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A Theoretical Evaluation of Secondary Atomization Effects on Engine Performance for Aluminum Gel Propellants
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Abstract
A one-dimensional model of a gel-fueled rocket combustion chamber has been developed. This model includes the processes of liquid hydrocarbon burnout, secondary atomization, aluminum ignition, and aluminum combustion. Also included is a model of radiative heat transfer from the solid combustion products to the chamber walls. Calculations indicate that only modest secondary atomization is required to significantly reduce propellant burnout distances, aluminum oxide residual size and radiation heat losses. Radiation losses equal to approximately 2-13% of the energy released during combustion were estimated. A two-dimensional, two-phase nozzle code was employed to estimate radiation and nozzle two-phase flow effects on overall engine performance. Radiation losses yielded a 1% decrease in engine $I_{sp}$. Results also indicate that secondary atomization may have less effect on two-phase losses than it does on propellant burnout distance and no effect if oxide particle coagulation and shear induced droplet breakup govern oxide particle size. Engine $I_{sp}$ was found to decrease from 337.4 to 293.7 seconds as gel aluminum mass loading was varied from 0-70 wt%. Engine $I_{sp}$ efficiencies, accounting for radiation and two-phase flow effects, on the order of 0.946 were calculated for a 60 wt% gel, assuming a fragmentation ratio of 5.

Nomenclature

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
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<tbody>
<tr>
<td>$a$</td>
<td>absorption coefficient</td>
</tr>
<tr>
<td>$A$</td>
<td>area</td>
</tr>
<tr>
<td>$c_p$</td>
<td>specific heat</td>
</tr>
<tr>
<td>$C_D$</td>
<td>drag coefficient</td>
</tr>
<tr>
<td>$d$</td>
<td>diameter</td>
</tr>
<tr>
<td>$F_{n}(T)$</td>
<td>fraction of total radiation emitted by a blackbody at temperature $T$ in the wavelength interval $n$</td>
</tr>
<tr>
<td>$h$</td>
<td>average convective heat transfer coefficient</td>
</tr>
<tr>
<td>$i$</td>
<td>specific enthalpy</td>
</tr>
<tr>
<td>$I$</td>
<td>radiant intensity</td>
</tr>
<tr>
<td>$I_0$, $I_1$</td>
<td>modified Bessel functions</td>
</tr>
<tr>
<td>$j$</td>
<td>droplet size class</td>
</tr>
<tr>
<td>$k$</td>
<td>thermal conductivity</td>
</tr>
<tr>
<td>$L_*$</td>
<td>number of wavelength intervals</td>
</tr>
<tr>
<td>$m$</td>
<td>mass</td>
</tr>
<tr>
<td>$m_*$</td>
<td>droplet mass flux or mass flux through the combustion chamber</td>
</tr>
<tr>
<td>$M_*$</td>
<td>number of droplet size classes</td>
</tr>
<tr>
<td>$n$</td>
<td>wavelength interval</td>
</tr>
<tr>
<td>$N/\tau$</td>
<td>number of droplets passing through combustor control volume per unit time</td>
</tr>
<tr>
<td>$r_s$</td>
<td>droplet/particle surface radius</td>
</tr>
<tr>
<td>$R$</td>
<td>chamber radius</td>
</tr>
<tr>
<td>$R_*$</td>
<td>chamber radius in terms of optical thickness</td>
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<tr>
<td>$q_r$</td>
<td>radiation heat flux from condensed combustion products to wall</td>
</tr>
<tr>
<td>$t$</td>
<td>time</td>
</tr>
<tr>
<td>$T$</td>
<td>temperature</td>
</tr>
<tr>
<td>$u_p$</td>
<td>particle velocity</td>
</tr>
<tr>
<td>$x$</td>
<td>axial location in the combustor model</td>
</tr>
<tr>
<td>$\alpha_1$, $\alpha_2$</td>
<td>control switches</td>
</tr>
<tr>
<td>$\beta$</td>
<td>fragmentation ratio (number of secondary droplets produced per initial droplet)</td>
</tr>
<tr>
<td>$\phi$</td>
<td>angular coordinate</td>
</tr>
<tr>
<td>$\eta$</td>
<td>fraction of aluminum droplet mass that forms as $\text{Al}_2\text{O}_3$ residual on droplet surface</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>wavelength</td>
</tr>
<tr>
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<tr>
<td>$\sigma$</td>
<td>Stefan-Boltzmann constant</td>
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<tr>
<td>$\sigma_s$</td>
<td>scattering coefficient</td>
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<td>$\theta$</td>
<td>angular coordinate</td>
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<tr>
<td>$\Omega_o$</td>
<td>scattering albedo</td>
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Subscripts

