Unified Navier-Stokes Flowfield and Performance Analysis of Liquid Rocket Engines

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In an effort to improve the current composite solutions in the design and analysis of liquid propulsive engines, a computational fluid dynamics (CFD) model capable of calculating the reacting flows from the combustion chamber, through the nozzle and the Space Shuttle main engine (SSME) fired at sea level are investigated. The CFD model, FDNS (finite difference Navier-Stokes), is a pressure-based, viscous, ideal gas/real gas, reactive flow code. An equilibrium chemistry algorithm is employed using the point implicit method. A conical nozzle with the same expansion ratio as the SSME nozzle is computed to study the shock formation in both the internal and external flowfields. The bell-shaped SSME nozzle is run at 100% power level at various flow conditions. The computed flow solutions and nozzle thrust performance are in good agreement with those of other standard codes and engine hot fire test data.

Nomenclature

\begin{align*}
A_i & = \text{chemical symbol of the } i\text{th species} \\
C_{pm} & = \text{heat capacity for species } n \\
C_p & = \text{turbulence modeling constant, 0.09} \\
C_v & = \text{element molar concentrations} \\
C_1 & = \text{turbulence modeling constant, 1.15} \\
C_2 & = \text{turbulence modeling constant, 1.9} \\
C_3 & = \text{turbulence modeling constant, 0.25} \\
c & = \text{local speed of sound} \\
c_i & = \text{species molar concentrations} \\
D_i & = \text{dissipation terms} \\
d_\rho & = \text{number of element in a species} \\
F & = \text{convection and diffusion fluxes} \\
G & = \text{geometrical matrices} \\
h & = \text{enthalpy} \\
J & = \text{Jacobian of coordinate transformation} \\
J_n & = \text{diffusion fluxes for species } n \\
K_r & = \text{equilibrium constant for } sth \text{ reaction} \\
k & = \text{turbulent kinetic energy} \\
M & = \text{total number of elements} \\
M_E & = \text{exit Mach number} \\
N & = \text{total number of species} \\
Pr & = \text{turbulent kinetic energy production} \\
P_{TCE} & = \text{total pressure at centerline exit} \\
P_{TWE} & = \text{total pressure at wall exit} \\
p & = \text{static pressure} \\
q & = \text{represents } 1, u, v, h, k, \varepsilon, \text{ and } p_t \\
R & = \text{gas constant} \\
S_q & = \text{local grid cell flow area} \\
T & = \text{static temperature} \\
U & = \text{transformed velocity} \\
\mu, \nu & = \text{mean velocities in } x \text{ and } y \text{ directions} \\
W_n & = \text{mass production rate for species } n \\
x, y & = \text{physical coordinates} \\
\alpha_n & = \text{mass fraction for species } n \\
\beta & = \text{pressure relaxation parameter} \\
\gamma & = \text{specific heat capacity ratio} \\
\varepsilon & = \text{turbulent kinetic energy dissipation rate} \\
\eta_1, \eta_2 & = \text{dissipation parameters} \\
\lambda & = \text{convective dissipation parameter} \\
\mu & = \text{effective viscosity} \\
\mu_t & = \text{turbulent kinetic energy production} \\
\nu & = \text{artificial dissipation parameter} \\
\xi & = \text{computational coordinates} \\
\rho & = \text{density} \\
\sigma_\varepsilon & = \text{turbulence modeling constants} \\
\Phi & = \text{energy dissipation function} \\
\end{align*}

Superscripts

\begin{align*}
a_n, b_n & = \text{stoichiometric coefficients} \\
n & = \text{time level} \\
' & = \text{perturbation} \\
\end{align*}

Introduction

A SERIES of procedures\textsuperscript{1-4} to determine the liquid rocket engine thrust chamber performance and plume flowfield have been developed by governmental and industrial organizations in the mid-sixties. Although improvements to the codes have been made, there are still limitations. The calculation is usually achieved by assembling composite solutions in series. For example, the entire combustion chamber is represented by a point chemical equilibrium analysis, the supersonic flow or the core flow region is solved by the inviscid method of characteristics (MOC) analysis, and the wall and shear layers are described by a separate analysis. The wall layer analysis is restricted to a thin boundary layer, flow separation cannot be predicted, and only axisymmetric flow and steady-state phenomenon can be described.

