AN INVESTIGATION OF SLURRY FUEL COMBUSTION

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January 1981

TECHNICAL REPORT AFWAL-TR-80-2125
Final Report for Period November 1979 - October 1980

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AIR FORCE/56780/18 March 1981 — 100
An Investigation of Slurry Fuel Combustion.

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F33615-77-C-2004
03480591
62203F

Jan 1981
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Abstract

A theoretical and experimental investigation of the combustion properties of carbon slurry fuels was performed. This involved observing the combustion of individual drops (400-1000 μm in diameter) supported at various positions within an open, turbulent diffusion flame. When a slurry drop was exposed to the flame, the liquid fuel in the slurry evaporated in the first stage, leaving a carbon agglomerate consisting of all the carbon particles originally in the slurry. The second stage of the process involved heat-up and reaction of the agglomerate (Continued on reverse).
Consumption of the agglomerate was the slowest step in the process, requiring 90-95 percent of the lifetime of the particle, even in regions where agglomerate reaction was fast. An analysis was developed to provide predictions of agglomerate reaction rates and lifetimes. The model was tested with measurements in the lean portions of the flame (equivalence ratios in the range 0.272-0.778). Good agreement between predictions and measurements was observed in this region, with respect to both particle temperature and size variations. Use of catalyzed slurry was found to increase burning rates and extend the lean limit for agglomerate reaction. It is recommended that further testing and model development be undertaken, considering combustion properties at higher equivalence ratios and for drops more representative of the sizes encountered in combustion chambers. The model should also be extended to consider the liquid gasification stage of slurry combustion.
PREFACE

This research supported by the Aero Propulsion Laboratory, Air Force Wright Aeronautical Laboratories, Air Force Systems Command, Wright-Patterson Air Force Base, Ohio, under Project 3048, "Fuels, Lubrication, and Fire Protection," Task 304805, "Aero Propulsion Fuels," Program Element 62203F. Dr Alvin F. Bopp of the Fuels Branch served as the Air Force project scientist.

The fuels used in the study were provided by R. S. Stearns, of Suntech, Inc., Marcus Hook PA. The authors wish to acknowledge useful discussions with R. S. Stearns of Suntech, and with T. W. Bruce and H. Mongia, of AiResearch Manufacturing Company of Arizona, Phoenix, Arizona.
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Summary

The results of a brief preliminary study of the combustion properties of carbon slurry fuels are reported. The carbon slurry fuels were formulated by mixing noncatalyzed and/or catalyzed medium thermal carbon black (roughly 50 percent by weight) with JP-10.

The combustion of individual supported slurry drops, 400-1000 μm initial diameter, was observed in an open turbulent diffusion flame at atmospheric pressure. The environment of the drops within the flame was known from measurements of mean velocity, temperature and composition. The drops were supported on the junction of a thermocouple. Shadowgraph motion pictures were also obtained for the process. This data provided the variation of particle temperature and diameter as a function of time.

A two-stage combustion process was observed. The first stage involved heat-up and evaporation of the liquid, leaving a solid agglomerate of the carbon particles in the slurry. The second stage involved heat-up and reaction or quenching of the agglomerate.

A model was constructed for the second stage of the process. This involved analysis of heat and mass transfer processes in the convective environment surrounding the agglomerate, and the chemical reaction of carbon with oxygen, carbon dioxide, and water vapor at the surface of the particle. Radiative heat transfer between the agglomerate and its surroundings was also considered. The predictions of the model were compared with measurements in the lean portions of the flame (equivalence ratios in the range 0.272-0.778).

The major findings of the study can be summarized as follows:

1. For present test conditions, all the solid carbon in the slurry remained in the agglomerate and heat-up and reaction of the agglomerate dominated the lifetime of the particle—usually requiring 90-95 percent of the particle lifetime, even in locations where agglomerate reaction was nearly diffusion controlled.

2. While liquid heat-up and gasification occurs in all regions of the flame, agglomerate reaction was limited to regions with equivalence ratios in the range 0.37-4.0 for noncatalyzed slurries and 0.21-4.0 for catalyzed slurries. The carbon reaction was quenched at other locations.

3. The model yielded good predictions of agglomerate reaction rates in the range tested (equivalence ratios of 0.272-0.778, where the main reactant is oxygen). Conditions near both the diffusion and reaction rate controlled limits were included in the comparison. This agreement was obtained by selecting a single empirical, area/reactivity multiplication factor, which allows for the pore and surface structure of the carbon agglomerate and the effect of a catalyst.
4. The model yielded good predictions of agglomerate-life-histories over the same range of conditions. Reasonable predictions of heat-up times and particle temperatures indicate that the thermal aspects of the model are adequate over the present test range.

5. Adding catalyst increased agglomerate burning rates, for conditions not too near the diffusion controlled limit. This effect could be treated in the theory by selecting a higher area/reactivity multiplication factor for the catalyzed slurry. The presence of catalyst also extended the lean limit for quenching.

6. Some particle shattering was observed with catalyzed slurries in the lean region of the flame.

These conclusions are based on test results with relatively large particles, 400-1000 μm in diameter. In particular, the effect of pore diffusion is likely to vary with particle size. Therefore, the area/reactivity multiplication factors selected to correlate the present data are not considered to be appropriate for other test conditions, without further testing. The effect of pressure and combustion properties in the fuel rich region (equivalence ratios of 0.778-4.0) were also not assessed in this study. This implies considerable uncertainty concerning agglomerate reaction with carbon dioxide, water vapor, and other species present at high equivalence ratios. In order to remove these uncertainties, it is recommended that additional work be undertaken using particle sizes more representative of combustion chamber conditions, as well as test environments extending to richer mixture ratios.
1. Introduction

1.1 General Objectives

The propulsion system of airbreathing missiles is frequently volume limited. This implies that the volume of the propulsion system is a critical factor in determining missile range and performance—as opposed to propulsion system weight. Since fuel volume comprises a significant fraction of propulsion system volume, fuels that have a large chemical energy release per unit volume (high volumetric energy density) are particularly attractive for such applications.

There has been continuous development of high volumetric energy density liquid fuels to meet these needs [1].* However, it has become increasingly difficult to achieve further improvements using liquids alone. This has resulted in increased consideration of slurries, since these fuels can be handled and burned similar to liquids and yet have increased density due to the presence of solids [2-5]. The objective of the present investigation was to conduct a preliminary investigation of the combustion properties of carbon slurry fuels of interest to the Air Force [2,3].