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
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<tbody>
<tr>
<td>$\text{Al}$</td>
<td>aluminum</td>
</tr>
<tr>
<td>$b$</td>
<td>boiling, propellant burnout, or blackbody</td>
</tr>
<tr>
<td>$f$</td>
<td>aluminum oxide fume particles or aluminum flame</td>
</tr>
<tr>
<td>$\text{fg}$</td>
<td>liquid-gas</td>
</tr>
<tr>
<td>$g$</td>
<td>gas</td>
</tr>
<tr>
<td>$\text{LH}$</td>
<td>liquid hydrocarbon</td>
</tr>
<tr>
<td>$o$</td>
<td>initial</td>
</tr>
<tr>
<td>$\text{Ox}$</td>
<td>Aluminum oxide residual</td>
</tr>
<tr>
<td>$p$</td>
<td>particle/droplet</td>
</tr>
<tr>
<td>$w$</td>
<td>wall</td>
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Introduction
Gel propellants, in which fine particles are suspended in a gelled combustible liquid, offer potential performance and/or safety advantages over conventional liquid and
solid propellants in rocket applications. Theoretical performance evaluations show that gel propellants may provide increases in specific impulse \( I_{sp} \) and/or propellant density over conventional liquid propellants. A number of potential liquid-solid constituent combinations were theoretically evaluated over a range of mixture ratios and solid constituent loadings for use in a LEO-GEO mission with fixed propellant tankage volume and vehicle dry-mass with no limit on initial gross mass. Based on propellant toxicity and vehicle payload considerations, aluminum was predicted to be the best candidate solid constituent.

More detailed analyses of earth-to-LEO, LEO-to-GEO, LEO-to-Moon, LEO-to-Mars, and several planetary fly-by missions, incorporating appropriate total vehicle mass/volume limitations and propellant density effects on vehicle dry-mass, also were performed to compare various aluminum gels with their liquid bi-propellant counterparts. These studies indicate that gel propellants are likely to provide the greatest benefit on earth-to-LEO, and high-energy planetary missions with NTO/MMH/Al gels also providing significant performance gains over NTO/MMH in LEO-to-GEO and LEO-to-Moon missions. For example, replacing the space shuttle solid boosters with an RP-1/O\(_2\)/Al or NTO/MMH/Al propellant combination, while maintaining the same booster dimensions, could theoretically increase maximum payload by 14-35% over the current shuttle payload. Similarly, the addition of aluminum to nitrogen tetraoxide/monomethyl hydrazine (NTO/MMH) may permit planetary missions that would be impossible with neat NTO/MMH.

In the area of propellant rheology, recent gains have been made in the development of stable gels that are easily sprayed, but more work in this area is still required. In related work, testing has demonstrated that expulsion pumping is feasible in a small engine system and that gel-fueled engines can be throttled over a range of operating conditions, shut-down, and restarted as desired. In addition, experimental results indicate that using aluminum can reduce combustion instabilities associated with liquid hydrocarbon propellants.

Although the above investigations are promising, it should be noted that the theoretical performance studies do not incorporate propellant combustion such as increased propellant combustion times, radiation heat transfer from condensed combustion products to the chamber walls, and nozzle two-phase flow losses due to these same condensed products. Because efficiency losses of only 1.5-4% may be sufficient to eliminate the payload benefits of using aluminum gels, minimization of aluminum combustion related losses is desirable. In addition, these performance losses must be determined before the performance of gel propellants can be accurately evaluated.

Since previous work has shown that the individual aluminum particles in a gel droplet can form an agglomerate that burns as a single aluminum droplet, it is apparent that small gel droplets are beneficial in reducing performance losses. Fine atomization of gel propellants, however, can be difficult to achieve due to their viscous nature. Fortunately, secondary atomization, in which a droplet spontaneously shatters into a number of smaller droplets due to internal vaporization of the liquid carrier, may produce the desired small droplets.