Accurate numerical prediction of the thrust performance and nozzle/plume flowfield can be achieved with recent ad-
vancements in CFD technology without aforementioned limitations. Transient flow can be simulated, and flow reversal has been predicted. Furthermore, advanced CFD technology has three-dimensional capability with applications to dual throat engines, scarfed, integrated nozzles, and cluster nozzles. In Ref. 7, a density-based code PARCH has been applied to a variety of rocket and scramjet propulsive flowfields, albeit no comparison with measurement was made. Reference 8 has used another density-based code RPLUS to calculate the performance and flow for the thrust chamber of a small scale thruster, while performance data was compared. In this study, a pressure-based CFD model is developed for the first time to calculate a unified liquid rocket engine flowfield which describes the main combustion chamber, nozzle, and plume regions simultaneously. Full-scale hot-firing test data on performance and wall pressure were collected and compared. Code validation was achieved by systematically comparing the computational results of the thrust performance and flowfields to those of various industrial codes and hot fire experiments.

**Governing Equations**

The basic equations employed in this study to describe a unified liquid rocket engine flowfield are the general-coordinate, multicomponent transport equations. The classification of the governing equations changes from one point to another in the unified flowfield. That is, mixed parabolic- hyperbolic type for subsonic chamber flow, and mainly hyperbolic for supersonic nozzle flow. A generalized form of these equations written in curvilinear coordinates is given by

\[
\left( \frac{1}{J} \right) \frac{\partial}{\partial t} \left[ \frac{\partial U_i}{\partial x_j} \right] = -\rho U_i U_j + \sigma \frac{\partial G_{ij}}{\partial x_j} + \left( \frac{1}{J} \right) S_i
\]

where \( J, U_i, \) and \( G_{ij} \) are written as

\[
J = \frac{\partial (\xi, \eta)}{\partial (x, y)}
\]

\[
U_i = \left( \frac{u_i}{\partial \xi} \right) \frac{\partial \xi}{\partial x_j}
\]

\[
G_{ij} = \left( \frac{1}{J} \right) \left( \frac{\partial \xi}{\partial x_j} \right) \left( \frac{\partial \xi}{\partial x_i} \right)
\]

\( \mu = (\mu_t + \mu_k)/\sigma_t \) is the effective viscosity when the turbulent eddy viscosity concept is employed to model the turbulent flows. \( \mu = \rho C_{\mu} k^2/\varepsilon \), is the turbulence eddy viscosity. \( \sigma_t \) and \( S_i \) are given in Table 1. \( \sigma_t = 0.95 \) is the turbulent Prandtl number that is widely used for nozzle flows. An extended two-equation k-\( \varepsilon \) turbulence model is used to describe the turbulence, while \( \sigma_t \) and \( \sigma_t \) are taken from that model closure.

The equation of state for an ideal gas is employed for the closure of the above system of equations. A four-step reversible hydrogen/oxygen equilibrium chemistry model is used to close the chemistry system.

### Numerical Schemes

An adaptive dissipation scheme was employed to approximate the convective terms of the momentum, energy, and continuity equations; the scheme is based on second- and fourth-order central differencing with artificial dissipation. First-order upwind scheme is used for the species and turbulence equations, since the parameters involved must have positive quantities. Different eigenvalues are used for weighting the dissipation terms depending on the conserved quantity being evaluated, in order to give correct diffusion fluxes near wall boundaries. This procedure is different from those proposed in other works \( ^{10-12} \) in which the sum of the absolute value of the convection velocity and the local speed of sound is used to weigh the dissipation terms. Adding the dissipation term to the convective fluxes \( F \) in \( \xi \) produces

\[
\frac{\partial F}{\partial \xi} = \frac{(F_{i+1} - F_{i-1})}{2} - (D_{i+1/2} - D_{i-1/2})
\]