The carbon slurry fuels considered in this investigation were formulated by Suntech, Inc., Marcus Hook, PA, for use during combustor tests by the AiResearch Manufacturing Company of Arizona [2,3]. While these fuels are representative of candidate carbon slurry fuels, they do not necessarily represent the most attractive formulations since fuel development efforts are still in progress. Nevertheless, the materials should still provide a reasonable indication of carbon slurry combustion properties, useful for subsequent fuel development efforts.

The properties of the fuel considered during the present investigation are summarized in Table 1. The fuels consist of a medium thermal carbon black, with JP-10 as the liquid fuel. Two slurries were tested, each containing roughly 50% carbon by weight. The ultimate carbon particle size was 0.3 μm. One slurry was catalyzed with a proprietary lead compound.

The overall combustion characteristics of carbon slurry fuels have been examined in gas turbine combustors [3], and in well-stirred reactors [5]. The objective of the present investigation was to examine slurry combustion on a more fundamental level, by observing the combustion of individual slurry drops. The experimental technique involved observations of single slurry drops (400-1000 μm in diameter). The drops were supported at various positions within a turbulent diffusion flame in order to provide an environment representative of actual combustion chamber conditions. Measurements were made of the variation of particle diameter and temperature as a function of time. The process was also modeled theoretically and predictions and measurements were compared.

*Numbers in brackets denote references.
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<th>Catalyzed&lt;sup&gt;b&lt;/sup&gt;</th>
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<td>790-942</td>
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<tr>
<td>Liquid</td>
<td>JP-10</td>
<td>JP-10</td>
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<tr>
<td>Dispersed Carbon&lt;sup&gt;c&lt;/sup&gt; (wt %)</td>
<td>50.4</td>
<td>49.2</td>
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<sup>a</sup>Fuels supplied by R. S. Stearns, Suntech, Inc., P. O. Box 1135, Marcus Hook, PA 19061.

<sup>b</sup>The catalyst is a proprietary lead compound [6].

<sup>c</sup>The dispersed carbon is a medium thermal carbon (carbon black) having an ultimate particle size of 0.3 μm. Further properties of this material are provided in Ref. 2.
1.2 Previous Studies

The present study is an extension of earlier work in this laboratory [3,4]. The test fuel and many aspects of the experiments are similar. The general test procedure involved observations of slurry drops supported in an open turbulent diffusion flame.

The major findings of Ref. 4 are as follows:

1. The combustion of a slurry drop occurred in four stages:
   1) heat-up of the drop and, at times, ignition of liquid fuel vapors; 2) evaporation or combustion of the liquid fuel accompanied by agglomeration of the carbon particles in the slurry; 3) further heating of the carbon agglomerate once all the liquid has evaporated; and 4) fragmentation, reaction or quenching of the agglomerate.

2. Combustion of the carbon agglomerate only occurred over a limited equivalence ratio range (0.21-4 for the catalyzed slurry, 0.37-4 for the noncatalyzed slurry).

3. Even in regions where the carbon agglomerate reacted, the heat-up and reaction of the agglomerate was the slowest step in the combustion process, generally requiring 20 times longer than the time required to gasify the liquid.

Figure 1 is a scanning electron microscope photograph of the surface of a partly reacted agglomerate. The light areas on the photograph are the surface of the solid agglomerate, the dark areas represent pores. The agglomerate is clearly very porous, with the carbon surface area available for reaction much greater than the apparent outer surface area of the agglomerate.

These general observations concerning the combustion properties of carbon agglomerates are similar to results found for coal slurries [7,8]. In the case of coal, however, the presence of ash and volatile material in the coal, materially influences the details of the process.

1.3 Specific Objectives

Previous investigations have demonstrated the importance of agglomerate formation and reaction during the combustion of carbon slurries. Some information regarding the regions of flames where agglomerates react has also been generated. However, existing data on agglomerate reaction rates are very limited. Furthermore, no attempt to model the agglomerate reaction process has been reported.

With this status in mind, this investigation extended the results of Refs. 3 and 4, employing a similar experimental arrangement. The specific objectives of the study were as follows:
Figure 1. SEM photograph of agglomerate from a catalyzed slurry drop in the glowing region. x1500. 100x magnification.
1. Complete additional measurements of the variation of particle size and temperature with time for drops supported at various positions within a turbulent diffusion flame. In particular, generate more complete information on agglomerate burning rates as a function of position in the flame.

2. Formulate a model for the reaction of the carbon agglomerate, using existing information on carbon reaction rates. Employ the model to compute agglomerate-life-histories and compare predictions and measurements.

The supported particles used during this investigation had diameters in the range 400-1000 μm, which is much larger than typical combustor applications. For preliminary work, however, this approach had advantages since more detailed measurements can be made with large particles, helping to provide more insight concerning slurry combustion properties.

2. Test Arrangement

2.1 Apparatus

The test apparatus involved modification of an earlier arrangement employed for a study of gas and spray diffusion flames [9,10]. A sketch of the modified apparatus appears in Fig. 1. The arrangement consists of an injector flowing propane gas, which is burned as a diffusion flame in stagnant room air. The test stand was supported from either a thermocouple or a quartz probe at various locations in the flame. The drops are also photographed with a motion picture camera, as shadowgraphs.

The propane gas injector was positioned near the bottom of the test stand and oriented vertically upward. The test stand was an area 1.2 m square and 3 m high enclosed with a single thickness of 16 mesh screen. The test stand was located within a room having dimensions 4 m x 7 m x 4 m high. Combustion products were removed through an exhaust fan located near the ceiling of the test cell. The injector was secured on a support which allowed a vertical movement of 1.2 m. Two traversing mechanisms were used to position the nozzle in the horizontal plane.

The nozzle was a full-cone air atomizing injector, with no swirl, manufactured by the Spraying Systems Company (Model 1/4"1200 fluid nozzle and Model 67147 air nozzle). The outlet diameter of the nozzle was 1.194 mm.

