T$ increases with $u' \equiv \sqrt{k}$. Often, the increase is linear, and

$$S_T = A S_L + B u'$$

provides as good a correlation as anything else*.

3. Sometimes $S_T$ is insensitive to the scale of approach flow turbulence.
4. In open flames, the variation of $S_T$ with composition is much the same as for $S_L$.
5. In flames in ducts, $S_T$ is larger than for open flames and is sometimes insensitive to approach flow turbulence and composition.

Our model represents an attempt to adhere to these properties, except for the latter clause in Number 5. It does not necessarily apply to our experimental configuration, as it was inferred from experiments on steady-state flames stabilized in small ducts.

To see how to embed the above properties 1 through 4 in a combustion model, we note that, for the case of mixing-limited combustion models such as Magnussen-Hjertager or eddy breakup, simple flat flames will be expected to have the following properties (see, for example, the discussion of laminar deflagrations in Section 5.1.2 of Reference 19):

The turbulent burn velocity will scale as

$$S_T \sim \sqrt{\chi/t_T}$$  \hspace{1cm} (13)

where $\chi$ is the thermal diffusivity (assumed proportional to the diffusion coefficient—see Equations (6) and (7)) and $t_T$ is the turbulent time scale. (This is commensurate with the assumption of burning limited by thermal mixing, where the rate of combustion is proportional to $1/t_T$).

The turbulent flame thickness will scale as

$$\delta \sim \sqrt{\chi t_T}$$  \hspace{1cm} (14)

From the $k$-$\epsilon$ turbulence model,

$$\chi \cong C_\mu \sqrt{k L / Pr}$$

and

* Note that $u'$ is defined here as the average turbulent velocity, and not as the instantaneous velocity fluctuation.
\[ t_T = \frac{L}{\sqrt{k}} \]  

where

\[ L = k^{3/2}/\epsilon \]

is the integral length scale of turbulence. From (13) and (14), then

\[ S_T \propto \sqrt{k} = u' \]

\[ \delta \propto L \]

In other words, if the turbulence intensity \( \sqrt{k} \) and the length scale \( L \) are given for the gas, one can achieve a model which complies with the linear part of Property 2 and with Property 3.

**Development of a Model With Appropriate Properties**

In applying a mixing-limited combustion model to a computation such as this, it must be recognized that difficulties may occur when starting up a problem when the turbulence level is low. Initially, one expects the flame to be laminar, but in our experimental configuration this will be quickly superseded by a flame which exhibits some form of turbulence, e.g., a wrinkled laminar flame.\textsuperscript{13,19} Data from a variety of experiments in the FLAME facility suggest that the hydrogen-air flames studied in the facility achieve a relatively constant burn velocity which is greater than laminar and which persists until large-scale instabilities and/or obstacles cause the flame to accelerate.\textsuperscript{2,3,25}

The approach that we have developed is to endow the gas with minimum values of \( k \) and \( L \), denoted \( k_m \) and \( L_m \), which are used for the computation of thermal conductivity and diffusivity only. In that way, according to Equations (17) and (18), there will be minimum values of burn velocity and flame thickness. The minimum burn velocity provides a term like \( AS_L \) in property 2, although not in such a way that the equation there is exactly satisfied. The scheme which we employ is to use \( k_m \) and \( L_m \) in the formulas for \( k_T \) and \( D \) unless the true turbulence intensity \( k \) or length scale \( L \) predicted by the \( k-\epsilon \) model exceeds those threshold values. Then we use \( k \) and/or \( L \). It is crucial to note that only the true values \( k \) and \( L \) (or, equivalently, \( \epsilon \)) are used in the viscosity. In particular, they are used in the transport equations (10) and (11) for \( k \) and \( \epsilon \) themselves, and in (12) for \( \mu_T \), so that \( k_m \) and \( L_m \) affect the generation of turbulence only indirectly. (It may appear that this indirect effect is a very strong one, since the gas velocities appear in the turbulence production terms in (10) and (11). However, if \( k_m \) and \( L_m \) are chosen correctly, these gas velocities would be essentially the same as those occurring in a perfectly resolved flame.)

Now, how should \( k_m \) and \( L_m \) be determined? It would be desirable that \( k_m \) correspond to something physically realistic, because the point at which \( k \) exceeds \( k_m \) should represent some real point in parameter space at which the flame starts to propagate more rapidly.
We have extracted a reasonable choice from the work of Libby, Bray, and Moss (LBM).\(^{26}\) In the limit of large turbulent Reynolds number \(R_T \equiv \rho u'L/\mu\), they obtain a formula

\[
S_T = S_L + 1.14\sqrt{k}
\]  

(19)

This formula is derived by making certain modeling assumptions regarding physical processes in flames. It is not intended to be universally valid, even in the limit \(R_T \to \infty\). In fact, References 13 and 26 demonstrate clearly that no universal formula exists. But we choose to use this as a starting point. Given that we can extract an initial turbulent burn velocity \(S_{T_i}\) from experimental data, we obtain a provisional estimate for \(k_m\) by inverting (19):

\[
k_m = \left( \frac{S_{T_i} - S_L}{1.14} \right)^2
\]  

(20)

We emphasize that this is only an estimate. As discussed below, it is impossible to satisfy all the desired flame properties and still adhere to this value of \(k_m\).

We now give our formula for the rate of consumption of fuel. (As noted in the discussion following Equation (8), for a one-step reaction, specification of the rate for only one species suffices to specify all chemical rates.) Except for a dependence on pressure and unburned gas temperature to be discussed later, the combustion rate is

\[
\dot{\rho}_f = -A_c \rho f(T) C(1 - C)/t_T
\]  