Although secondary atomization has been proposed as a means of reducing gel-related performance losses, previous research has focused on the secondary atomization mechanism itself rather than its effects on engine performance. The focus of our present investigation, therefore, is to theoretically examine the effects of secondary atomization on engine performance losses and, incorporating these performance loss estimates, evaluate the performance potential of gel-fueled engines. To accomplish these objectives, a one-dimensional combustor model has been derived to evaluate secondary atomization effects on propellant burnout distances and radiation heat losses in a rocket combustion chamber. This combustor code is used in conjunction with the Solid Propellant Rocket Motor Performance Prediction Computer Program (SPP), a two-dimensional, two-phase rocket code, to provide a preliminary evaluation of nozzle two-phase flow losses and overall engine performance.

**Combustion Chamber Model**

**One-Dimensional Model Description**

In brief, a radially uniform spray, consisting of four droplet size classes, enters the combustion chamber and burns in a process incorporating liquid carrier burnout, droplet secondary atomization, aluminum agglomerate heat-up and combustion, two-phase particle flow, and radiation heat transfer from solid combustion products to the chamber walls. The propellants used in the combustor model are a JP-10/Al gel and a preheated gaseous O\(_2\) oxidizer. JP-10, a pure hydrocarbon (C\(_{10}H_{16}\)), was chosen as the gel hydrocarbon component instead of RP-1 to avoid the complexity of modeling multi-component droplet combustion. The combustor flow is modeled using a single product-phase containing both gaseous and small Al\(_2\)O\(_3\) fume particles, and three additional flow phases for each droplet size class: a liquid hydrocarbon phase, an aluminum phase, and a phase containing large Al\(_2\)O\(_3\) residuals. The Al\(_2\)O\(_3\) must be separated into two flow phases because of the two oxidation mechanisms inherent in aluminum combustion. In the first oxidation mechanism, large Al\(_2\)O\(_3\) residuals are formed through droplet surface condensation/oxidation, and in the second, very small Al\(_2\)O\(_3\) fume particles are produced through vapor-phase oxidation. Since the Al\(_2\)O\(_3\) residuals are much larger than the fume particles and are attached to the surface of the aluminum droplets it is necessary to model the Al\(_2\)O\(_3\) residuals separately from the fume particles. Including the fume particles in the gas-phase flow requires the assumptions of no temperature or velocity slip between the fume particles and the gases, which are reasonably valid, and greatly simplifies the problem solution.

As an Al droplet burns, it continually produces Al\(_2\)O\(_3\) fume particles at the oxide boiling point which subse-
quently equilibrate with the gas-phase temperature and velocity. This process results in a range of fume particle temperatures and velocities as new particles are created and older particles continue to equilibrate with the gases. If the temperature and velocity slip were not neglected, many additional flow phases would be required to accurately model this range of fume particle temperatures and velocities.

Mass Conservation: Using the definition of velocity, \( u = \frac{dx}{dt} \), to relate the differential variables, \( dx \) and \( dt \), the following equation can be written for system mass conservation for \( M \) droplet size classes:

\[
\frac{dm}{dx} = -\sum_{j=1}^{M} \left[ \frac{N}{\tau} \left( \frac{dm_{\text{LH}}}{dt} + \frac{dm_{\text{Al}}}{dt} + \frac{dm_{\text{Ok}}}{dt} \right) \right]
\]

(1)

where it is assumed that no mass is added to or removed from the chamber except at the injector face and the chamber exit, and that steady-state operation prevails.

The three time derivatives on the right hand side of Eq. 1 are found from hydrocarbon droplet gasification and aluminum droplet combustion models which are discussed later in this paper. The value of \( N/\tau \) for each droplet size class can be determined from the total initial gel mass flux and a normalized droplet size distribution.

Energy Conservation: The steady-state energy balance for the system, including radiation heat losses, can be expressed as

\[
\frac{d(m_i)}{dx} = -\sum_{j=1}^{M} \left[ \frac{d(m_i)_{\text{LH}}}{dx} + \frac{d(m_i)_{\text{Al}}}{dx} + \frac{d(m_i)_{\text{Ok}}}{dx} \right] - 2\pi Rq'
\]

(2)

Since gas flow optical properties are dominated by the small \( \text{Al}_2\text{O}_3 \) fume particles, the radiation term is independent of the individual droplet size classes and is not included in the summation term. Each of the terms on the right hand side of Eq. 2 can be represented as follows by expanding the derivatives and assuming that the only changes in specific enthalpy are due to convective heat transfer with the gaseous product-phase:

\[
\frac{d(m_i)}{dx} = \left[ \frac{N}{\tau} \left( \alpha_1 \frac{A_k h(T_e - T_p)}{m_p} + \alpha_2 \frac{dm_p}{dt} \right) \right]
\]