The gas flame was stabilized near the exit of the injector using an array of four hydrogen capillary flames. The flame tubes were mounted in a symmetrically opposed pattern injecting toward the centerline of the injector passage. The flow passages for the hydrogen were slots 0.4 mm wide and 2 mm long, with the long axis parallel to the injector centerline. The lower edges of the slots were 1.8 mm above the face of the injector. The exit planes of the slots were 0.4 mm apart.
The propane gas flow rate was metered with a Matheson Model 604 rotameter and controlled by a Harris Model 2515 pressure regulator with an output capacity of 0-0.69 MPa. The flow rate of the hydrogen gas was metered with a Matheson Model 601 rotameter and controlled with a Matheson Model 11 pressure regulator having an 0-1.38 MPa output capacity. The rotameters for both the propane and hydrogen flows were calibrated with a Precision Scientific Company wet-test meter (2.83 ml/rev).

The fuel drops were mounted with a hypodermic needle. The flame was deflected from the mount region until the droplet was in place. The deflector was constructed from a 200 x 305 mm sheet of stainless steel, having a thickness of 3 mm. The drop combustion process was initiated by removing the deflector with a pneumatic cylinder arrangement.

2.2 Instrumentation

2.2.1 Drop Environment Measurements

The drop environment at each test location within the flame was measured. This involved measurements of mean velocity, temperature and composition. Mean velocities were measured with a laser-Doppler anemometer, which employs frequency shifting so that flow reversals could be detected. Mean temperature was measured with a 50 μm diameter platinum/platinum-10% rhodium thermocouple (with a radiation error of less than 35°C over the test range). Mean concentrations were measured by extracting gas samples isokinetically and analyzing them with a gas chromatograph. The details of these techniques are presented elsewhere [9,10].

2.2.2 Drop Measurements

The drop combustion process was observed with a 16 mm, Photosonics, Model 16-B, motion picture camera. The camera optics provided a 2:1 magnification. The camera was powered with a Kepco, SM 36-5 AM d.c. power supply. The film speed was indicated with a timing light on the camera, activated with an Adtrol Electronic pulse generator, Model 501. Kodak Plus-X reversal film was used for all tests.

Backlighting for shadowgraph measurements was provided by a PeK, Model 401A, 75W mercury arc lamp. The light from the arc was collimated and directed toward the drop location. A diffuser screen was employed behind the drop to equalize the light intensity of the background. The background intensity was adjusted so that envelope flames around the drop could also be observed.

The film records were analyzed on a frame by frame basis, using a Vanguard motion picture analyzer. Photographs of objects of known size at the drop location provided a calibration of distances on the film.
Particle temperatures were measured by mounting the test drop at the junction of a thermocouple. A Pt/Pt-10% Rh thermocouple was used, constructed of 75 μm diameter wires. In order to help support the drop, the junction was placed within a round bead, formed with Sauereisen cement (300-400 μm in diameter). The output of the particle thermocouple was recorded digitally on a Nicolet Explorer Oscilloscope, Model 206, and reduced to temperature-time data on a computer.

2.3 Test Conditions

Tests were conducted using the burner operating condition summarized in Table 2. The drops were mounted along the centerline of the flow. Relatively complete structure measurements have been reported for this flame, as noted earlier [9]. Details of interest to this investigation will be discussed later.

3. Theoretical Considerations

3.1 Description of the Model

Results obtained during this and earlier investigations indicate that slurry drop combustion is primarily a two-stage process involving gasification of liquid in the first stage and heat-up and reaction of the agglomerate in the second stage. Since the agglomerate does not actively participate in the first stage, other than influencing the density of the particle, it is convenient to treat these two stages separately.

The following analysis will be limited to the second stage—involving heat-up and reaction of the agglomerate. The present investigation emphasized this aspect of the problem since reaction of the agglomerate is the slowest step in the process. Therefore, agglomerate reaction generally controls combustor performance. Furthermore, agglomerate reaction presents the greatest uncertainties; in contrast, gasification of liquid drops is reasonably well understood [11].

The initial condition of the present analysis is the point where all the liquid has vaporized and the agglomerate is at the wet-bulb temperature appropriate for liquid vaporization (the wet-bulb temperature is slightly below the boiling temperature of the liquid for a pure fuel like JP-10 [11]). The objective of the analysis is to predict the subsequent heat-up and reaction of the agglomerate. This involves determining the variation of particle temperature and diameter as a function of time.

3.2 Agglomerate Reaction Model

Since the test droplets were supported at fixed locations, the motion of the particle is not considered. The formulation was limited to determining the rate of consumption of carbon in the
Table 2
Burner Flame Characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuel:</td>
<td>propane</td>
</tr>
<tr>
<td>Fuel flow rate:</td>
<td>176 mg/s</td>
</tr>
<tr>
<td>Initial fuel jet velocity:</td>
<td>88.7 m/s</td>
</tr>
<tr>
<td>Initial fuel jet diameter:</td>
<td>1.194 mm</td>
</tr>
<tr>
<td>Jet Reynolds number:</td>
<td>23600</td>
</tr>
<tr>
<td>Orientation:</td>
<td>vertical (upward)</td>
</tr>
<tr>
<td>Flame Height:</td>
<td>460 mm (visual)</td>
</tr>
<tr>
<td>Injector Thrust:</td>
<td>15.6 mN</td>
</tr>
<tr>
<td>Hydrogen flow rate:</td>
<td>0.126 mg/s</td>
</tr>
<tr>
<td>Ambient and injector inlet temperature:</td>
<td>296 K</td>
</tr>
<tr>
<td>Ambient pressure:</td>
<td>97 kPa</td>
</tr>
</tbody>
</table>

*Nozzle is Spraying Systems Co. Model 1/4 J 2050 fluid nozzle and Model 67147 air nozzle, air atomizing injector.*
aggomerate, the rate of heat transfer to the particle, and the rate of chemical energy release at the surface of the particle.

A sketch of the model appears in Fig. 3. The major assumptions of the analysis are as follows:

1. The particle is assumed to be spherical and particle spacing is large so that the surroundings can be taken as an infinite medium having uniform properties.

2. The gas flow field around the particle is assumed to be quasisteady, i.e., the flow field has properties equivalent to a steady state flow field, for the same boundary conditions, at each instant of time.

3. Flow fields around spheres are too complex to analyze exactly; therefore, the conventional film theory approximation is employed to represent the effect of convection [11]. This assumption implies that gas phase transport processes can be represented as a spherically symmetric flow through a stagnant layer whose outer boundary is fixed empirically.

