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Aerothermodynamic Analysis of Stardust Sample Return Capsule with Coupled Radiation and Ablation

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An aerothermodynamic analysis of the forebody aeroshell of the Stardust Sample Return Capsule is carried out by using the axisymmetric viscous shock-layer equations with and without fully coupled radiation and ablation. Formulation of the viscous shock-layer equations with shoulder radius as the length scale and implementation of the Vigneron pressure condition allow resolution of the flowfield over the shoulder. With a predominantly supersonic outflow over the shoulder, a globally iterated solution of viscous shock-layer equations can be obtained. The stagnation-point results are obtained along a specified trajectory, whereas detailed calculations along the body are provided at the peak-heating point. The equilibrium calculations with ablation injection are the focus of the present study because of the lack of a general chemical nonequilibrium analysis that accounts for both surface and flowfield effect. The equilibrium calculations also provide a simple way to conserve surface (and flowfield) elemental composition for the current small ablation injection rates, where the surface elemental composition is a mixture of freestream and ablator elements. Therefore, the coupled laminar and turbulent flow solutions with radiation and ablation are obtained by using the equilibrium flow chemistry, whereas a nonequilibrium chemistry model is used for solutions without ablation and turbulence. Various computed results are compared with those obtained by the other researchers.

**Nomenclature**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_i$</td>
<td>mass fraction of species $i$</td>
</tr>
<tr>
<td>$C_k$</td>
<td>mass fraction of element $k$</td>
</tr>
<tr>
<td>$C_p$</td>
<td>frozen specific heat of mixture, J/kg-K</td>
</tr>
<tr>
<td>$D_{ij}$</td>
<td>binary diffusion coefficient, m$^2$/s</td>
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<tr>
<td>$h_a$</td>
<td>enthalpy of undecomposed ablation material, J/kg</td>
</tr>
<tr>
<td>$h_i$</td>
<td>enthalpy of species $i$, J/kg</td>
</tr>
<tr>
<td>$h_i$</td>
<td>enthalpy of species $i$, $h_i/V_0^2$</td>
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<tr>
<td>$K$</td>
<td>thermal conductivity of mixture, W/m-K</td>
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<tr>
<td>$K$</td>
<td>thermal conductivity of mixture, $K/\mu_{ref}C_p_\infty$</td>
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<td>$L_e$</td>
<td>Lewis number, $\rho D_{ij}C_p/K$</td>
</tr>
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<td>$m$</td>
<td>mass injection rate, kg/m$^2$-s</td>
</tr>
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<td>coordinate measured normal to body, m</td>
</tr>
<tr>
<td>$n$</td>
<td>coordinate measured normal to body, $n/R_C1$</td>
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<tr>
<td>$n_{sh}$</td>
<td>shock standoff distance, m</td>
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<tr>
<td>$Pr$</td>
<td>Prandtl number, $\mu C_p/K$</td>
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<td>pressure, N/m$^2$</td>
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<td>heat flux, W/m$^2$</td>
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<td>$g_{\text{cond}} + q_{\text{conv}} + q_{\text{abl}}$, W/m$^2$</td>
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<td>component of radiative flux toward shock, W/m$^2$</td>
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<td>nose radius, m</td>
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<td>$T_{\text{REW}}$</td>
<td>radiative equilibrium wall temperature, K</td>
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<tr>
<td>$T_{\text{ref}}$</td>
<td>reference temperature, $V_\infty^3/C_p_\infty$, K</td>
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$T_{\text{sub}}$ = sublimation temperature, K
$l$ = time, s
$V_\infty$ = freestream velocity, m/s
$\alpha_{1,j}$, $\alpha_{2,j}$, $\alpha_{3,j}$ = curve-fit coefficients for sublimation temperature, $T_{\text{sub}}$
$\beta_{1,j}$, $\beta_{2,j}$, $\beta_{3,j}$ = curve-fit coefficients for heat of ablation, $\Delta H_a$
$\Delta H_a$ = heat of ablation, MJ/kg
$\varepsilon$ = char emmissivity, 0.9
$\tilde{\varepsilon}$ = Reynolds-number parameter, $(\mu_{\text{rel}}/\rho_0 V_\infty R_C1)^{1/2}$
$\mu$ = viscosity of mixture, N-s/m$^2$
$\mu_{\text{rel}}$ = reference viscosity, $\mu(T_{\text{ref}})$, N-s/m$^2$
$\rho$ = density of mixture, kg/m$^3$
$\sigma$ = Stefan–Boltzmann constant, 5.668 x 10$^{-8}$ W/(m$^2$·K$^4$)