(3)

The coefficients \( \alpha_1 \) and \( \alpha_2 \), equal to 0 or 1, are used to provide the correct terms, depending on whether the liquid hydrocarbon, aluminum, or \( \text{Al}_2\text{O}_3 \) mass flux is being examined. In the hydrocarbon vaporization model, the bulk droplet temperature is assumed to remain at the initial droplet temperature while only a thin surface layer of hydrocarbon is heated to the hydrocarbon boiling temperature before being vaporized. Since the liquid hydrocarbon enthalpy does not vary with chamber axial location, \( \alpha_1=0 \) and \( \alpha_2=1 \). In this case, the enthalpy, \( i \), equals the initial liquid hydrocarbon specific enthalpy, and \( dm_p/dt \) is the hydrocarbon mass vaporization rate from a single droplet.

Following hydrocarbon burnout, an agglomerate of aluminum particles remains. The aluminum agglomerate temperature rises from the hydrocarbon boiling temperature, through the aluminum melting point, to the aluminum ignition temperature as heat is transferred from the gas flow to the agglomerate, at which point aluminum combustion commences. In the combustor model, this agglomerate ignition/heat-up process is approximated as convective heat transfer from the gas flow to a spherical, uniform-temperature agglomerate. Heat transfer to the agglomerate causes the agglomerate temperature to rise from the hydrocarbon boiling temperature to the aluminum melting temperature. The agglomerate temperature is then held constant until sufficient energy to melt the entire agglomerate has been transferred from the gas-flow, at which point it is assumed that aluminum combustion begins. Heat-up of the molten aluminum droplet to the aluminum boiling temperature is accounted for in the aluminum combustion model, as discussed below. Since it is assumed that no aluminum vaporization occurs during agglomerate heat-up and melting, and because the specific enthalpy does not change, \( \alpha_1=1 \) and \( \alpha_2=0 \) during these stages of the aluminuin ignition/combustion process. Once the agglomerate melts, however, the aluminum specific enthalpy remains constant for the same reasons as discussed for the liquid hydrocarbon, and combustion begins to occur. During this portion of the ignition/combustion process, \( \alpha_1=0 \), \( \alpha_2=1 \), and the specific enthalpy, \( i \), equals that of liquid aluminum at its melting temperature.

Since the \( \text{Al}_2\text{O}_3 \) residual forms on the aluminum droplet surface, the \( \text{Al}_2\text{O}_3 \) residual is maintained at the aluminum boiling temperature, and therefore maintains a constant specific enthalpy, as long as any aluminum remains in the droplet. For the time period prior to aluminum burnout, \( \alpha_1=0 \), \( \alpha_2=1 \), and the specific enthalpy is that of \( \text{Al}_2\text{O}_3 \) at the aluminum boiling temperature. Once aluminum combustion and \( \text{Al}_2\text{O}_3 \) formation ceases, \( \alpha_1=1 \), \( \alpha_2=0 \), and the particle temperature is allowed to equilibrate with the gas temperature. The total system energy balance is represented by the substitution of Eq. 3 with appropriate values of \( \alpha_1 \), \( \alpha_2 \), and \( i \) into Eq. 2.

Radiation Heat Transfer: Radiation from the solid combustion products to the chamber walls is a participating media phenomenon, requiring a solution of the radiative transfer equation. Expressed in cylindrical coordinates this equation is
\[
\sin \phi \left[ \cos \phi \frac{\partial}{\partial \phi} - \sin \phi \frac{\partial}{\partial r} \right] I + l = (1 - \Omega_s)I_{b,w}(r) + \frac{\Omega_s}{2\pi} \int I \sin \theta \, d\theta \, d\phi
\]

where \( l = l(r, \theta, \phi) \). It should be noted that the radial position, \( r \), is based on optical thickness and is not a physical position (i.e., \( r = \int (a + \sigma_s) \, dr \)). The scattering albedo, \( \Omega_s \), is defined as

\[
\Omega_s = \frac{\sigma_s}{a + \sigma_s}
\]

where \( a \) and \( \sigma_s \) are determined from Mie theory. In the Mie scattering calculations the fume particles are assumed to be uniform in size (1 \( \mu m \)) and the refractive indices are those of Al\(_2\)O\(_3\) smoke.\(^{23}\)