(21)

where \(A_c\) is a dimensionless constant, and

\[
f(T) = \begin{cases} 
1 & T_a < T \\
\frac{T - T_o}{T_a - T_o} & T_o < T \leq T_a \\
0 & T < T_o
\end{cases}
\]

where \(T_a\) is adiabatic flame temperature and \(T_o\) is the initial gas temperature. Note that the factor \(C(1 - C)\) is similar to that appearing in approximations to the eddy breakup model (see Equation (4.72) of Reference 13) and to the minimum normalized mass fraction appearing in the Magnussen-Hjertager model\(^{11}\). The function \(f(T)\) serves a special numerical purpose. It does not really represent any physical process, except that it provides a qualitative increase in combustion rate with temperature up to the adiabatic flame temperature. It is inserted in Equation (21) so that the flame profile will be determined by thermal diffusion in such a way as to obey Equations (13) and (14), with the result that (17) and (18) are satisfied. Lacking the factor \(f(T)\), Equation (21) results in flame zones that tend to diffuse too far ahead of and behind the flame, and which do not exhibit the functional behavior specified in (13) and (14). This spreading is predictable on the basis of comparison with solutions to the Kolmogoroff-Petrovsky-Piscounoff (KPP) equation\(^{27-29}\).
At this point we have three adjustable parameters: \( k_m, A_c \) and \( L_m \). They are fixed by simultaneously specifying the following three flame properties: The initial turbulent burn velocity \( S_{T_i} \), the derivative \( S_T' = \frac{dS_T}{d\sqrt{k}} \) for adiabatic flames, and an appropriate numerical flame thickness. Our initial choice for \( dS_T/d\sqrt{k} \) is 1.14, in accordance with (19). A variation on this value will be discussed in Section IV. (The actual values of \( k_m, A_c \), and \( L_m \) are determined by adjusting them in a series of one-dimensional flat flame calculations.) In most of our calculations, we have attempted to maintain a 10-90% thickness of 4 grid points. As noted above, our model does not precisely satisfy Equation (19). It does, however, provide values for the initial burn velocity and slope of the \( S_T \) vs \( \sqrt{k} \) curve which are physically well-justified. (See Figure 5.)

**Dependence on Pressure and Preflame Temperature**

Finally, we describe the way in which the combustion model is augmented to include a dependence on pressure and the temperature of the unburned gas through which the flame is propagating. As the flame travels down the obstructed channel of the FLAME facility, the unburned gas in front of it is compressed due to the overall drag forces which build up. This compression results in an increase in pressure and temperature. Since the laminar burn velocity of a hydrogen-air mixture depends on the pressure and temperature of the unburned gas, the turbulent burn velocity will also exhibit such a dependence. In the spirit of property 4 which is desired for the combustion model, we simply choose to impose the same functional dependence of \( S_T \) on \( p \) and unburned gas temperature \( T_u \) as is expected for the laminar burn velocity \( S_L \). Warnatz\(^30\) has shown that in certain regimes \( S_L \) varies according to

\[
S_L \propto p^m T_u^n
\]

Consideration of Equations (13) and (14) indicates that, all else remaining constant, \( S_T \) will exhibit the same behavior if the burn rate, thermal diffusivity, and diffusion coefficient are multiplied by a factor

\[
h(p,T_u) = \left( \frac{p}{p_0} \right)^m \left( \frac{T_u}{T_0} \right)^n
\]

where \( p_0 \) is the initial pressure and \( T_u \) is the unburned gas temperature. We have used the values \( m = 0.2, n = 1.64 \) as being representative of Warnatz’ results; this choice is by no means clear-cut, however.\(^{30}\)

We have employed \( T_u \), the *unburned* gas temperature, to alter the burn rate and diffusivities. But \( \hat{\rho}, \chi, \) and \( D \) vary *through* the flame. How do we know what \( T_u \) is at points inside the flame? This is determined by extrapolating back to an unburned mixture at the same pressure. (It is assumed that the pressure does not vary significantly through the flame.) We write

\[
\int_{T_u}^{T} c_p(T) dT = \frac{\Delta H}{m_p} Y_p
\]
Figure 5. Turbulent burn velocity vs turbulence intensity for the Libby-Bray-Moss model\(^{26}\) (chain-dashed line with slope 1.14) and for the present model (solid line). The difference between the two at low turbulence intensity results from the choice of numerical mechanism available in the present case, and not from any suggestion that the physical situation is better represented. The dashed line represents a variation in the model in which the combustion rate is reduced (see discussion in Section IV). The scales in the figure represent a realistic range of values encountered in the calculations; typical values of \(\sqrt{k}\) in the choking regime are about 30 times \(\sqrt{k_m}\).
where \( c_p(T) \) is the specific heat of the mixture and \( \Delta H \) is the molar heat of reaction (57.8 Kcal/mole for \( H_2O \)). We have obtained a linear fit to the specific heat from data available in the CHEMKIN code. This approximation has the form

\[
c_p = c_{po} + c'_p(T - T_u)
\]  

(24)

When (24) is substituted into Equation (23), there results a quadratic equation for \( T_u \); the solution is substituted into Equation (22) to obtain the function \( h(p,T_u) \).

### Summary of the Model

This completes the description of the combustion model. The implementation of the material presented in the previous section requires a redefinition of the thermal conductivity and diffusion coefficient and an alteration of the formula for combustion rate given in equation (21). For convenience, the appropriate formulas are collected in the following:

\[
k_T = \frac{c_p\mu'}{Pr}
\]  

(6')

\[
D = \frac{\mu'}{\rho S_c}
\]  

(7')

\[
\dot{p}_f = -A_c \rho f(T) h(p,T_u) C (1 - C) / t'^T
\]  

(21')

where

\[
\mu' = \mu_L + \mu'_T
\]  

(5')

\[
\mu'_T = C \mu \rho \sqrt{k_e} L_e
\]  

(12')

\[
t'^T = L_e / \sqrt{k_e}
\]  

(15')

and \( k_e \) and \( L_e \) are effective values of \( k \) and \( L \), defined as

\[
k_e = \max\{k,k_m\}
\]  

(25)

\[
L_e = \max\{L,L_m\}
\]  

(26)

The model has been tested by running one-dimensional calculations of adiabatic flames at constant pressure. For this purpose, the \( k-\epsilon \) model was turned off so that \( k_m \) and \( L_m \) would
determine the flame properties. In such calculations the flame achieves a self-similar profile propagating at an approximately constant velocity. As indicated in the foregoing discussions, the desired behavior of the flame velocity and the computational flame thickness \( \delta_c \) vary according to

\[
S_T \propto \sqrt{k_e} p^m T_u^n
\]  

(27)

\[
\delta_c \propto L
\]  

(28)

Furthermore, \( S_T \) and \( \delta_c \) should be independent of the grid size. Table I gives some results obtained by varying the parameters around a representative baseline set. (Anticipating the simulations to be discussed in Section IV, the baseline calculation corresponds to a 14.5% hydrogen-air burn at atmospheric pressure and 300°K with \( k_m, L_m, \) and \( A_e \) chosen appropriately for one of the calculations.) The agreement between the computational values and the desired values is quite good, except possibly for the pressure and temperature dependences. The reason for the discrepancy in the case of the temperature variation is that there is some inaccuracy in the determination of \( T_u \) via Equations (23) and (24). Since there is some arbitrariness in the choices for \( m \) and \( n \) which are used in Equation (22), these discrepancies will be accepted for the remainder of this paper. For the most part, the model achieves its goals for the simple flat (one-dimensional) flame. In particular, it may be noted that the first three lines in the table verify that the slope \( dS_T/d\sqrt{k} \) is approximately 1.14, as required by Equation (19).