**Subscripts**

$A$ = ablator
$\text{abl}$ = ablation
$\text{cond}$ = conduction
$\text{conv}$ = convection
$\text{diff}$ = diffusion
$\text{Eq}$ = equilibrium
$i$ = $i$th species
$j$ = $j$th species
$k$ = $k$th element
$r$ = radiation
$w$ = wall value
$\omega$ = values for solid ablation material at surface
$\infty$ = freestream value

**Introduction**

The Stardust mission, part of NASA's Discovery Program, plans to fly a spacecraft through the tail of the comet Wild-2 and bring samples of cometary material as well as interstellar dust to Earth for analysis. The collected cometary particles and the dust will be contained in the Stardust Sample Return Capsule (SRC), which must survive an intense Earth entry heating. At 12.6 km/s the SRC entry is the fastest ever attempted into the Earth's atmosphere. This paper focuses on the aerothermodynamic issues concerning the flow environment around the SRC forebody during such an entry. Since Stardust was scheduled for launch in early 1999, the work presented here was not available in time to impact the SRC design.
However, a better understanding of the SRC entry environment and the computational tools employed for its analysis will help in promoting a better and more efficient design of the thermal protection system (TPS) for future sample return vehicles such as MUSES-C, Genesis, Champollion (DS-4), and Mars Sample Return. Improved computational tools will also be useful in postflight evaluation of the TPS and other measured quantities.

Previous Work

Recently, calculations have been done\(^2\) for the Stardust SRC forebody TPS with an axisymmetric Navier-Stokes flow solver, loosely coupled to the radiation and ablation modules. The thermochemical nonequilibrium flowfield calculations with ablation are based on an 18-species chemical model. The ablation boundary condition (namely, the blowing rate, species mass fractions, and wall temperature) for the flowfield solution are obtained iteratively by assuming the surface composition to be in equilibrium at the temperatures and pressures predicted from a material response code (with inputs of wall heat transfer rate and pressure from the flowfield solution).

Employing a methodology similar to that of Ref. 2, Ref. 3 has recently obtained stagnation-point heat transfer rates for the Pioneer-Venus probes,\(^4\) whose flight environment resembles that of current sample return vehicles.\(^2\)\(^5\)\(^6\) A 17-species nonequilibrium chemistry model is used for the shock-layer flow. The pyrolysis gas composition at the surface is obtained by assuming that the surface is in equilibrium at the local temperature and pressure.

A recent analysis\(^7\) of the MUSES-C asteroid sample return mission has considered a 19-species nonequilibrium chemistry model (with the thermal equilibrium assumption) both for the shock-layer flow and the ablating surface for a true ablation calculation. The chemistry model consists of 11 air species and 8 carbon-containing species. The hydrocarbon species are not included, and the pyrolysis process is not considered to keep the analysis simple. Because the 19-species finite rate chemistry model is implemented both at the surface and in the shock layer, it includes all of the species considered throughout the computational domain. Thus, the analysis is consistent both at the surface and through the shock layer for the chemistry model considered.

Present Work

In the present work the Stardust SRC entry flowfield is investigated by assuming complete thermal equilibrium. Fully coupled radiation solutions with and without ablation injection are obtained by using an axisymmetric viscous shock-layer method by assuming chemical equilibrium both in the flowfield and at the surface. The element continuity equations are solved iteratively for each element to determine the appropriate mix of ablative and freestream elemental composition at the surface (adjacent to the ablator) and through the flowfield. With information concerning the elemental composition, pressure, and temperature, the species concentration at the surface (adjacent to the ablator) and through the flowfield is obtained by using the free energy minimization procedure.

With the calculation of both the flowfield and surface ablation species from the equilibrium assumption, the transition from ablation to freestream elements (and the corresponding species) is obtained continuously\(^8\) through the solution of elemental continuity equations. Further, the fully equilibrium calculations also bypass the entire discussion about governing processes and intermediate steps concerning the number of species, reaction mechanisms, and the associated reaction rates, especially for the complex flowfields with ablation. Therefore, the flowfield calculations with the thermochemical equilibrium model would appear adequate for the TPS design (for the convective-dominated heating environment), especially if the ablation boundary condition is specified with the equilibrium assumption. Although a nonequilibrium overall solution including surface and flowfield would be desirable, a full equilibrium calculation further provides a conservative estimate of the surface heating with the desired safety factor for the TPS design.