4. The particle is assumed to consist of pure carbon and has a uniform temperature at each instant of time.

5. Solid carbon is only assumed to react at the apparent surface of the particle. The effect of pores and reaction within the particle is treated by introducing an empirical area/reactivity multiplication factor. Reaction of solid carbon with oxygen, carbon dioxide, and water vapor is considered.

6. The gases surrounding the particle are assumed to be represented by their mean properties and the effect of turbulent fluctuations is ignored. The actual gas mixture was approximated by oxygen, nitrogen, carbon dioxide, carbon monoxide, water vapor, hydrogen and hydroxyl radicals—in thermodynamic equilibrium.

7. Only concentration diffusion is considered, employing an effective binary diffusion coefficient for all diffusing species. All species are assumed to have equal specific heats (assumed constant) and molecular weights, for the purpose of computing diffusion and energy transport rates in the flow. Constant average values are used for all properties in the gas phase.

8. The gas is assumed to be transparent to radiation, however, provision is made for particle radiation to the confinement surrounding the flow.

9. The pressure within the flow field is assumed to be constant.

10. The gases maintain thermodynamic equilibrium at the particle surface however, the flow is chemically frozen elsewhere.
These assumptions are generally similar to those employed in recent analyses of carbon particle reaction [12,13].

3.2.1 Gas Phase Transport

The governing equations consist of equations for conservation of mass, energy and species. The latter equation can be represented more conveniently in the present analysis, where equal binary diffusivities are assumed, as equations for conservation of elements—since elements are conserved during chemical reactions. Employing the assumptions of the analysis, the following equations apply [14].

\[ \rho vr^2 = \frac{m'' r^2}{p} \]  

(1)

\[ \frac{m'' r^2}{p} \frac{d}{dr} \left( \frac{2dh}{dr} \right) = \frac{\lambda}{\rho c_p} \left( \frac{d}{dr} \left( \frac{r^2 dh}{dr} \right) \right) \]  

(2)

\[ \frac{m'' r^2}{p} \frac{d}{dr} \left( \frac{dY_i}{dr} \right) = \rho D_i \left( \frac{d}{dr} \left( \frac{dY_i}{dr} \right) \right), \quad i = 1, 4 \]  

(3)

Four element fractions are considered in the analysis, \( Y_1, Y_2, Y_3, Y_4 \) and \( Y_5 \).

Seven species were considered: \( O_2, N_2, CO_2, CO, H_2O, H_2 \) and \( OH \). Following Refs. 12 and 13, \( O \) was ignored on the grounds that its concentration is too small, for temperatures below 4000K, for it to contribute significantly to the reaction of carbon. Species and element concentrations are related by the following mass balances

\[ \sum_{j=1}^{7} \nu_{ij} Y_j = \nu_i, \quad i = 1, 4 \]  

(4)

where \( \nu_{ij} \) is the mass fraction of the \( i \)th element in the \( j \)th species.
The enthalpy appearing in Eq. (2) is given by

\[ h = \sum_{i=1}^{7} Y_i h_i \]  

(5)

where

\[ h_i = C_{p_i} (T-T_{ref}) + h_i^{\infty}_{ref} \]  

(6)

The boundary conditions for Eqs. (2) and (3) are as follows:

\[ r=r_p, \ h=h_p, \ \bar{Y}_i=\bar{Y}_i_p; \ r=r_\infty, \ \bar{Y}_i=\bar{Y}_i_\infty; \ r=r_\infty, \ h=h_\infty \]  

(7)

The ambient conditions are specified. The conditions at the particle surface must be determined from the analysis of surface reaction. This will be discussed later.

The general case is treated in Eq. (7), where the film radii for mass and heat transfer, \( r_\infty \) and \( r_\infty \), are not the same. These radii are found employing the usual correlations from film theory [11]. For mass transfer,

\[ \frac{r_\infty}{r_p} = \frac{Sh}{(Sh-2)} \]  

(8)

while for heat transfer

\[ \frac{r_\infty}{r_p} = \frac{Nu}{(Nu-2)} \]  

(9)

The Sherwood and Nusselt number correlations employed in Eqs. (8) and (9) are found from the following correlation [11].

\[ (Nu \text{ or } Sh)-2 = \frac{0.555 \ Re^{1/2} (Pr \text{ or } Sc)^{1/3}}{(1+1.232/(Re(Pr \text{ or } Sc)^{4/3}))^{1/2}} \]  

(10)
Equations (2) and (3) can be integrated, subject to the boundary conditions of Eq. (4), to yield

\[ \frac{\dot{y}_i}{Y_i} = \frac{(\dot{Y}_{i_p} - \dot{Y}_{i_p}) \exp(-K/\zeta) + Y_{ip} \exp(-K/\delta) - \dot{Y}_{i_p} \exp(-K)}{\exp(-K/\delta) - \exp(-K)} \tag{11} \]

\[ h = \frac{(h_i - h_p) \exp(-K/\zeta L_e) + h_p \exp(-K/\gamma L_e) - h_o \exp(-K/L_e)}{\exp(-K/\gamma L_e) - \exp(-K/L_e)} \tag{12} \]

where

\[ \zeta = r/r_p \tag{13} \]

\[ \delta = r_{oo} / r_p \tag{14} \]

\[ \gamma = r_{oo} / r_p \tag{15} \]

and \( K \) is a nondimensional particle mass loss parameter, defined as

\[ K = m'' r_p / \rho D \tag{16} \]

A relationship between \( K \) and the \( Y_{ip} \) can be developed by noting that there is no net mass flux of the elements oxygen, hydrogen or nitrogen at the particle surface. Furthermore, the net mass flux of carbon at the surface is equal to \( m'' \), since the particle is assumed to be pure carbon. Therefore, at \( \zeta=1 \), we have

\[ \frac{d\dot{Y}_i}{d\zeta} = K_\gamma Y_i \quad i = 1, 4 \quad ; \quad i \neq C \tag{17} \]
Taking the derivative of Eq. (11) and substituting into Eqs. (17) and (18) yields the following:

\[
\frac{d\hat{Y}_C}{d\sigma} = -K(1-\hat{Y}_C)
\]  \hspace{1cm} (18)