In summary, the characteristics of the model are as follows: It is designed to provide for the consumption of fuel at a rate consistent with certain assumptions as to the phenomenological behavior of turbulent flame propagation. It permits the computation of the behavior of a compressible fluid ahead of and behind the flame front; it thereby accounts for the interaction of turbulence with the flame insofar as the assumptions relating the turbulent time scale to the combustion rate are correct. It does not permit a computation of burn velocity from a knowledge of only the state of the gas and the turbulent flow structure; it requires prior assumptions as to what the burn velocity would be in a given situation.

It should also be noted that, in the form presented here, the fluid model will exhibit a non-physical heat conduction and particle diffusion everywhere because of the universal application of the (artificially high) threshold values of \( k \) and \( L \) in Equation (12'). For the present application, errors due to this effect are negligible. To see this, consider the energy conservation equation (3). We wish to show that the artificial heat conduction is negligible on the time scales typical of changes in the specific energy. Suppose that the typical time scale \( t_t \) is the time for the initial flame to propagate the width of the channel:

\[
t_t = \frac{w}{v_{fo}}
\]

where \( w \) is channel width and \( v_{fo} \) is the initial flame speed. If \( w \) is used as the thermal diffusion length scale, the ratio of the diffusion time \( t_x \) to \( t_t \) is
Table 1.

Results of tests of the combustion model. The quantity $\Delta x$ is the grid spacing. The burn velocity and flame thickness resulting from the model are $S_T$ and $\delta_c$; the values of these quantities expected on the basis of simple theory are denoted $S_T^c$ and $\delta_c^c$. The constant $A_c$ (see Equation (21)) is equal to 30.8. The exponents $m$ and $n$ (see Equation (22)) are equal to 0.2 and 1.64.

<table>
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<th>$k_m$(J/kg)</th>
<th>$L_m$(cm)</th>
<th>$p$(atm)</th>
<th>$T_a(°K)$</th>
<th>$\Delta x$(cm)</th>
<th>$S_T^c$(m/s)</th>
<th>$S_T$(m/s)</th>
<th>$\delta_c^c$(cm)</th>
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<td>300</td>
<td>7.62</td>
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<td>16</td>
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<tr>
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</table>
For typical values \( v_f = 16 \) m/s, \( w = 1.83 \) m, \( \Pr = 0.52 \), \( C_{\mu} = 0.09 \), \( k_m = 4.7 \) J/kg, \( L_m = 0.68 \) m, this ratio is 115. Hence, the diffusion time is long compared to the time during which other processes are occurring, and the artificial diffusion is negligible (except in the flame, of course). As time elapses and the flame accelerates, \( t_t \) decreases and the relative error incurred becomes smaller. Inspection of the dispersion relation for acoustic waves shows that the attenuation of such waves is similarly negligible over a propagation distance of a channel width. The attenuation of acoustic waves that propagate the entire length of the FLAME Facility might not be negligible, but they would not be expected to contribute significantly to the results of the present study.

The effects of this nonphysical diffusion could probably be eliminated entirely if necessary. All that would be required is to remove the thresholds on \( k \) and \( L \) away from the flame. This would require some care, as it would be necessary to ensure that the resulting decrease in diffusivities was kept far enough away from the flame zone that the flame propagation velocity was not affected.

Finally, we note that no attempt has been made in this work to account for the effects of flame stretch and flame curvature in detail. (A general discussion is given in, e.g., Reference 19.) It is impossible to compute such effects in this case, since the flame is artificially thickened, but one might attempt to model them (see, e.g., the model for flame quenching in Reference 32).

IV. COMPARISON OF COMPUTATIONAL RESULTS WITH EXPERIMENT

Experimental Considerations

A total of 9 experiments in which obstacles were present have been performed in the FLAME facility. Table II gives a brief description of these tests and their results. In Tests F-21 through F-23, the facility was unvented; in F-24 through F-29, one-half of the top plates were removed, resulting in 50% venting. In the present report, we are considering only the unvented experiments. In experiment F-21, failure of the mixing fans resulted in uncertain hydrogen concentrations. Hence, only experiments F-22 and F-23 are available to us for comparison with the type of computations with which we are concerned.

The only difference in the experimental arrangement in F-22 and F-23 is that the initial hydrogen concentrations were 15% and 14.5%, respectively. In experiment F-23, the mixture appeared to detonate; in F-24 it did not. Since turbulent flows and conditions leading to detonation are random in nature, the 0.5% difference in concentrations in these two experiments
Table II.

Summary of FLAME tests with obstacles. (From Reference 34.) A total of 16 pairs of obstacles with a blockage ratio of 1/3 were used in the experiments. (See Figures 1 and 2.)

<table>
<thead>
<tr>
<th>Test Number</th>
<th>Top Venting Area, %</th>
<th>Hydrogen Mole Fraction, %</th>
<th>Peak Overpressure, atm</th>
<th>Peak Planar Flame Speed m/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>F-21</td>
<td>0</td>
<td>10-15(^a)</td>
<td>6.5</td>
<td>580</td>
</tr>
<tr>
<td>F-22</td>
<td>0</td>
<td>15.0</td>
<td>31.0</td>
<td>700</td>
</tr>
<tr>
<td>F-23</td>
<td>0</td>
<td>14.5</td>
<td>12.0</td>
<td>540</td>
</tr>
<tr>
<td>F-24</td>
<td>50</td>
<td>15.5</td>
<td>(b)</td>
<td>46</td>
</tr>
<tr>
<td>F-25</td>
<td>50</td>
<td>19.7</td>
<td>16.5</td>
<td>(\bullet) 890</td>
</tr>
<tr>
<td>F-26</td>
<td>50</td>
<td>28.5</td>
<td>19.7</td>
<td>1860</td>
</tr>
<tr>
<td>F-27</td>
<td>50</td>
<td>13.1</td>
<td>.09?</td>
<td>15?</td>
</tr>
<tr>
<td>F-28</td>
<td>50</td>
<td>14.9</td>
<td>.09?</td>
<td>33.4</td>
</tr>
<tr>
<td>F-29</td>
<td>50</td>
<td>18.5</td>
<td>.22?</td>
<td>130</td>
</tr>
</tbody>
</table>

\(^a\) - mixing fans did not operate.

\(^b\) - signal in noise level.