Since the flow is likely to be optically thick,\(^{23,25}\) a cylindrical \( P_1 \) diffusion approximation is used to simplify the radiative transfer equation. By assuming that \( I_{b,w} \) is independent of radial location, the following equation for radiative heat flux is obtained:\(^{25}\)

\[
q'(R) = \frac{4\pi l_1 \left( \frac{R_s}{\xi} \right) (I_{b,w} - I_{b,.w})}{3\xi I_0 \left( \frac{R_s}{\xi} \right) + 2l_1 \left( \frac{R_s}{\xi} \right)}
\]

The variable \( \xi \) is based on the flow optical properties and is defined as

\[
\xi = \frac{1}{\sqrt{3(1 - \Omega_s)}}
\]

If \( \xi(\lambda) \) is assumed to be constant over a wavelength interval, Eq. 6 can be expressed as the following summation:

\[
q'(R) = \sum_{n=1}^{\infty} \frac{4l_1 \left( \frac{R_s}{\xi} \right)_{n}}{3\xi I_0 \left( \frac{R_s}{\xi} \right)_{n} + 2l_1 \left( \frac{R_s}{\xi} \right)_{n}} \left[ F_n(T_s) \sigma T_s^4 - F_n(T_w) \sigma T_w^4 \right]
\]

**Momentum Conservation:** The gas-phase momentum equation in the combustion chamber is trivial assuming a negligible chamber pressure gradient and no body forces. Neglecting virtual mass and Bassett forces, the particle momentum equation for a given droplet size class can be expressed as

\[
\frac{du_z}{dx} = -\frac{3 \rho \gamma C_p \left( u_z - u_g \right) u_z}{4 \gamma_p d_r u_p}
\]

**Gel Combustion and Secondary Atomization:** The physics of gel droplet combustion and probable secondary atomization processes are described in other work.\(^{12-19}\) In the present model, rigid-shell induced secondary atomization\(^{17-19}\) is assumed. In the combustor code, secondary atomization is presumed to occur when the droplet diameter reaches the predicted rigid-shell diameter. Although secondary atomization actually occurs sometime after rigid shell formation, the time interval is currently unknown and is therefore neglected in the combustor model. Particle size distribution after secondary atomization is also presently unknown and is treated as a system variable. In the combustor model, a droplet undergoing secondary atomization is assumed to shatter into a specified number of equal-size secondary droplets. Defining the fragmentation ratio, \( \beta \), as the number of secondary droplets produced per initial droplet, a new value of \( N/\tau_i \) can be expressed as

\[
\frac{N}{\tau} \bigg|_{\text{new}} = \beta \frac{N}{\tau} \bigg|_{\text{old}}
\]

Equation 10 is then used to determine a new droplet diameter, \( d_j \), given the gel mass flux of the \( j^{th} \) size class.

**Hydrocarbon Vaporization:** Based on the relative proximity of the gel droplets to each other, it is unlikely that the droplets are surrounded by individual flames. Therefore, hydrocarbon combustion is represented using a spherically symmetric droplet evaporation model with convective effects incorporated through film theory.\(^{26}\) As mentioned previously, droplet heat up is approximated by assuming that heat transfer from the gas flow only affects a thin liquid layer at the droplet surface. Assuming quasi-steady, spherically symmetric droplet vaporization with unity Lewis number, uniform droplet temperature, and constant thermophysical properties, the hydrocarbon vaporization rate can be expressed as

\[
\frac{dm_{d,LH}}{dt} = -\frac{2\pi \kappa_w \mu u_r \theta}{c_p} \left[ \frac{c_p}{\rho} \left( \frac{T_r}{T_{b,LH}} - 1 \right) \right] + 1
\]

**Aluminum Combustion:** As mentioned previously, aluminum combustion proceeds through two oxidation mechanisms: aluminum vapor oxidation and droplet surface condensation/oxidation. This aluminum combustion process is treated using a simple combustion model in which a specified fraction of the vaporizing aluminum is involved in the surface oxidation/condensation mechanism. It is assumed that all heat released by this surface mechanism goes into the droplet with none lost to the sur-
rounding gases. Using the same assumptions as the hydro-carbon model and assuming that Al₂O₃ does not interfere with aluminum vaporization, the droplet vaporization rate can be expressed as

\[
\frac{dm_{Al}}{dt} = \frac{2\pi k_A Nu_r c_{pe}}{c_{pe}} \left[ \frac{c_{pe}(T_f - T_{Al})}{1 + \int_{T_{Al}}^{T_f} c_{pe}dT - \eta_{Al,O_3}} \right]
\]