The dissipation terms are constructed such that a fourth-order central and fourth-order damping scheme is activated in smooth regions, and a second-order central and second-order damping scheme is used near shock waves. Since the Jacobian matrices of the Euler fluxes have eigenvalues of \( U, U + c, \) and \( U - c, \) it may be sufficient to use the magnitudes of these eigenvalues to weight the dissipation terms. \( U - c \) is not desirable, however, due to the possibility of \( U \) changing signs. To maintain the smoothness of the solution with improved accuracy, \( |U| + c \) was used for the continuity equation and the minimum damping \( |U| \) was used for other transport equations in this study. General forms of the dissipation terms are given for the continuity equation by

\[
D_{i+1/2} = D_1 (\rho_{i+1} - \rho_i) + D_2 (\rho_{i-1} - 3\rho_i + 3\rho_{i+1} - \rho_{i+2})
\]

and for other transport equations

\[
D_{i+1/2} = D_1 (q_{i+1} - q_i) + D_2 (q_{i-1} - 3q_i + 3q_{i+1} - q_{i+2}) + (1 - \varepsilon_1) (|U_{i+1/2}|/16)(q_{i-1} - q_i) - (q_{i+2} - q_i)
\]

where

\[
D_1 = 0.25 \nu_{i+1/2} (|U| + c)_{i+1/2} \quad D_2 = \max(0, 0.01 - 0.25 \nu_{i+1/2}) (|U| + c)_{i+1/2} \quad D_3 = 0.05 \varepsilon_1 |U|_{i+1/2} \quad D_4 = \varepsilon_2 (1 - \varepsilon_1) \max(0.01 \varepsilon_1, 2|\nu|_{i+1/2})
\]

\[
\varepsilon_1 = \max(\lambda, \min(1.0, 25 \nu_{i+1/2})) \quad \varepsilon_2 = 0.015 \quad \nu_{i+1/2} = \max(|a_i|, \min(1, 10, 25 \nu_{i+1/2}))
\]

In the above formulations, the values of \( a_i \) and \( \nu_{i+1/2} \) approach zeros in smooth regions and reach unitary near high pressure gradients. For the continuity equation, second-order damping dominates when \( \nu_{i+1/2} \) approaches unity and fourth-order damping dominates when \( \nu_{i+1/2} \) approaches zero. This is the basis of the adaptive dissipation scheme. For the energy and momentum equations, and in addition to the adaptive dissipation scheme, a parameter \( \lambda \) can be specified to further direct the upwindness, depending on the physics of the flow. An unity \( \lambda \) corresponds to a full upwind scheme, and a vanishing \( \lambda \) corresponds to a central difference scheme in smooth regions. \( \lambda \) is unity for the turbulence and species equations; this is to prevent any cosmetic blemishes near regions of large gradients for the scalar quantities.

A pressure-based solution method was selected so that a wide range of flow speeds could be analyzed with the same code. Successful results of viscous flow computations using pressure-based methods have been reported. \( ^{13,14} \) For high-

<table>
<thead>
<tr>
<th>( q )</th>
<th>( \sigma_q )</th>
<th>( S_q )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>( u )</td>
<td>1.00</td>
<td>( p_n + \nabla</td>
</tr>
<tr>
<td>1.00</td>
<td>( p_n + \nabla</td>
<td>\mu</td>
</tr>
<tr>
<td>( h )</td>
<td>0.95</td>
<td>( D_p \Delta t + \Phi + \Sigma L_{cp} \Delta T - \Sigma h \Delta W_n )</td>
</tr>
<tr>
<td>( k )</td>
<td>0.89</td>
<td>( \rho (\rho - \varepsilon) )</td>
</tr>
<tr>
<td>( \varepsilon )</td>
<td>1.15</td>
<td>( \rho \varepsilon (C_P - C_P e + C_P \Delta T/\varepsilon) )</td>
</tr>
<tr>
<td>( a_n )</td>
<td>1.00</td>
<td>( \Delta W_n, n = 1, \ldots, N )</td>
</tr>
</tbody>
</table>
speed flow cases, a hyperbolic pressure correction equation was employed by perturbing the density in the mass conservation equation. This provides a smooth transition from low- to high-speed flow characteristics. For time accuracy, a time-centered, time-marching scheme with a multiple pressure correctors algorithm was employed. In general, a non-iterative time-marching scheme was used for time-dependent flow computations; however, subiterations can be used if necessary. The multigrid procedure is described below.