? - analysis not complete.
is not necessarily significant. In other words, it is not necessarily true that a 15% concentration would always lead to a deflagration-to-detonation transition (DDT), while a 14.5% concentration would not. Our computations are not capable of the spatial and temporal resolution or the detailed physics and chemistry required to provide a true simulation of DDT. Nonetheless, it will be seen that we observe a "numerical" transition. Hence, we will concentrate on a comparison with experiment F-23, with a brief observation on detonation with a view to experiment F-22.

In the FLAME facility, flame arrival times are measured by detecting temperature increases at 5 thermocouple "rakes" positioned vertically on the channel midplane. (See Figure 2.) In the present set of experiments, each rake has thermocouples at 7 vertical locations. From this arrival time data, it is possible to construct rough vertical profiles of the flame as it propagates down the channel. The arrival time data for experiments F-22 and F-23 and the corresponding flame front profiles are given in Figures 6-9. The flame velocity obtained from the slope of the line connecting the last two data points in Figure 6 is only 700 m/s. This is below the Chapman-Jouguet (CJ) detonation velocity of 1520 m/s, but it is believed that the discrepancy is due to the poor resolution available from the thermocouple data. Presence of a detonation was inferred from visual observation. Note also the rapid propagation of the flat profiles which begins sometime between 300 ms and 320 ms in Figure 7. This is indicative of a detonation.

In addition to the thermocouples, there are pressure transducers located at various sites on the side walls. Data from these transducers will be compared with the computational results below.

**Comparison of Flame Trajectories and Pressure Histories**

The first calculation carried out to simulate Experiment F-23 was made with the coefficients in the combustion model selected precisely as described in Section III. (It will be referred to as the baseline calculation.) Parameters pertinent to the test configuration and the computation are collected in Table III. Ignition in these computations is accomplished by forcing a controlled burn over a small region at the closed end of the channel. This ignition process ends at a time of 50 ms. Some care must be taken in implementing ignition in these simulations; a discussion of this point is given later in this section. The initial computational burn velocity $S_{Ti}$ was chosen as 2.5 m/s so that the initial computational flame speed would correspond to the experimental value of 20 m/s. It is to be emphasized that, for this calculation, this is the only parameter adjusted to agree with experiment. Note that the flame speed expected for an adiabatic flat flame with a constant burn velocity in an unobstructed tube would be

$$v_T = rS_{Ti}$$

where $r$ is the volume expansion ratio. For a 14.5% hydrogen-air mixture, $r$ is 4.45. Hence, according to the above formula, $v_T$ would be 11 m/s. The reason that the experimental flame speed is greater than this is that the flame is not flat, but is stretched out, primarily by the obstacles. As an additional point of reference, it may be noted (see Table III) that the laminar burn velocity at this concentration is 0.9 m/s. The existence of a wrinkled laminar flame or a weakly turbulent flame with burn velocity equal to, say, 2.5 times the laminar burn velocity is consistent with other experimental results and theoretical considerations.13,19.
Figure 6. Flame arrival time data for Experiment F-22. (This and the following 3 figures were provided by Reference 34.) The elevations of the thermocouples are given in the key. The distances of the thermocouple rakes from the closed end of the channel are: 1.0 m, 3.8 m, 11.6 m, 19.8 m, and 37.8 m (see the locations of the sets of data points).
Figure 7. Flame front profiles for Experiment F-22. A side view of the facility is shown (the section is through the midplane, where the thermocouple rakes are located). The upper figure is to scale. The lower figure is expanded in the vertical direction, showing the times in milliseconds to which the contours correspond. The data points indicate the vertical positions of the thermocouples (but not their axial locations—these profiles are obtained by interpolation).
DEVELOPMENT AND APPLICATION OF
A COMPUTER MODEL FOR LARGE-SCALE
FLAME ACCELERATION EXPERIMENTS

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Figure 8. Flame arrival times for Experiment F-23. (See Figure Caption 6 for explanation.)
Figure 9. Flame front profiles for Experiment F-23. (See Figure Caption 7 for explanation.) No detonation occurred in this case, although in general the flame propagation rates appear to be only slightly slower than for F-22.
Table III.
Summary of parameters pertinent to Experiment F-23 and the numerical simulation.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen mole fraction</td>
<td>0.145</td>
</tr>
<tr>
<td>Initial density</td>
<td>1.014 kg/m³</td>
</tr>
<tr>
<td>Initial temperature $T_i$</td>
<td>300°K</td>
</tr>
<tr>
<td>Initial pressure</td>
<td>1 atm</td>
</tr>
<tr>
<td>Initial sound speed</td>
<td>374 m/s</td>
</tr>
<tr>
<td>Adiabatic flame temperature</td>
<td>1438 °K</td>
</tr>
<tr>
<td>Volume expansion ratio $r$ for adiabatic flame</td>
<td>4.45</td>
</tr>
<tr>
<td>Laminar burn velocity $S_L$</td>
<td>0.9 m/s</td>
</tr>
<tr>
<td>Isobaric sound speed</td>
<td>758 m/s</td>
</tr>
<tr>
<td>Isochoric temperature</td>
<td>1757 °K</td>
</tr>
<tr>
<td>Isochoric pressure</td>
<td>5.43 atm</td>
</tr>
<tr>
<td>Isochoric sound speed</td>
<td>894 m/s</td>
</tr>
<tr>
<td>Chapman-Jouguet detonation velocity</td>
<td>1500 m/s</td>
</tr>
<tr>
<td>Chapman-Jouguet pressure</td>
<td>11 atm</td>
</tr>
<tr>
<td>Initial experimental flame speed</td>
<td>20 m/s</td>
</tr>
<tr>
<td>Combustion rate coefficient $A_c$</td>
<td>30.8</td>
</tr>
<tr>
<td>Threshold turbulent kinetic energy $k_m$</td>
<td>4.74 J/kg</td>
</tr>
<tr>
<td>Threshold turbulent length scale $L_m$</td>
<td>0.68 m</td>
</tr>
<tr>
<td>Pressure exponent $m$</td>
<td>0.2</td>
</tr>
<tr>
<td>Temperature exponent $n$</td>
<td>1.64</td>
</tr>
<tr>
<td>Prandtl number $Pr$</td>
<td>0.52</td>
</tr>
<tr>
<td>Schmidt number $Sc$</td>
<td>0.52</td>
</tr>
<tr>
<td>Initial turbulent burn velocity $S_{Ti}$</td>
<td>2.5 m/s</td>
</tr>
<tr>
<td>Initial computational flame speed</td>
<td>20 m/s</td>
</tr>
</tbody>
</table>
It is interesting to note that the burn velocity of 2.5 m/s for a hydrogen concentration of 14.5% is between 1/2 to 2/3 of the value computed independently from an analysis of experiments without obstacles\textsuperscript{25}. Since non-planarity of the flame was ignored in obtaining the latter results, this lower value is reasonably consistent.