Employing the usual sign convention that convection to the surface of the particle is positive, and recalling that equilibrium has been neglected in the gas phase, yields

\[
\dot{Y}_{p, i} = \hat{Y}_{i, \infty} \exp(K(\delta^{-1}-1)) , \hspace{0.5cm} i = 1,4 \hspace{0.5cm}, \hspace{0.5cm} i \neq C
\]  \hspace{1cm} (19)

\[
Y_{p, C} = 1-(1-Y_{C, \infty}) \exp(K(\delta^{-1}-1))
\]  \hspace{1cm} (20)

where the second part of Eq. (21) follows from the assumption of constant gas phase specific heat. Taking the derivative of Eq. (12) and evaluating the resulting expression at \( r = r_p \), yields

\[
\frac{q''_{c, p}}{K(h_{\infty} - h_p)} = \frac{-\exp(-K/Le)}{\exp(-K/\gamma Le) - \exp(-K/Le)}
\]  \hspace{1cm} (22)

Given \( K \), Eqs. (19) and (20) yield the four element mass fractions at the surface of the particle. Equation (11) then provides four of the equations to determine the seven unknown species mass fractions. Imposing the condition of chemical equilibrium, yields the three more equations needed to determine the \( Y_i \). The following dissociation reactions were employed...
This yields the following equilibrium requirements

\[
\begin{align*}
K_y_1 & \quad H_2O \rightleftharpoons H_2 + \frac{1}{2} O_2 \\
K_y_2 & \quad H_2O \rightleftharpoons \frac{1}{2} H_2 + OH \\
K_y_3 & \quad CO_2 \rightleftharpoons CO + \frac{1}{2} O_2
\end{align*}
\] (23)

\[
\begin{align*}
K_y_1 p^{-1/2} &= Y_{H_2} Y_{O_2}^{1/2} / Y_{H_2O} \\
K_y_2 p^{-1/2} &= Y_{H_2}^{1/2} Y_{OH} / Y_{H_2O} \\
K_y_3 p^{-1/2} &= Y_{CO} Y_{O_2}^{1/2} / Y_{CO_2}
\end{align*}
\] (24) (25) (26)

Since the \( K_y \) are known functions of temperature, Eqs. (4), (19), (20) and (24)-(26), allow \( Y_i \) and \( Y_{ip} \) to be determined, given the ambient composition, \( K \), and the surface temperature. The next section will consider the relationship between \( K \) and the \( Y_{ip} \).

Constant average properties were employed at each instant of time, using methods similar to earlier work in this laboratory [9]. The methods used to compute properties are described in Appendix A.

3.2.2 Surface Reactions

The reaction of carbon is a complex process, particularly when the material is in a combustion product mixture containing many species. Reaction mechanisms of carbon involving \( O_2 \), \( CO_2 \), \( H_2O \), \( H_2 \), \( OH \) and \( O \) can play a role [15,16]. During the present analysis, only the first three species \( O_2 \), \( CO_2 \) and \( H_2O \) were considered. The present methodology, however, is applicable to more complex treatment of reaction as new information becomes available.
Other problems that must be considered, include: 1) reaction is
not just limited to the apparent outer surface, since pores are present;
and 2) the presence of a catalyst influences reaction rates. The effect
of both these factors was treated in the present analysis by intro-
ducing empirical area/reactivity multiplication factors for each
reaction contributing to the consumption of carbon. The values of the
area/reactivity factors were specified by matching predictions and
measurements.

The reaction rates for carbon with oxygen and carbon dioxide em-
ployed the same kinetic parameters as Libby and Blake [13]. The
reactions are assumed to proceed by the following global expressions

\[ R_1 = \frac{1}{2} \text{C} + \text{O}_2 \rightarrow \text{CO} \quad (27) \]

\[ R_2 = \text{C} + \text{CO}_2 \rightarrow 2\text{CO} \quad (28) \]

where the specific rate constants are

\[ R_1 = K_1 p_{O_2}, \quad R_2 = K_2 p_{CO_2} \quad (29) \]

where

\[ K_i = A_i \exp(-E_i/RT), \quad p_i = p_{MY_i}/M_i \quad (30) \]

The specific values of the chemical parameters appearing in Eqs. (29)
and (30) are summarized in Table 3.

The reaction rate expression for water vapor employs the results
of Johnstone, et al. [17]. The reaction mechanism is as follows [15]
Table 3

**Summary of Reaction Rate Parameters**

<table>
<thead>
<tr>
<th>i</th>
<th>A</th>
<th>E (kcal/gmol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>87100 kg/m²s atm</td>
<td>35.8</td>
</tr>
<tr>
<td>2</td>
<td>2470 kg/m²s atm</td>
<td>41.9</td>
</tr>
<tr>
<td>3</td>
<td>5.15 x 10^{-3} kg/m²s atm</td>
<td>32.7</td>
</tr>
<tr>
<td>4</td>
<td>9.42 x 10^{-11}</td>
<td>-60.8</td>
</tr>
<tr>
<td>5</td>
<td>7.07 x 10^{-16}</td>
<td>-79.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Slurry</th>
<th>Reactivity Factor, a_i</th>
</tr>
</thead>
<tbody>
<tr>
<td>noncatalyzed</td>
<td>58.2</td>
</tr>
<tr>
<td>catalyzed</td>
<td>70.6</td>
</tr>
</tbody>
</table>

*a* Assuming a surface area of reaction of 1.15 m²/µ of carbon, which is the average surface area over the period of reaction [17].

*b* Reactivity factor taken to be the same for reaction with O₂, CO₂ and H₂O.
The reaction rate of carbon due to the presence of water vapor can then be expressed as

\[
R_j = \frac{K_3P_{H_2O}}{1 + K_4P_{H_2} + K_5P_{H_2O}}
\]  

The parameters in this case, appearing in Table 3, were obtained from Ref. 17. In order to obtain the rate constants of Table 3 for these reactions, it was necessary to estimate the actual surface area for reaction for the experiments of Ref. 17. The surface areas are taken as the average over the range of reaction, measured by nitrogen adsorption [17].

The total reaction of carbon was obtained as the sum of the rates or O_2, CO_2 and H_2O, ignoring potential interactions between the reactants. This yields

\[
K = \sum_{i=1}^{r_j} \frac{P_i}{D_i} R_i
\]

where the \(a_i\) appearing in Eqs. (34) are the area/reactivity multiplication factors. These parameters were varied in the present investigation in order to match burning rate results to predictions. In general the presence of pores and a catalyst result in \(a_i\) values greater than unity.
3.2.3 **Agglomerate Life-History Computations**

Given the temperature and size of the particle, and the ambient conditions, solution of Eqs. (4)-(31) are sufficient to provide \( Y_p, Y_p', q'' \) and \( K \). The variation of the size and temperature of the particle are determined by solving equations for conservation of particle mass and energy.