A simplified momentum equation was combined with the continuity equation to form a pressure correction equation. The simplified momentum equation can be written as

$$\frac{\partial p}{\partial t} = -\nabla p'$$

or, in discrete form

$$u'_t = -\beta(\Delta t/\rho)\nabla p'$$

The velocity and density fields in the continuity equation for time level \(n + 1\) are then perturbed to form a correction equation. That is

$$\left[\frac{1}{\nabla l} + \nabla(pu_i)\right]^{n+1} = \delta(p^n + p')$$

$$+ \nabla[(\rho^n + \rho')(u'^n + u'_t)] = 0$$

By neglecting the \(\rho'u'_t\) terms, the following equation results:

$$\frac{\partial p'}{\partial t} + \nabla(pu_i'\rho) + \nabla(pu_i) = -\left[\frac{\partial p}{\partial t} + \nabla(pu_i)\right]^n$$

Substituting Eq. (1) into Eq. (2) and letting \(\rho' = p'/RT\), the following pressure correction equation is obtained:

$$\frac{\partial (p'/RT)}{\partial t} + \nabla[(u_i/RT)p'] - \nabla(\beta \Delta T \nabla p') = -\left[\frac{\partial p}{\partial t} + \nabla(pu_i)\right]^n$$

To provide smooth shock solutions the adaptive dissipation terms described above are added to the right side of Eq. (3). Once Eq. (3) is satisfied, the velocity field and the pressure field are updated through Eq. (1) and the following relation:

$$p^{n+1} = p^n + p'$$

The density field is then updated by applying the equation of state. To ensure that the updated velocity, density, and pressure fields satisfy the continuity equation, the above pressure correction solution procedure is repeated several times before marching to the next time step. This represents a multigrid corrector solution procedure.

The chemistry source terms were evaluated with a point implicit procedure before the species equations were solved. The equilibrium chemistry source terms were based on the CHMQGM algorithm.\(^{15}\)

**Equilibrium Chemistry Algorithm**

The chemistry source terms must be computed for each grid point at each time step. The point implicit procedure allows the equilibrium calculation for each grid point to be independent from that of its neighbors and may be treated in isolation. Since kinetics loss was estimated to be negligible in the SSME thrust chamber and nozzle,\(^{16}\) equilibrium is assumed to exist for the entire computational domain, including the exhaust plume region.

In general, if there are \(N\) distinct chemical species composed of \(M\) chemical elements, then the algebraic system to be solved consists of \(N-M\) nonlinear equilibrium equations and \(M\) linear element-conservation relations. The equilibrium reactions are of the form

$$\sum_{i=1}^{N} a_i A_i = \sum_{i=1}^{N} b_i A_i \quad s = 1, \ldots, N-M$$

The equilibrium reactions lead to algebraic relations of the form

$$K_s [c_i]^{m_s} = \sum_{i=1}^{N} c_i^{p_i} \quad s = 1, \ldots, N-M$$

The linear element-conservation reactions are of the form

$$\sum_{i=1}^{N} d_{mn} c_i = c_m \quad m = 1, \ldots, M$$

CHMQGM algorithm has shown that a reduced system can be obtained by substituting the \(M\) linear relations, Eq. (5), into the \(N-M\) nonlinear relations, Eq. (4). As a result, an iterative Newton-Raphson technique was used to solve a system of \(N-2M\) equations rather than the larger equivalent \(N \times N\) system. The equilibrium constant approach of CHMQGM is not as general as a minimization of Gibb's free energy approach\(^{17}\) for an arbitrarily chosen chemical system. However, for a generally well-known hydrogen/oxygen system such as the one being used in SSME, the equilibrium constant approach appears to be more efficient.

**Hydrogen/Oxygen Equilibrium Chemistry**

The hydrogen/oxygen equilibrium reactions used in this study are a subset reduced from the hydrocarbon combustion partial equilibrium system.\(^{18}\) It consists of six species (Table 2) and four reactions [Eqs. (6–9) as shown in Table 3].