A comparison of the computed and experimental flame trajectories of one calculation with the corresponding experimental data taken from Experiment F-23 (14.5% hydrogen) is given in Figure 10. Considering the size of the experiment and the complexity of the computational problem, the agreement is quite good. To some degree, the discrepancies in the computed flame trajectory must be viewed in light of the fact that the flame velocity is extremely sensitive to the burn velocity. (Because of the geometrical factors and the feedback loop involving gas flow velocities, turbulence intensities, and combustion rates, the dependence is nonlinear.) This sensitivity notwithstanding, the computation underestimates the rapid acceleration of the burn rate in the region from 200 to 275 milliseconds. The reasons for this are not known. Precise comparisons with experiment are not possible, as no measurements of turbulence levels are available and it is not possible to instrument the experiments sufficiently to make detailed spatial plots of such flow parameters as flame position, pressure, and temperature.

At a time of about 300 ms into the calculation, however, the computation is accelerating rapidly. Eventually, the computational flame trajectory essentially parallels that of the experiment while the flame traverses through axial positions from about 17 m through 22 m.

Figures 11–14 give a comparison of the computed and experimental pressures at the locations indicated in Figure 10. Data from two pressure transducers at axial positions 18.6 m, 21.5 m, and 25.6 m are shown. Table IV gives complete information on the locations of all transducers pertinent to this paper. The computed pressures are taken from the finite-difference zone next to the channel wall. (Note: All of our calculations terminated when their flames reached the end of the channel. That is why the computed pressure histories end abruptly at 320 ms in Figures 11–14.)

The agreement between computation and experiment at 14.6 m is rather poor, as seen in Figure 11. This simply illustrates the point that the computed combustion rate is significantly slower than that occurring in the experiment at that location. (Compare the flame speeds, i.e., the slopes of the trajectories in Figure 10.) It is important to realize that the agreement with the experimental pressures in this figure could easily be improved by adjusting the parameters $k_m$, $L_m$, and $A_c$ until the flame speed in the region around $z = 14.6$ m approximately coincided with the experimental flame speed, but then the agreement would not be as good elsewhere (see below).

A significant improvement is apparent in Figure 12. The computed pressures agree reasonably well with those measured experimentally. The reason for this is that the structure of the pressure history is characterized by the timing of the burnout of the gas in the successive chambers. The fact that the flame velocities approximately agree means that the chambers are burning out at about the correct rate. It should be noted that the flame velocity of about 540 m/s is greater than the sound velocity in the unburned gas, but less than that in the burned gas. This is to be expected if the flow rates are reduced due to choking\textsuperscript{6}. Note also that the experimental pressure signals from different transducers at the same axial location are somewhat different.
Figure 10. Comparison of computed and experimental flame trajectories for Experiment F-23. The experimental data has been shifted 100 ms to the left (relative to Figures 5-9) to facilitate comparison. (In view of the lack of a detailed simulation of ignition, the relative origins of the computational and experimental time scales is somewhat arbitrary.) The calculation used the baseline parameters as indicated in the text and the first line of Table I. The horizontal dotted lines show the location of the pressure transducers (at approximately 14.6 m, 18.6 m, 21.5 m, and 25.6 m down the channel) which provided the data discussed in this paper. (The complete array of experimental pressure data is not restricted to these locations.)
Figure 11. Pressure histories at 14.6 m from the baseline calculation and from transducer PKU4 in the experiment. (See Figure 10 and Table IV for a precise description of transducer location.) In this and all other pressure-time histories, the experimental signal has been shifted in time so that the times at which the flame passes the pressure transducer agrees with that obtained from the computation.
Figure 12. Pressure histories at 18.6 m.
Figure 13. Pressure histories at 21.5 m. In this and the following figure, pressures computed on the axis of the facility are included for comparison (see text).
Figure 14. Pressure histories at 25.6 m.
Table IV.

Location of pressure transducers in Experiment F-23. (From Reference 34.) The coordinates $x$, $y$, and $z$ correspond to spanwise, vertical, and axial directions. (Positive and negative values of $x$ indicate transducers mounted on opposite sides of the channel.) All distances are in meters.

<table>
<thead>
<tr>
<th>Transducer</th>
<th>$x$</th>
<th>$y$</th>
<th>$z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>PKU4</td>
<td>-0.914</td>
<td>0.30</td>
<td>14.6</td>
</tr>
<tr>
<td>PKU5</td>
<td>-0.914</td>
<td>0.31</td>
<td>18.6</td>
</tr>
<tr>
<td>PKU6</td>
<td>0.914</td>
<td>2.12</td>
<td>21.5</td>
</tr>
<tr>
<td>PKU7</td>
<td>0.914</td>
<td>0.29</td>
<td>25.5</td>
</tr>
<tr>
<td>PKU8</td>
<td>-0.914</td>
<td>2.12</td>
<td>25.6</td>
</tr>
<tr>
<td>PKU9</td>
<td>-0.914</td>
<td>2.11</td>
<td>21.4</td>
</tr>
<tr>
<td>PKU10</td>
<td>0.914</td>
<td>0.21</td>
<td>18.7</td>
</tr>
</tbody>
</table>
This illustrates the complexity of the problem; these signals may differ because of different proximity to the floor or ceiling, or simply because of random fluctuations in the combustion and pressure waves.

It may be noted that the computed pressure signals do not show as much high-frequency structure as the experimental data. This is undoubtedly due to some numerical dissipation exhibited by the computational algorithm. This will be discussed below with regard to a computation performed on a finer grid.

The order of magnitude of the pressure pulses which occur when the flow is choked, or nearly so, can be estimated as follows: The peak burn rate in the corners in front of the obstacles is very high. Hence, it is reasonable to suppose that the burn there is essentially isochoric. As noted in Table III, the isochoric pressure for 14.5% \( H_2 \) at 3000 K is 5.4 atm (65 psi). This is seen to be roughly verified both experimentally and computationally in Figures 12 and 13. There is naturally some spreading due to dispersion, especially at locations far from the flame, and this reduces the amplitude of the pressure signals.