(12)

It should be noted that \(i_{Al,O_3}\) is not a true enthalpy of vaporization since \(Al_2O_3\) does not exist in a vapor state. Therefore, \(i_{Al,O_3}\) is actually the enthalpy change for the chemical reaction \(2Al(g) + \frac{3}{2}O_2 \rightarrow Al_2O_3(l)\) occurring at the droplet surface temperature. Droplet lifetime comparisons with an empirical correlation for aluminum droplet combustion in solid rockets demonstrated good agreement. Average thermophysical properties are estimated as recommended in previous droplet combustion work.

**Solution Method:** Equations 1, 2, 9, 11, and 12 form the governing equation set and are numerically integrated in the axial direction using the IMSL Dverk integration routine. Gas-phase temperature, density, and composition are calculated using the STANJAN chemical equilibrium subroutine. The temperature dependent gas-phase conductivity and viscosity are approximated as those of \(O_2\) to avoid the complexities of calculating gas mixture properties. Details of these and other properties used in the combustor model can be found in reference 30.

**One-Dimensional Combustor Model Results**

The one-dimensional combustor code was exercised using the conditions presented in Table 1. These values were chosen to simulate an upper-stage booster and represent the maximum \(I_{sp}\) operating point for a 60 wt% aluminum gel. Because propellant mass flow rate in a rocket is governed in part by nozzle geometry, the SPP nozzle code was used to determine the propellant mass flow rates. Since an appropriate spray distribution remains to be determined from gel atomization research, the arbitrary normalized droplet size distribution presented in Fig. 1 was used.

Gas temperature and composition, assuming a fragmentation ratio, \(\beta\), of 5 are presented in Fig. 2. The jagged shapes of both the temperature and composition profiles arise from using only four droplet size classes instead of a continuous droplet size distribution. The slow rate of temperature increase in the region between 0.1 and 0.25 m is caused by the large enthalpy transfer from the gas flow to heat the aluminum after hydrocarbon burnout.

In Fig. 3, propellant burnout distance is plotted versus fragmentation ratio, \(\beta\), to illustrate the potential benefits of

<table>
<thead>
<tr>
<th>Table 1. Model Operating Conditions</th>
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<tr>
<td>Chamber Diameter</td>
</tr>
<tr>
<td>Chamber Pressure</td>
</tr>
<tr>
<td>Liquid Carrier</td>
</tr>
<tr>
<td>Gel Aluminum Loading</td>
</tr>
<tr>
<td>Gel Flow Rate</td>
</tr>
<tr>
<td>Oxidizer Flow Rate</td>
</tr>
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</table>

Figure 1. Normalized droplet size distribution used in the combustor code.

Figure 2. Gas temperature and major species mass fluxes versus axial location. Data are for a 60 wt% aluminum gel assuming a fragmentation ratio of 5.
secondary atomization in reducing required engine residence times. It is readily apparent that only slight secondary atomization is required to significantly reduce propellant burnout distance, and that higher secondary atomization intensities, represented by larger fragmentation ratios, have a lesser marginal effect, primarily due to the fact that droplet lifetime is inversely proportional to droplet surface area, which increases as \((\beta)^{2/3}\).

Final oxide residual diameter as a function of fragmentation ratio is shown in Fig. 4. Similar to the trend seen in propellant burnout, small secondary atomization intensities significantly reduce final \(\text{Al}_2\text{O}_3\) residual diameter, with greater atomization intensities providing decreasing marginal reductions in residual diameter. Final residual particle diameter, however, is proportional to \((1/\beta)^{1/3}\) instead of \((1/\beta)^{2/3}\) as is droplet lifetime.

Radiation losses from condensed combustion products to the chamber walls as functions of fragmentation ratio and aluminum mass loading are shown in Fig. 5. Chamber operating conditions are those that produce maximum \(I_{gp}\) for each aluminum loading. Radiation losses were found to be strongly influenced by the fragmentation ratio, primarily due to changes in propellant burnout distance. Increasing the gel aluminum mass loading for a given fragmentation ratio was also found to increase radiation losses because of increased chamber temperature and slight increases in propellant burnout distance with increasing aluminum mass loading. Radiation losses were approximately equal to 2-13% of the energy released during combustion, depending on the aluminum mass loading and fragmentation ratio.