The element-conservation equations are

$$C_0 = c_1 + c_3 + c_4 + 2c_6$$

$$C_0 = 2c_1 + c_3 + c_4 + 2c_5$$

Substitute Eqs. (6) and (7) into Eqs. (8) and (9), we have

$$c_6 = \frac{R_1}{c_1^2}$$

$$c_5 = \frac{R_1}{c_1^2}$$

$$c_4 = \frac{R_1}{c_1^2}$$

$$c_3 = \frac{R_1}{c_1^2}$$

The equilibrium reactions are

$$K_{1} c_1 c_2 = c_3$$

$$K_{2} c_1 c_3 = c_3$$

$$K_{3} c_1 c_3 = c_3$$

$$K_{4} c_1 c_3 = c_3$$

**Table 2** Species and reactions considered

<table>
<thead>
<tr>
<th>Subscript</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>H_2O Water vapor</td>
</tr>
<tr>
<td>2</td>
<td>H Atomic hydrogen</td>
</tr>
<tr>
<td>3</td>
<td>O Atomic oxygen</td>
</tr>
<tr>
<td>4</td>
<td>OH Hydroxyl radical</td>
</tr>
<tr>
<td>5</td>
<td>H_2Molecular hydrogen</td>
</tr>
<tr>
<td>6</td>
<td>O_2 Molecular oxygen</td>
</tr>
</tbody>
</table>

**Table 3** Reactions and equilibrium equations

<table>
<thead>
<tr>
<th>Reactions</th>
<th>Equilibrium equations</th>
</tr>
</thead>
<tbody>
<tr>
<td>O_2 = 2O</td>
<td>(K_{1} c_1 c_2 = c_3)</td>
</tr>
<tr>
<td>H_2 + 2H_2O = 2H + O_2</td>
<td>(K_{2} c_1 c_3 = c_3)</td>
</tr>
<tr>
<td>2O + 2H_2O = 2H_2 + O_2</td>
<td>(K_{3} c_1 c_3 = c_3)</td>
</tr>
<tr>
<td>2H_2O = 2H_2 + O_2</td>
<td>(K_{4} c_1 c_3 = c_3)</td>
</tr>
</tbody>
</table>

\(^{15}\) CHMQGM: CH quasigeminal method.

\(^{16}\) Exclusion of the exhaust plume is justified for a large area engine, since the exhaust area is a very small part of the entire computational domain.

\(^{17}\) Free energy approach: The minimization of Gibb's free energy is a thermodynamic approach to determine the equilibrium state of a chemical system.

\(^{18}\) Hydrocarbon combustion: The combustion of hydrocarbons involves the reactions of hydrocarbon molecules with oxygen to form carbon dioxide, water, and possibly other products.
Substitute the linear term in Eqs. (12–15) into the linear element-conservation Eqs. (10) and (11), yielding two nonlinear equations and three unknowns in \( c_2, c_3, \) and \( c_5 \):

\[
C_0 = R_1 c_2 c_3 + c_3 + R_2 c_3^2 + 2R_1 (c_4)^2 \tag{16}
\]

\[
C_H = 2R_1 c_2 c_3 + c_2 + R_2 c_3^2 + 2R_2 (c_5)^2 \tag{17}
\]

The final reduced equations can then be obtained by substituting Eq. (16) into Eq. (17), and to eliminate \( c_5 \) with Eq. (13), we have

\[
R_1 R_4 (c_5)^2 c_3 + R_2 c_3^3 + 2R_1 (c_4)^2 + c_3 - C_0 = 0
\]

\[
R_2 c_3 + 4R_1 (c_4)^2 + 2c_4 - 2R_2 (c_5)^2 - c_4
\]

\[
+ (C_H - 2C_0) = 0
\]

This reduced system consists of one cubic equation and one quadratic equation with two unknowns, and can readily be solved by Newton-Raphson's iterative method.

**Boundary Conditions**

Fixed chamber total conditions were used at the inlet of the combustion chamber. Subsonic boundary conditions were used at the inlet of the external ambient air and the pressure was extrapolated. This is to allow the transient disturbances sent from downstream to permeate through the boundary. Flow properties at the wall, centerline and exit were extrapolated from those of the interior domain. To obtain a unique solution for the sea level SSME hot firing sample case, a fixed pressure was applied to the outermost point of the ambient exit boundary.

**Sample Cases for SSME at 100% Power Level**

SSME operating conditions at 100% power level were used for the calculations. The actual values used are shown in Table 4.