The agreement between computational and experimental pressures is not as good at the 21.5 m station (see Figure 13). The reason for this is that the numerical flame is burning more rapidly than was the case in the experiment; in fact, at this point the numerical flame is very close to a transition to detonation. It is interesting to note that for the case shown in Figure 13, the pressure computed on the axis of the facility agrees more closely with experiment than that computed in the zone next to the wall, where the experimental pressure is measured. The reason for this is believed to be that the computed combustion rate at the wall is too high, and that this is causing the transition to detonation.

Figure 14 shows the pressures at 25.6 m down the channel. Detonation has occurred in the computation. As noted above, no detonation was observed in the experiment, but the peak pressure of 13 atm (12 atm overpressure, as indicated in Table II) exceeds the Chapman-Jouguet pressure of 11 atm. The peak computed pressure of 48 atm far exceeds the CJ pressure as well. This may be due in part to numerical overshoot, but it is also true that in the computation, symmetric waves are reflecting off the side walls. Since the pressure in a reflected wave adds in phase and multiple reflected waves can occur, there is a strong mechanism leading to the computation of pressures higher than CJ. It should be noted that such collisions of strong waves would probably not be observed very often experimentally, since the symmetry imposed in the computation would not occur. The effect is enhanced because the calculation is carried out in two dimensions.

Further Discussion of the Computational Results

All of the behavior described above can be seen from a different perspective in Figures 15–18, which show the computed pressures, temperatures, turbulent kinetic energy, and rate of consumption of hydrogen in a series of three-dimensional plots over 7 of the chambers at 5 different times. These plots illustrate the progress of the flame as it propagates down the channel from an axial position of about 16.5 m to 22 m. The flame position is defined by the temperature plots in Figure 16. Note that burning continues for several chambers behind the flame front. Note also in Figure 17 that the flow ahead of the flame has generated turbulence,
primarily because of the strong shear in the flow behind the obstacles. Recall that the threshold turbulent kinetic energy $k_m$ is 4.74 J/kg. The turbulence levels shown in Figure 17 (peaks of about 10 kJ/kg) far exceed the threshold level. This provides justification for introducing the threshold as a startup mechanism to simply provide a minimum burn velocity.

Because of the choking effect, the flame does not actually accelerate rapidly in the region shown in these figures until DDT occurs. However, the pressures are still rising rapidly in the choked regime. It is important to note the mechanism leading to this pressure rise. As the pressure wave propagates down the channel, it hits the obstacles. Reflection of these shocks off the front faces of the obstacles results in a large increase in pressure and some shock heating (see Figures 15 and 16). In accordance with Equation (27), this increases the burn rate (see Figure 18), which feeds back into an increase in the overall pressure level of the wave.

There is a further mechanism contributing to this feedback in the computation. When the gas burns in the corners in front of the obstacles, the increase in pressure there forces the gas out past the corner of the obstacle and into the central flow. (See the flow pattern in Figure 19.) The resulting shear stresses contribute to a strong increase in the production of turbulence. This, in turn, results in a further increase in the burn rate.

The computation exhibits the deflagration to detonation transition at about 315 ms, when the flame has reached an axial position of 22 m. Note that a sharp pressure ridge has formed all the way across the channel in the uppermost plot in Figure 15; i.e., the pressure spike is not restricted to the region just in front of an obstacle.

As noted above, detonation was not observed experimentally at the concentration of 14.5% $H_2$ used in this calculation, but was observed at 15% $H_2$. It is likely that the actual concentrations at which DDT is or is not experimentally observed would fluctuate from experiment to experiment, so the computational prediction of a transition at 14.5% $H_2$ for the particular modeling parameters used is not regarded as being very conclusive. The fact is that we do not claim to be able to simulate the detailed physical processes which occur in DDT. But it is reasonable to assume that the computed trends in pressure, temperature, and turbulence levels are indicative of real processes, and that what is being calculated is the evolution of a system that is progressing toward a tendency to detonate. (Note that it was assumed in Section III that the pressure was constant through the flame when determining the dependence of the combustion rate on temperature. This is not true for a detonation; however, the propagation velocity for a detonation is determined by the conservation laws and driving forces, and not the combustion rate, so this discrepancy is irrelevant for the numerical detonation, regardless of its validity on other grounds.)

**Results of Variations in the Computational Parameters**

In order to assess the accuracy of the calculation, a second calculation was performed on a refined grid. The baseline computation required about 3.5 hours of computer time on a Cray-1S computer. Because of the expense of these calculations, it was not possible to reduce the grid size enough to make a thorough study of convergence. What was done was to decrease the spacing by a factor of 2/3. (The grid for the complete domain was changed from 13 x 397 to 19 x 595.)
Figure 15. Three-dimensional plots of pressure from the baseline calculation shown in Figures 10–14. Only the region between the 7th through the 14th obstacles along the FLAME facility channel are shown. Coordinates are indicated in centimeters, and absolute pressure is given in atmospheres. The configuration has a symmetry plane at $x = 0$. The times corresponding to each plot appear on the right side of the figure. The axial positions of the pressure transducers which provided the pressures shown in Figures 11–14 are indicated by bullets (●). (Note: In all three-dimensional plots, the values of the dependent variables (pressure, temperature, turbulent kinetic energy, and rate of combustion of hydrogen) have been set equal to zero in the region occupied by the obstacles. This is done simply to make the obstacles easily identifiable; the variables are actually undefined there.)
Figure 16. Three-dimensional plots of temperature (in degrees Kelvin) from the baseline calculation.
Figure 17. Three-dimensional plots of turbulent kinetic energy (in kJ/kg) from the baseline calculation.
Figure 18. Three-dimensional plots of the hydrogen combustion rate (in kg/(m³·s)) from the baseline calculation.
Figure 19. Computed flow pattern between the 9th and 11th obstacles at $t = 310$ ms. The calculation is actually restricted to the domain $x \geq 0$, with a symmetry boundary condition at $x = 0$. However, in this figure the velocity vectors are reflected across the midplane to show the full width of the channel for clarity.
Some of the results are presented in Figures 20-22. The flame trajectory (see Figure 20) is seen to be quite similar to that obtained from the previous calculation, although the flame accelerates faster on the finer grid and detonates sooner. These differences are the result of relatively small errors in the numerical solution on the coarse grid compared to the fine grid. Because of the feedback mechanisms described above, any errors in the flow variables or turbulence parameters become amplified.

To illustrate this sensitivity, an additional simulation was carried out (on the coarse grid) in which the only difference from the baseline calculation was that the rate of combustion during ignition was increased by 31%. The results are given by the dashed line in Figure 20. It is seen that, even though ignition ends before the flame has propagated very far, the gas flow has been given a boost sufficient to generate a higher initial turbulence level and cause considerably greater acceleration of the flame. This example shows how susceptible the progress of such experiments can be to initial conditions. If the burn in the first chamber proceeds differently from shot to shot because of, e.g., different manifestations of fluid instabilities, the end results may vary significantly. For example, if the concentration is marginally detonable, the startup of the burn may determine whether detonation actually occurs.