Computed results for an ablating surface include those with the laminar flow assumption as well as those for a fully turbulent flow immediately downstream of the stagnation line. For a nonablating surface results have been obtained with nonequilibrium chemistry and fully catalytic\(^2\) as well as equilibrium catalytic wall boundary condition. Only at lower temperatures (i.e., \(T < 2000\) K) would a fully catalytic boundary condition (with complete recombination to freestream value) be realistic to use.

Analysis

Flowfield Model

The viscous shock-layer (VSL) equations employed are those of a multicomponent reacting-gas mixture under conditions of chemical nonequilibrium\(^9\) and equilibrium\(^8\)\(^10\) with thermal equilibrium. These equations are the same as those given in Refs. 9 and 10, and, therefore, they are not given here. The chemistry model, boundary conditions, and thermodynamic and transport properties employed are similar to those of Refs. 9–11, whereas the ablation injection, radiative transport, and turbulence models (used with equilibrium chemistry only) are those of Refs. 8 and 10–13. These models, boundary conditions, and the properties are briefly described here.

Chemistry Model

For calculations of airflow over a nonablating surface, an 11-species (\(N_2, O_2, N, O, NO, NO^+, e^-, N^+, O^+\)) chemical model is used for nonequilibrium calculations, whereas a 9-species (\(N_2, O_2, N, O, NO, NO^+, e^-, N^+, O^+\)) chemical model is used for the equilibrium flow. For the equilibrium ablation injection calculation, 20 chemical species are used: the 7 equilibrium air species (without \(NO^+\) and \(NO^+\)) plus \(C, C_2, C_3, CO, CN, C_2H, C_2H_2, C_2H_3, C^+\), \(H, H_2\), and \(HCN\). The equilibrium composition is determined (for a given temperature, pressure, and elemental composition) by using the free-energy minimization method of Ref. 14.

Boundary Conditions

The boundary conditions at the shock are obtained by using the Rankine–Hugoniot relations. The flow behind the shock is assumed to be in chemical equilibrium or frozen at the freestream composition for equilibrium and nonequilibrium calculations, respectively.\(^9\) No-slip continuum boundary conditions are employed at the surface. The surface temperature with no ablation injection is assumed to be the radiative equilibrium wall value obtained from

\[
T_{REW} = \left[ \frac{\partial}{\partial t} \left( T \right) \right]^{-1} \left( \sum_{i=1}^{NS} \frac{h_i}{\theta_i} \right) \frac{\rho_w V_w^2}{\varepsilon} \left( \frac{1}{c_p} \right)
\]

(1)

For ablation injection cases steady-state ablation is assumed. However, the species surface concentrations, ablation rates, and surface temperatures, in general, can be obtained from a material response code (such as FIAT of Ref. 2), by employing input surface heat flux and pressure from an equilibrium flowfield code. For the surface ablation cases considered in the present study, an energy balance at the flowfield-ablator interface gives the coupled mass injection rate for quasi-steady ablation:

\[
m = \frac{-q_{w} - q_{w}}{\sum_{i=1}^{NS} (h_i - h_a)}
\]

(2)

The surface temperature for the present calculations with ablation injection is that at which the quasi-steady ablation occurs. The expression used for surface temperature for the Phenolic Impregnated Ceramic Ablator (PICA)\(^15\) has been obtained by curve fitting these values (computed from the charring material and ablation thermal\(^16\)\(^17\) response code) in the pressure range 0.001 atm < \(p < 1.00\) atm. The elemental composition of PICA is similar to that of a carbon-phenolic ablator. It is, however, less dense and has much lower thermal conductivity. The expressions for the sublimation temperature and heat of ablation for PICA (with 92% carbon, 4.9% oxygen, 2.2% hydrogen, and 0.9% nitrogen by mass) are

\[
T_{abl} = \sum_{j=1}^{5} \alpha_{abl,j} \left( \frac{\log C_a}{\log C_x} \right)^{j-1} + \left( \log p_a \right) \sum_{j=1}^{5} \alpha_{abl,j} \left( \frac{\log C_a}{\log C_x} \right)^{j-1}
\]

(3)
where $p_w$ is the wall pressure in atmospheres and $C_A$ is the ablator mass fraction at the surface. The values for $\alpha_{1,j}$ are given in Table 1. For the case when the gas species adjacent to the surface are caused solely by the ablation species (i.e., $C_A = 1.0$), Eq. (3) with the values of $\alpha_{1,j}$ in Table 1 gives

$$T_{abl} = 3790.0 + 329.94(\log p_w) + 20.386(\log p_w)^2 \tag{4}$$

The surface temperature and the coupled mass injection rate are calculated by iterating the solution of the governing flowfield equations and the boundary conditions.