**Results of the Inviscid, Ideal Gas, Adiabatic Wall Calculation**

FDNS calculations were made for the inviscid, ideal gas and adiabatic wall conditions in order to compare to the equivalent MOC solutions. The domain of computation was started from the combustor injector faceplate, through the throat, and ended at the nozzle exit plane. Computational results obtained from running the PARC code\(^6\) and RAMP code\(^2\) were also used for comparison. Cases for different specific heat capacity ratios (\( \gamma = 1.14, 1.1875, \) and \( 1.25 \)) were performed. Figure 1 shows the comparison of vector and Mach number contour for FDNS and PARC solutions at \( \gamma = 1.1875 \).

The vectors and Mach numbers show a transition from subsonic flow in the main combustor to sonic flow at the throat, and to supersonic flow in the nozzle. While FDNS is a pressure-based method, PARC is a density-based procedure. Both codes have captured the nozzle shock stemming downstream from the throat, and exhibited similar Mach number contours throughout the computational domain. A sonic line can be observed near the throat region. The computations were made on the same grid with a grid size of \( 111 \times 65 \).

Figures 2 and 3 show the comparison of centerline and wall pressure distributions for \( \gamma = 1.1875 \) and \( \gamma = 1.25 \), respectively. The FDNS predictions compare well with those of the MOC and PARC results. The sonic start lines obtained from FDNS calculations were used for MOC calculations. The computational time for a typical FDNS ideal gas calculation was estimated as \( 1.03E^{-4} \) CPU s/grid/step on a NASA/MSFC CRAY-XMP. Five hundred iterations were required for near convergence.

Performance calculations of the present method gave specific impulses (ISP) of 426.6, 452.5, and 472.3 for \( \gamma = 1.25, 1.1875, \) and 1.14, respectively. Real gas calculations indicated that the heat capacity ratio ranged from 1.14 at the combustor inlet, to about 1.25 at the nozzle exit. The thermodynamic properties of the combusting gases are very important parameters:

**Table 4: Operating parameters**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixture ratio</td>
<td>6.000</td>
</tr>
<tr>
<td>O(_2) inlet temp.</td>
<td>90.56</td>
</tr>
<tr>
<td>H(_2) inlet temp.</td>
<td>20.56</td>
</tr>
<tr>
<td>Chamber pressure, psia</td>
<td>2925.7</td>
</tr>
<tr>
<td>Chamber temperature, K</td>
<td>3639.0</td>
</tr>
<tr>
<td>Geometric area ratio</td>
<td>77.5</td>
</tr>
</tbody>
</table>
eters to the accurate prediction of the liquid rocket engine flowfields and performance. A real gas thermodynamics calculation is therefore in order.

Results of the Inviscid, Real Gas, Adiabatic Wall Calculation

The computational domain was extended to include the exhaust plume and the ambient air for the low altitude FDNS real gas calculations. Seven species were considered, including $\text{H}_2$, $\text{O}_2$, $\text{H}_2\text{O}$, $\text{O}$, $\text{H}$, $\text{OH}$, and $\text{N}_2$. Chemical equilibrium was assumed to be valid for the entire flowfield. Figure 4 shows the comparison of FDNS and MOC predicted centerline and wall Mach numbers. MOC used an equilibrium properties table obtained from Chemical Equilibrium Calculations (CEC).

The FDNS predicted centerline Mach numbers agreed well with those of the MOC predictions. The FDNS prediction of wall Mach number matched the MOC calculation from approximately $x = 5.5$ ft to the exit plane, but underprediction was revealed near the throat. The underprediction may have been caused by two reasons.

1) A constant perpendicular sonic start line ($\text{Mach number} = 1.01$) was assumed for the MOC calculation that was different from the FDNS calculated sonic line.

2) The artificial dissipation used to capture shock by FDNS may have generated excessive total pressure loss, near the starting point of the nozzle shock at the wall.

Nevertheless, the Mach number comparisons for the FDNS and MOC were reasonably good. Figure 5 shows comparisons of the centerline and wall pressure distributions. The FDNS predicted centerline and wall pressure distributions agreed well with those of the MOC predictions. RAMP code overpredicted the centerline pressure slightly near the exit plane.

The FDNS prediction of the SSME nozzle specific impulse is compared to that of MOC and TDK predictions in Table 5. TDK calculation used an eight-reaction finite rate kinetics mechanism. The closeness of all three ISP values indicated that the kinetics loss and shock-kinetics interaction is probably negligible under these circumstances. The computational time for a typical FDNS real gas calculation was estimated to be $2.36\times10^{-4}$ CPU s/grid/step. Approximately 2500 iterations were required for near convergence. The grid size used in this calculation was $201 \times 81$.