Conservation of mass for the particle is

\[
\frac{d}{dt} \left( \frac{4}{3} \pi r_p^3 \rho_s \right) = -4 \pi r_p^2 m''_p
\]

(35)

Assuming that the particle density is constant,

\[
\frac{dr_p}{dt} = -\rho DK/\rho_s r_p
\]

(36)

introducing \( K \) from Eq. (16).

Conservation of energy at the surface of the particle yields

\[
\frac{d}{dt} \left( \frac{4}{3} \pi r_p^3 \rho_s h_s \right) = 4 \pi r_p^2 \left( \Sigma m''_l h_l + q'' + q''_r \right)
\]

(37)

where

\[
q''_r = \sigma \varepsilon (T_p^4 - T_w^4)
\]

(38)

and \( T_w \) is the temperature of the enclosure surrounding the hot gas. Simplifying Eq. (34) by introducing conservation of mass, Eq. (17), yields

\[
\frac{dT_p}{dt} = \frac{-3}{(\rho_s c_s r_p)} \left[ m''_p (h_p - h_s) + q''_c + q''_r \right]
\]

(39)

During some experimental conditions, the rate of radius change of the particle was relatively slow. In this case, the particle would reach a quasisteady equilibrium condition where its rate of temperature
change with time was small. These quasisteady particle temperatures were determined by setting the left-hand side of Eq. (39) equal to zero and solving the equation for particle temperature.

Conservation of particle momentum would yield an additional equation to determine the motion of the particle. Since supported particles were considered during this investigation, these results are not needed and will not be considered here.

The initial conditions for Eqs. (36) and (39) are

\[ t = 0, \quad r_p = r_p^o, \quad T_p = T_p^o \]  

(40)

Equations (36) and (40) were numerically integrated on the computer, using Gear's method. The nonlinear system of algebraic equations which must be satisfied in order to determine \( Y_{i_p}^p, Y_{i_p}^q, \) etc., were solved using the Newton-Raphson method, given the current values of \( T \) and \( r_p \), and the ambient conditions, at each instant of time. Quasisteady burning rates were determined by iterating on \( T_p \), until the left-hand side of Eq. (39) was zero.

4. Results and Discussion

4.1 General Observations

Figure 4 is an illustration of the variation of mean properties along the centerline of the propane flame [9]. Near the injector exit, gas velocities and the concentration of propane are relatively high, while the temperature is low and oxygen is absent. Moving downstream, the gas velocity and the concentration of propane decrease monotonically. The temperature increases at first, reaching a maximum at the point where the fuel has disappeared and combustion product concentrations are highest. Beyond this location of the flame, the flow decays. In this region, temperatures and product gas concentrations decrease, while oxygen and nitrogen concentrations increase.

Similar to our previous slurry combustion study [4], present measurements were obtained along the centerline of this flame. Also marked on Fig. 4 are various combustion regions determined for both catalyzed and noncatalyzed slurry drops during our earlier work [4]. Combustion performance for each of these regimes described in Table 4.
Figure 4. Flow properties and slurry drop combustion regions in the turbulent flame [4].
Table 4
Summary of Slurry Drop Combustion Regions in the Turbulent Flame

<table>
<thead>
<tr>
<th>Region</th>
<th>Designation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fragmentation</td>
<td>F</td>
<td>The liquid fuel evaporated without an envelope flame; however, a diffuse luminous wake was observed. When all the liquid was gone, some of the carbon agglomerate was fragmented into 3-5 large flakes. A carbon agglomerate also remained on the probe.</td>
</tr>
<tr>
<td>Noncombusting</td>
<td>NC</td>
<td>The liquid fuel evaporated without an envelope flame; however, a diffuse luminous wake was observed. When all the liquid had evaporated, a luminous wake or reduced intensity was still observed. A carbon agglomerate remained on the probe.</td>
</tr>
<tr>
<td>Full combustion</td>
<td>FC</td>
<td>An envelope flame was observed around the particle until the liquid fuel was consumed. After a dark period, the carbon agglomerate began to glow. Glowing continued until virtually all the carbon was consumed.</td>
</tr>
<tr>
<td>Glowing</td>
<td>G</td>
<td>The liquid fuel evaporated with no envelope flame observed. Some time after the liquid was gone (2-15 s for present test conditions), the carbon agglomerate began to glow. Glowing persisted until virtually all the carbon was consumed. This region was only observed for the catalyzed slurry.</td>
</tr>
<tr>
<td>Evaporation</td>
<td>E</td>
<td>The liquid fuel evaporated with no envelope flame observed. Glowing of the carbon agglomerate was not observed. No change of the particle size was observed once the liquid was gone for the noncatalyzed slurry. Some reaction of the catalyzed slurry was observed in the upstream end of this zone.</td>
</tr>
</tbody>
</table>

\(^{a}\)In order of increasing distance from the injector.
The present measurements were confined to the range $225 \leq x/d \leq 310$. This includes the full combustion (FC), glowing (G) and evaporation (E) regions. In the first two regions, combustion of the carbon agglomerate is observed, although the second region is only observed for the catalyzed slurry. In the evaporation region, combustion of the agglomerate is quenched. This region is of marginal interest for the present study, however, it is desirable to predict its onset. Table 5 is a summary of flow properties at each of the drop test locations.

Figure 5 is an illustration of a noncatalyzed slurry drop-life-history at $x/d = 297.5$, where agglomerate reaction rates are reasonably high. Particle diameter and temperature are plotted as a function of time. The time scale has been expanded during the first portion of the plot, when the liquid is gasified, since this period is relatively short.