It should be cautioned that the above radiation heat transfer calculations are only approximate. Axial radiation heat transfer, which has been neglected, may alter the droplet combustion process and/or the actual radiation losses. In addition, research has indicated that the optical properties of condensed combustion products can vary considerably depending on propellant composition and oxidizer/fuel mixture ratio.
SPP Modeling of Engine Performance

Nozzle Two-Phase Flow

Two-phase flow effects in the engine nozzle were estimated using the SPP rocket code. Nozzle inlet conditions were determined using a chemical equilibrium module of the SPP code that provides a mechanism to account for the radiation losses predicted by the one-dimensional combustor code.

Because particle interaction and mass transfer between the particles and gas are neglected in the SPP code, the AI2O3 residual size distribution predicted by the combustor code may be incorrect for nozzle performance calculations. Since the AI2O3 particles are molten throughout most of the nozzle, and because small particles accelerate more quickly than large particles, particle size may increase through coagulation. Similarly, additional AI2O3 may be produced through the recombination of gas-phase radicals as the exhaust gases cool during expansion, resulting in the nucleation of additional particles and/or growth of previously formed particles. Particle size may also decrease due to shear breakup of droplets, particularly in the throat region of the nozzle.

Because of these uncertainties in AI2O3 particle size, two methods of estimating particle size, which should bound the true particle size, are used in the evaluation of nozzle performance. In the first method, we use an AI2O3 particle size distribution that is determined by the one-dimensional combustor code, making secondary atomization the primary mechanism governing particle size. In the second method, secondary atomization is assumed to have no effect on mean particle size; rather, coagulation, particle surface growth, and shear induced droplet breakup are assumed to be the dominant mechanisms affecting particle size. Assuming that these coagulation and breakup mechanisms are comparable to those in solid nozzles, the following solid motor correlation of mass median AI2O3 particle diameter, Dₐ₀, as a function of nozzle throat diameter was employed:

\[ D_{a0} = 3.63D_1^{0.932} \]  \hspace{1cm} (13)

This results in a particle mass median diameter of 5.6 \( \mu \)m.

Engine Performance Results

A comparison of two-phase flow effects on Isp for the above methods of determining AI2O3 particle size are presented in Fig. 6 for a 60 wt% aluminum gel. As seen here, secondary atomization may reduce two-phase flow losses but not as significantly as propellant burnout distance (4% versus 62% decrease, cf. Fig. 3). To illustrate the separate contributions of radiation and two-phase flow losses, radiation losses were incorporated in the case represented by the dashed line. Given a fragmentation ratio of 5, radiation losses yield a decrease in Isp of approximately 1% compared to the 4% resulting from two-phase losses.

Using the solid motor correlation for particle size, Isp was calculated for a range of aluminum mass loadings and propellant mixture ratios and compared with Isp calculations for a JP-10/O₂ bi-propellant. Figure 7 shows that Isp decreases with increasing aluminum loading and that the maximum Isp mixture ratio becomes richer, as has been predicted by other studies, although the Isp decrease is
much greater with the incorporation of radiation and two-phase flow losses.

Conclusions

Based on the above analysis of an Al/hydrocarbon/O₂ fueled rocket engine, the following results were obtained:

1. Exercise of the one-dimensional combustor model predicts that only moderate secondary atomization (β=5) is required to reduce overall propellant burnout distance by 62% and final Al₂O₃ residual diameter by 41%.

2. Radiation losses for a 60 wt% gel, assuming β=5, are estimated to be approximately 5% of the energy released during combustion, resulting in a 1% decrease in engine Iₚₑ. Secondary atomization may reduce radiation heat transfer losses, primarily due to decreases in the propellant burnout distance. For example, a fragmentation ratio of five results in a 61% decrease in radiation losses. Radiation losses were also found to be a function of gel composition and engine operating conditions, as a result of changes in chamber temperature.

3. Two-dimensional, two-phase nozzle code results indicate that secondary atomization may have little effect on nozzle two-phase flow losses. Specifically, a fragmentation ratio of 5 decreases two-phase flow losses by only 4% compared to the above 62% reduction in propellant burnout distance. Furthermore, secondary atomization may have no effect on two-phase flow losses if particle coagulation, surface growth, and shear induced breakup are the dominant mechanisms controlling oxide particle size.

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