Results of the Viscous, Real Gas, and Cooled Wall Calculation

The simulation of 100% power level SSME fired at sea level, was completed by adding the viscous flow and a specified (measured) wall temperature distribution to the calculation. The extended k-ε turbulence model and wall function approach were used to calculate the viscous flow. Figure 6 shows the comparison of predicted nozzle wall pressure distribution with that of the NASA Marshall Space Flight Center Technology Test Bed (TTB) hot fire test data (test numbers 021, 023, and 025). The three TTB pressure taps were located near the end of the nozzle. The prediction agreed well with the measurement, with a maximum difference of less than one PSI. The error bound for the test data ranged from 10 to 15% for these taps among test firings.

Figure 7 shows the Mach number contour for a calculated conical nozzle flow. The overall features of the nozzle and plume such as nozzle shock, Mach disc, lip shock, plume slp
stream, and Mach disc slip stream were captured quite well. The operating conditions were identical to that of the bell-shaped SSME nozzle as indicated in Table 4, including the nozzle length and the area ratio. This was to see if the proposed method can capture a normal shock for a nearly one-dimensional nozzle flow. From Fig. 7, a nozzle shock formed after the inflection point, hit the centerline, and reflected inside the nozzle; a lip shock formed at the nozzle exit and extended to the triple point in the plume region, where a Mach disc formed beneath the triple point and perpendicular to the axis.

The computed Mach number contour of the bell-shaped SSME nozzle flowfield is shown in Fig. 8. Unlike the nearly one-dimensional conical nozzle flow, a distinctive two-dimensional flow behavior is observed. A slightly curved Mach disc developed in the plume region, the chamber total temperature was recovered behind the Mach disc. The disc location was estimated to be 1.1 nozzle exit radii from the exit plane, with a width of about 1.1 nozzle exit radii. The location and width of the computed normal disc agreed well with those of a hot fire test photograph. Comparisons of the thrust performance of the experimental data, the results of this study, and a TDK/BLM1 calculated result are shown in Table 6. BLM is a boundary-layer code that provides TDK with the boundary-layer solution. The FDNS calculated ISP value compared very well with that of the experimental data and the TDK/BLM calculation.

The typical grid size used in this calculation was similar to that of the inviscid case, i.e., 201 \times 81. The computational time used for one typical run was 2.77E-4 CPU s/grid/step. Approximately 3000 iterations were required for near convergence. The CPU time for a nonreacting case was 2.32E-4 s/grid/step. That means only about 20% increase for an equilibrium chemistry calculation.

**Grid Dependency Study**

Extensive grid dependency studies were carried out in order to find the best possible computational grids for the liquid rocket engine flowfield and performance calculations. For instance, an inviscid Mach 4 source flow calculation was performed on several grid systems. This grid study was designed to find out general grid clustering strategy for FDNS to preserve total pressure in the absence of a nozzle shock. The velocity vector and pressure contour for the one-dimensional source flow is shown in Fig. 9. A theoretical exit Mach number of 4 and total pressure of 100,000 should be obtained for a grid-independent CFD solution. Table 7 indicates that good solutions can be obtained with grid clustering near the flow inlet.
A pressure-based reactive Navier-Stokes CFD model FDNS has been developed to analyze a unified SSME liquid rocket engine flowfield. The flow domain included the combustion chamber, nozzle, exhaust plume, and the ambient. The code was validated through several systematic stages. For example, the FDNS predicted inviscid nozzle flowfields were compared to those of the MOC, PARC, and RAMP solutions. The comparisons showed that the FDNS predictions were reasonably accurate for inviscid ideal gas and real gas cases. The viscous, real gas, and specified wall temperature case also shows excellent comparison in terms of nozzle wall pressure, Mach disc location, and width to those of the hot fire test. Furthermore, the FDNS predictions of the SSME ISP values were in excellent agreement with those of the MOC and TDK predictions, and the Rocketdyne experimental data. This study demonstrated that the FDNS code predicts the thrust performance and the nozzle/plume flowfield accurately, and it
can be used as a design and analysis tool for general liquid rocket engines.

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References