The four main stages of slurry drop combustion are quite apparent in Fig. 5. The initial phase of the process involves heat-up of the particle, approaching the wet-bulb temperature for liquid evaporation (which is somewhat below the boiling temperature of the liquid [11]). In this period, the rate of liquid evaporation is relatively low and the particle diameter remains nearly constant. As the surface temperature nears the wet-bulb temperature, the rate of gasification increases, beginning the second stage of the process. During the second stage, the particle diameter decreases while the particle temperature remains nearly constant. The third stage involves heat-up of the agglomerate, once all the liquid has evaporated. The variation of particle diameter is relatively slow in this period, due to low carbon reaction rates at low temperatures. As particle temperatures increase, reaction rates increase as well, signaling the onset of the fourth stage of the process. During the fourth stage, the agglomerate reacts, with particle temperatures near the gas temperature. In this regime, the particle temperature can be either above or below the gas temperature, depending upon the magnitudes of energy released by reaction and heat transport by convection and radiation. These quantities tend to vary with particle size, so there is no fixed particle temperature in the fourth regime, even though ambient conditions are fixed. As the particle becomes small, however, convection heat transfer dominates, and the particle tends to approach the local gas temperature.

It is evident from the results illustrated in Fig. 5, that heat-up and reaction of the agglomerate comprises the bulk of the lifetime of the particle.

4.2 Agglomerate Burning Rates

Figure 6 is an illustration of predicted and measured agglomerate burning rates for noncatalyzed drops. Results are shown for various positions in the range $297 \leq x/d \leq 510$. This region is in the lean portion of the propane flame, corresponding to an equivalence ratio range of 0.761-0.272. For these conditions, reaction of carbon with oxygen dominates the process, and the effects of $CO_2$ and $H_2O$ are small.
Table 5
Summary of Flame Properties at Droplet Test Locations

<table>
<thead>
<tr>
<th>x/d</th>
<th>T (K)</th>
<th>u (m/s)</th>
<th>( \phi )</th>
<th>O₂</th>
<th>CO₂</th>
<th>CO</th>
<th>N₂</th>
<th>H₂O</th>
</tr>
</thead>
<tbody>
<tr>
<td>170</td>
<td>1841</td>
<td>9.93</td>
<td>1.350</td>
<td>0.006</td>
<td>0.101</td>
<td>0.079</td>
<td>0.653</td>
<td>0.161</td>
</tr>
<tr>
<td>212.5</td>
<td>1903</td>
<td>8.14</td>
<td>0.939</td>
<td>0.021</td>
<td>0.094</td>
<td>0.051</td>
<td>0.689</td>
<td>0.145</td>
</tr>
<tr>
<td>255</td>
<td>1824</td>
<td>6.15</td>
<td>0.870</td>
<td>0.076</td>
<td>0.079</td>
<td>0.039</td>
<td>0.677</td>
<td>0.129</td>
</tr>
<tr>
<td>297.5</td>
<td>1541</td>
<td>4.82</td>
<td>0.761</td>
<td>0.093</td>
<td>0.075</td>
<td>0.031</td>
<td>0.688</td>
<td>0.113</td>
</tr>
<tr>
<td>340</td>
<td>1271</td>
<td>4.35</td>
<td>0.557</td>
<td>0.122</td>
<td>0.066</td>
<td>0.016</td>
<td>0.699</td>
<td>0.097</td>
</tr>
<tr>
<td>382.5</td>
<td>1101</td>
<td>4.21</td>
<td>0.484</td>
<td>0.136</td>
<td>0.057</td>
<td>0.013</td>
<td>0.705</td>
<td>0.0895</td>
</tr>
<tr>
<td>467.5</td>
<td>887</td>
<td>3.90</td>
<td>0.348</td>
<td>0.163</td>
<td>0.038</td>
<td>0.007</td>
<td>0.717</td>
<td>0.075</td>
</tr>
<tr>
<td>510</td>
<td>779</td>
<td>2.93</td>
<td>0.272</td>
<td>0.176</td>
<td>0.029</td>
<td>0.004</td>
<td>0.724</td>
<td>0.067</td>
</tr>
</tbody>
</table>
Figure 5. Slurry drop life-history at x/d = 297.5.
Figure 6. Quasisteady reaction rates for a noncatalyzed agglomerate.
In Fig. 6, predictions have been matched to the measurements at $x/d = 382.5$, by selecting a reactivity factor of $a_i = 58.2$ for all the reactions. This parameter was then held constant for the remaining computations, a procedure which yields a reasonably good correlation of the remaining measurements, including both effects of particle size (in the range 400-730 μm) and position in the flame. The low reaction rates indicative of the evaporation region were also predicted reasonably well. Thus the model demonstrates some potential for predicting quenching of agglomerate reaction.

There are two main combustion limits for carbon agglomerates. The first limit involves reaction rate control, where reaction rates are relatively low in comparison to diffusion rates, and the concentrations of gaseous reactants at the surface of the particle are nearly the same as in the surroundings. This regime is characterized by the burning rate being relatively independent of particle diameter, e.g., results for $x/d > 382.5$ in Fig. 6 approach the reaction rate limit.

The second limit involves diffusion control, where reaction rates are high in comparison to diffusion rates, and the concentrations of reactant gases at the surface of the particle are small. The variation of the burning rate is influenced by particle diameter in this region, showing a dependence

$$-\frac{dr}{dt} \sim 1/r_p \text{ (0.5 to 1.0)}$$

(41)

The power of $r_p$ in Eq. (41), tends to increase as the Reynolds number of the particle decreases. From this argument, it is apparent that the higher reaction rates illustrated in Fig. 6 ($x/d = 297.5, 340.0$) are approaching the diffusion controlled limit. Diffusion rates become large for small particles; therefore, all regions of the flow would eventually exhibit kinetic control for sufficiently small particles.

Figure 7 is an illustration of predicted and measured burning rates for a catalyzed agglomerate. These test results were limited to $x/d = 297.5, 340.0$ and 382.5. Reaction of the agglomerate is nearly diffusion controlled at the first two positions. Therefore, burning rates with and without catalyst are essentially the same at these locations. At $x/d = 382.5$, however, the burning rate is more nearly reaction rate controlled. At this position, the use of the catalyst results in a 20-25 percent increase in the burning rate. Naturally, the effect of the catalyst is even larger at positions farther downstream, e.g., adding catalyst reduces the lean limit for quenching from 0.37 to 0.21 [4].
Figure 7. Quasisteady reaction rates for catalyzed agglomerates.
The presence of catalyst is treated in the theory by adjusting the area/reactivity factor. Selecting $a_r = 70.6$, for all the reactions, yields the theoretical results illustrated in Fig. 7. With this choice, the predictions are in reasonably good agreement with the measurements.