The pressure histories in Figure 21 and the three-dimensional pressure plot in Figure 22 display the behavior suggested earlier; when numerical dissipation is diminished by lowering the grid size, the shocks and acoustic waves are resolved better and exhibit more structure.

Two more calculations were made to determine the sensitivity of the results to the parameters used in the combustion model. In the first, the values of $A_c, k_m$, and $L_m$ were adjusted so that the slope $dS_T/d\sqrt{k}$ was reduced to 0.82. (See Figure 5.) The initial burn velocity $S_{Ti}$ and the flame thickness $\delta_c$ were changed to 3.6 m/s and 17 cm, respectively. The new values of $A_c, k_m$, and $L_m$ were 18, 15.6 J/kg and 52 cm. For this run, the pressure and temperature exponents $m$ and $n$ were held at 0.2 and 1.64, as before.

For the second calculation, these same new values of $A_c, k_m$, and $L_m$ were used, but the temperature exponent $n$ was raised to 2.0. The pressure exponent $m$ remained at 0.2. Both these calculations were made on the 13 x 397 grid.

The results are compared with those from the original computation in Figures 23-25. Qualitatively, the effects on the flame trajectories are entirely predictable. The flame does not burn as rapidly as before with either of the new sets of parameters, but a more rapid burn is obtained for the higher value of $n$. In both cases, a numerical detonation occurs.

As seen in the sample pressure histories in Figures 24 and 25, the pressure levels are lower with the new parameters, also as expected.

V. CONCLUDING REMARKS

This paper has presented a method for simulating large-scale premixed turbulent flames. The emphasis here has been on developing a way to model the burn rate so that the gas flow driven by the expansion of the burned gases can be computed. A crucial aspect of the application of the model to the computation of the behavior of accelerating flames is that the resulting turbulence intensity and compression of the gas feeds back into the combustion rate.
Figure 20. Comparison of experimental flame trajectories with those obtained from the baseline calculation, from a similar calculation on a refined finite-difference grid, and from a calculation in which the ignition strength was increased by 31% (see text).
Figure 21. Pressure histories at 18.6 m: From the experiment and from the calculation on the refined grid.
Figure 22. Three-dimensional plots of pressure from the calculation on the fine grid. The bullet indicates the location of the pressure transducer corresponding to the results shown in Figure 21.
Figure 23. Comparison of the flame trajectory obtained from the baseline calculation and from the calculations in which the combustion model was altered (see text).
Figure 24. Pressure histories at 18.6 m: From the experiment and from the calculation with reduced combustion rate.
Figure 25. Pressure histories at 18.6 m: From the experiment and from the calculation with reduced combustion rate, but with temperature exponent $n = 2$. 
A second important aspect, pertinent to the simulation of large-scale flames, is that the flame thickness is artificially increased and controlled for satisfactory numerical behavior.

The development of the model has been guided by the desire to use as much general experimental information as possible to determine parameters in the model, while minimizing the amount of knob-twisting to tune to specific experiments. The application to the simulation of experiments in the FLAME facility has been quite successful. The simulated flame trajectories and pressures provide reasonably accurate reproductions of experimental behavior. Some of the computed pressure histories exhibit amplitudes and low frequencies which agree quantitatively with experimental data to within about 25%. It has been shown that computation of pressures which match experiment follows from a good reproduction of the rate of burnout of the individual chambers formed by the obstacles. These rates, in turn, depend on the accuracy of the burn velocities. We have been able to obtain adequate burn velocities by requiring the numerical flames to behave in a way consistent with known properties of turbulent flames and applying flame speed data from the weakly turbulent regime of the experiments.

The resulting combustion model should be of interest to those confronted with the type of simulation addressed here; viz., one in which the size of the computational domain greatly exceeds the flame thickness. In order to apply the method, the following are required: (1) Some experimental and/or theoretical information on turbulent flame speeds and their dependence on the state of the gas, and (2) A computer code which solves the equations of reacting flow (for a one-step reaction), computes the state of turbulence in the gas, and provides for flexibility in the specification of the burn rate and diffusion coefficients. Once a numerical flame with the desired properties has been developed, it can be used for the prediction of flame behavior in a variety of situations.

These computations are not intended to describe deflagration-to-detonation transitions. However, they do graphically demonstrate the way in which feedback mechanisms produce a combustion wave which propagates down the FLAME channel with increasing gas velocities and pressure levels. This suggests that there is a potential for the eventual development of the ability to numerically simulate the evolution of the state of the gas as it approaches a sensitivity to detonation. Given a suitable experimental data base, this might provide a means for predicting DDT.

The combustion model should still be regarded as being under development. In particular, there are two features specific to flame acceleration which need improvement. The first is that a mechanism should be provided that will simulate the sudden burst of acceleration observed in the experiments (the sharp knee in the experimental curve in Figures 10, 20, and 23). The second is the tendency of the numerical flame to exhibit a transition to detonation in the absence of corresponding experimental behavior. As noted earlier, however, this latter problem may not be particularly severe, in view of the demonstrated detonability of mixtures at concentrations close to that studied here.

One way to inhibit numerical DDT would be to include a mechanism for flame quench due to turbulent flame stretch, which is a plausible physical process. Simulating the rapid acceleration while still inhibiting detonation appears less straightforward. It may be that a more accurate treatment of thermal mixing due to turbulence is required, perhaps in concert with compressional heating, but this is merely speculation.
Future work should address these questions. In particular, a joint experimental and computational study of the effects of varying hydrogen concentration would be of interest. Also of interest to the reactor safety community would be the inclusion of gas venting in the simulations. It should be emphasized that the mechanism for the extremely rapid acceleration of the flame fronts which appears in the present work arises to a large extent from the fact that the channel is enclosed except for the opening at one end. Pressure relief due to venting can reduce flame speeds dramatically\(^3\,^5\). However, in some cases, venting may enhance the combustion rate, presumably by introducing additional turbulence\(^3\).