Another property of catalyzed slurries was their tendency to spontaneously shatter from time to time. This shattering occurred after all the liquid had apparently evaporated. This behavior could be influenced by the presence of the thermocouple support, and was not consistently observed. If shattering does persist for small unsupported particles, this would provide another (physical) mechanism for the improved combustion properties observed for catalyzed slurries [3]. Tests with unsupported particles having sizes more typical of sprays will be required to resolve this question.

4.3 Agglomerate-Life-Histories

Agglomerate-life-histories were computed at $x/d = 255-510$, for noncatalyzed slurries. The resulting comparison between predicted and measured particle diameters and temperatures as a function of time are illustrated in Figs. 8-13. For reference purposes, the local temperature of the flow is also shown on each plot.

The initial condition for the computations illustrated in Figs. 8-13, was the temperature and diameter of the agglomerate, when all the liquid had evaporated. The comparison between predictions and measurements is seen to be quite good. This is partly expected, since the model was capable of correlating the burning rate measurements for this range of flame conditions. However, the results show that the model gives a good representation of the heat-up of the particle as well, where reaction effects are small, for various particle sizes and ambient temperatures, compositions and velocities, c.f. Table 5. At higher temperature levels, heat-up is also influenced by reaction and radiation to the surroundings. It is promising that the model appears to be capable of treating these effects as well.

The range of conditions illustrated in Figs. 8-13 is such that particle heat-up is terminated at temperatures both above and below the local gas temperature. This behavior is due to the opposing effects of reaction, tending to increase particle temperatures above the gas temperature, and radiation, which tends to reduce the particle temperatures below the gas temperature. In all cases, however, convection would dominate the process as the particle becomes small and the particle would finally tend to approach the gas temperature. This is not observed in Figs. 8-13 due to the finite-sized thermocouple bead used to mount the slurry drop.

4.4 Limitations of the Study

While the model shows an encouraging capability to predict the combustion properties of carbon agglomerates, several limitations of the present evaluation should be recognized. The main limitation
Figure 8. Agglomerate-life-history at x/d = 255.0.
Figure 9. Agglomerate-life-history at $x/d = 297.5$. 

$\phi = 0.761$
Figure 10. Agglomerate-life-history at x/d = 340.0.
Figure 11. Agglomerate-life-history at x/d = 382.5.
Figure 12. Agglomerate-life-history at $x/d = 467.5$. 

$D_{avg} = 467.5, \varphi = 0.348$
Figure 13. Agglomerate-life-history at \( x/d = 510.0 \).
of this study is the large particle sizes examined, 400-1000 \( \mu \text{m} \), in comparison to the sizes expected in practical combustors. The major difficulty involves the present specification of area/reactivity multiplication factors—which were taken to be constant during the present study. For large particles, diffusion of reactant gases is not expected to penetrate very far into the particle, while very small particles would have reactant gases penetrated throughout their volume. Therefore, the relative proportions of apparent surface to actual reaction areas will tend to increase as particle sizes become small. Therefore, it is expected that the area/reactivity multiplication factor would tend to increase as particle sizes are reduced. The magnitude of this change can only be established by additional study. Naturally, this effect would only be important outside the diffusion controlled limit.

Present test conditions were not sufficiently varied to establish the appropriate area/reactivity multiplication factors for the carbon dioxide and water vapor reactions, since oxygen dominated carbon reaction when the basic reaction parameters of Table 5 are used in the analysis. It is unlikely that the effect of pores, surface properties or catalyst would act the same for each reactant. Therefore, use of the present findings should be limited to conditions where reaction of carbon with oxygen dominates the process. Possible interference effects between the various reactants, and whether all the appropriate reactants were considered also remains to be resolved.

Another variable of significant practical importance is the effect of pressure. The present results were limited to atmospheric pressure and whether the reaction expressions used here yield the proper pressure dependence, must still be evaluated.
Appendix A

Computation of Properties

A.1 Mean Property State

The average properties of the gas surrounding the droplet were computed at the mean state, defined as follows:

$$\theta_{\text{avg}} = \alpha \theta_p + (1-\alpha) \theta_\infty$$  \hspace{1cm} (A-1)

where $\theta$ is a generic property at a specific temperature and mass fraction. $\alpha$ can range between 0 and 1 and a value of 0.9 was selected for the present investigation as suggested by Faeth and coworkers [9,10].

A.2 Mixture Thermal Conductivity

The thermal conductivity of the gaseous mixture was calculated with the Mason and Saxena formulation of the Wassiljewa equation [18].

$$\lambda_m = \frac{\lambda_i}{N} \left[ \frac{x_i \lambda_i}{\sum_{j=1}^{N} x_j \phi_{ij}} \right]$$  \hspace{1cm} (A-2)

where

$$\phi_{ij} = \frac{1}{8^{1/2}} \left( 1 + \frac{M_i}{M_j} \right)^{-1/2} \left[ 1 + \frac{\mu_i}{\mu_j} \left( \frac{M_i}{M_j} \right) \right]^{1/4}$$  \hspace{1cm} (A-3)

The thermal conductivity and viscosity for gases other than JP-10 was obtained from Ref. 19. The thermal conductivity of JP-10 was determined using the Miscic and Thodos method [18] and the viscosity of JP-10 was determined using the Reichenberg method [18]. Equation (A-2) was evaluated at the average temperature and composition of the gas phase. In Eq. (A-3), $\phi_{ij}$ is set equal to 1, when $i = j$. 

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A.3 Mixture Viscosity

The viscosity calculations for the mixture employed the method of Wilke, cited in Ref. 18.

For a mixture of $N$ components

\[ \mu_m = \frac{1}{N} \sum_{i=1}^{N} \frac{x_i \mu_i}{\sum_{j=1}^{N} x_j \phi_{ij}} \]  

(A-4)

where $\phi_{ij}$ is given by Eq. (A-3). The viscosity data used in the calculations was obtained in the same manner as described in Section A.2.

A.4 Mixture Diffusivity

When all of the liquid fuel has evaporated and only a carbon agglomerated particle remains the diffusion coefficient of the gas was taken for simplicity as the binary diffusion coefficient of oxygen through nitrogen

\[ D = D_{O_2,N_2} \]  

(A-5)

Use of other species combinations yielded diffusivities of similar magnitude.
References


References (Continued)


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