Another direction that future research could take would be to break the computational problem with its large spatial domain up into smaller pieces. Two or three chambers could be considered with appropriately chosen inlet and outlet boundary conditions. This would permit grid refinement to ascertain the convergence properties of the solution. It would also allow better resolution of the turbulent flame fronts, recirculation regions and shock and acoustic waves.
APPENDIX A. INTEGRATION OF PRODUCTION-DISSIPATION TERMS

The following numerical scheme is used to ensure that numerical errors in the production and dissipation terms do not result in nonnegative $k$ or $\epsilon$. First, the production and dissipation terms are split off from the rest of the two equations. This means that in each cell, these terms are allowed to advance $k$ and $\epsilon$ within one computational time step according to

$$\frac{d}{dt}(\rho k) = P - \rho \epsilon \quad (A - 1)$$

$$\frac{d}{dt}(\rho \epsilon) = C_1 \frac{\epsilon}{k} - C_2 \rho \frac{\epsilon^2}{k} \quad (A - 2)$$

where

$$P = \sigma : \nabla \mathbf{u}$$

(Mesh subscripts are dropped for simplicity. Ordinary derivatives are indicated, since the quantities are decoupled from those in other computational cells. The convection and diffusion terms in Equations (10) and (11) are added separately.)

In the strong turbulence regime we have

$$\mu_T \gg \mu_L$$

This is essentially always true in our calculations. Then

$$\mu \approx C_\mu \frac{k^2}{\epsilon}$$

Define

$$Q \equiv \rho k$$

$$E \equiv \rho \epsilon$$

Then

$$\mu = C_\mu \frac{Q^2}{E}$$

and

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\[ P = C_\mu f_p(u,k,\epsilon) \frac{Q^2}{E} \]

where

\[ f_p(u,k,\epsilon) = \frac{\sigma \cdot \nabla u}{\mu} \]

The rationale behind separating out the term \( f_p(u,k,\epsilon) \) is that it will be held constant while equations (A-1) and (A-2) are integrated over one time step. This is justifiable in that \( f_p \) is not a strong function of \( k \) and \( \epsilon \). (In incompressible flow, it has no \( k \) or \( \epsilon \) dependence at all.)

Equations (A-1) and (A-2) can now be written

\[
\frac{dQ}{dt} = \frac{P'Q^2}{E} - E \quad (A - 3)
\]

\[
\frac{dE}{dt} = C_1 \epsilon P'Q - C_2 \epsilon \frac{E^2}{Q} \quad (A - 4)
\]

where

\[ P' = C_\mu f_p(u,k,\epsilon) \]

From (A-3) and (A-4) we obtain

\[
\frac{d}{dt} \left( \frac{Q}{E} \right) = \frac{1}{E} \frac{dQ}{dt} - \frac{Q}{E^2} \frac{dE}{dt}
\]

\[
= - \left( C_1 \epsilon - 1 \right) P' \frac{Q^2}{E^2} + \left( C_2 \epsilon - 1 \right)
\]

Let \( \xi = Q/E \). Then

\[
\frac{d\xi}{dt} = - \left( C_1 \epsilon - 1 \right) P' \xi^2 + \left( C_2 \epsilon - 1 \right)
\]

and we use the following implicit scheme to integrate \( \xi \):

\[
\xi^{n+1} = \frac{\xi^n + (C_2 \epsilon - 1) \Delta t}{1 + (C_1 \epsilon - 1) P' \xi^n \Delta t} \quad (A - 5)
\]

where superscripts \( n \) and \( n + 1 \) denote old and new time levels and \( \Delta t \) is the time step. Since \( C_1 \epsilon, C_2 \epsilon > 1 \), positive \( \xi^n \) insures positive \( \xi^{n+1} \).
From Equation (A-3),

\[ \frac{dQ}{dt} = P' \xi Q - \frac{Q}{\xi} \]

and the scheme

\[ Q^{n+1} = \frac{(1 + P' \Delta t \xi^{n+1}) Q^n}{1 + \Delta t \xi^n} \quad (A - 6) \]

is consistent and results in nonnegative \( Q^{n+1} \). Applying first (A-5), then (A-6), and using

\[ E^{n+1} = \frac{Q^{n+1}}{\xi^{n+1}} \]

permits evaluation of

\[ k^{n+1} = Q^{n+1} / \rho \]

\[ \epsilon^{n+1} = E^{n+1} / \rho \]

and these quantities are positive as long as \( k^n \) and \( \epsilon^n \) are positive.
APPENDIX B. DIFFERENCE FORMULA FOR $\epsilon$ AT WALLS

A function which varies as $1/y$ cannot be differentiated accurately by the usual second-order central difference formulas. Since $\epsilon$ has this behavior near walls, a special differentiation formula was derived for this purpose, as described below. A formula which is exact for $1/y$ is

$$\frac{\partial \epsilon}{\partial y} \approx \frac{3}{4} \left( \frac{\epsilon_{i+\frac{1}{2}} - \epsilon_{i-\frac{1}{2}}}{\Delta y} \right)$$

where $i$ is a grid index. (In Conchas-Spray, $\epsilon$ is a cell-centered quantity, so $\partial \epsilon/\partial y$ is centered on the cell faces.) We have found that such an alteration at only the first grid point in from the wall yields a more accurate solution; however, it is an inconsistent formula for the derivative and should not be carried into the interior of the fluid. The discontinuity in transferring from this formula to the second-order approximation

$$\frac{\partial \epsilon}{\partial y} \approx \frac{\epsilon_{i+\frac{1}{2}} - \epsilon_{i-\frac{1}{2}}}{\Delta y}$$

obviously leads to some inaccuracy, but it appears to be less severe than carrying the second-order approximation all the way to the wall.

Note added in proof: In the most recent version of the Kiva code (a descendant of Conchas-Spray\textsuperscript{17}), Los Alamos researchers are solving the $\epsilon$-equation by advecting the length scale $L$ rather than $\epsilon$. This alleviates the aforementioned difficulties with $\epsilon$ at wall boundaries.
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A new computational model for large-scale premixed flames is developed and applied to the simulation of flame acceleration experiments. The primary objective is to circumvent the necessity for resolving turbulent flame fronts; this is imperative because of the relatively coarse computational grids which must be used in engineering calculations. The essence of the model is to artificially thicken the flame by increasing the appropriate diffusivities and decreasing the combustion rate, but to do this in such a way that the burn velocity varies with pressure, temperature, and turbulence intensity according to prespecified phenomenological characteristics. The model is particularly aimed at implementation in computer codes which simulate compressible flows. To this end, it is applied to the two-dimensional simulation of hydrogen-air flame acceleration experiments in which the flame speeds and gas flow velocities attain or exceed the speed of sound in the gas. It is shown that many of the features of the flame trajectories and pressure histories in the experiments are simulated quite well by the model. Using the comparison of experimental and computational results as a guide, some insight is developed into the processes which occur in such experiments.