For ablation injection the elemental concentrations at the wall are governed by convection and diffusion and are obtained from the equation

$$\left( \frac{\partial C_i}{\partial t} \right)_w = - \frac{1}{\rho_\infty} \left( \frac{m}{\mu L} \right)_w \left[ (C_i)_w - (C_i)_\infty \right] = 0 \tag{5}$$

For the radiative transport calculations the bow shock is considered transparent, and the freestream is considered cold and transparent. Therefore, the precursor effects are neglected. Further, the body surface is assumed to be gray with a reflectivity of 0.1, emissivity of 0.9, and transmissivity of 0. The energy reradiated from the surface is included both in the radiation transport calculation as well in the surface energy balance [Eq. (2)]. The net radiative flux can be represented as

$$q_r = q_r^{(+) - q_r^{(-)} } \tag{6}$$

At the surface

$$q_r^{(+) - q_r^{(-)} } = \varepsilon \sigma T_w^4 \tag{7}$$

The heat transferred to the wall because of conduction, diffusion, and convection is

$$-q_{c,w} = \varepsilon^2 \rho_\infty V_\infty \left[ \tilde{K} \frac{\partial T}{\partial n} + \frac{\mu L e}{\rho_\infty} \sum_{i=1}^{N_b} \frac{\partial C_i}{\partial n} \right]$$

$$- \frac{m}{\rho_\infty V_\infty} \sum_{i=1}^{N_b} (C_i \tilde{h}_i - C_i \tilde{h}_i)_w \tag{8}$$

where $N_b$ is the number of species.

**Catalytic Wall Conditions**

For nonablating, nonequilibrium flow the following three catalytic wall boundary conditions are used.

1) **Noncatalytic wall (NCW):** Because no reactions occur at the surface in this case, the mass-fraction gradients for all species are zero at the surface, i.e.,

$$\left( \frac{\partial C_i}{\partial n} \right)_w = 0 \tag{9}$$

2) **Equilibrium catalytic wall (ECW):** The wall catalyzed reactions are assumed to occur at an infinite rate, and, therefore, the species mass fractions at the wall are those corresponding to their local equilibrium values, i.e.,

$$(C_i)_w = (C_i)_\text{eq} = f(p_w, T_w) \tag{10}$$

3) **Fully catalytic wall (FCW):** The gas species at the surface are assumed to recombine to the freestream composition, i.e.,

$$(C_i)_w = (C_i)_\infty \tag{11}$$

At low surface temperatures the surface condition of Eq. (10) reduces to Eq. (11).

**Radiative Transport**

The radiative transport code RADICAL\textsuperscript{18,19} has been used to compute radiative heat flux $q_r$. This code accounts for the effects of nongray self-absorption and includes the molecular band, continuum, and atomic line transitions. The ultraviolet properties for $C_3$ are taken from Ref. 20. The radiative transport is fully coupled with the flowfield solutions for equilibrium chemistry. The nonequilibrium total heat transfer rate (for a nonablating surface) is obtained by adding the equilibrium radiative component to the nonequilibrium conductive and diffusive components.

**Thermodynamics and Transport Properties**

Thermodynamic properties for specific heat, enthalpy and free energy, and transport properties for viscosity and thermal conductivity are required for each species considered. Values of these properties are obtained by using polynomial curve fits of Refs. 8 and 11. The equilibrium composition is determined by a free-energy minimization calculation procedure of Ref. 14. Mixture viscosity is obtained by the method of Armaly and Sutton,\textsuperscript{21} and mixture thermal conductivity is computed by the Mason and Saxena\textsuperscript{22} relation. Both constant and variable Lewis number values are employed in the computations. However, the results presented here are for a variable Lewis number based on the effective diffusion coefficient $D_w$ for the mixture.\textsuperscript{9}

**Turbulence Model**

A two-layer, eddy-viscosity, Cebeci–Smith turbulence model\textsuperscript{13,23,24} is employed in the present investigation. Reference 24 gives a detailed description of the model and various expressions for it. The boundary-layer edge definition used in the current study is based on an index of diffusion, conduction, and dissipation.\textsuperscript{25} The transition to turbulent flow is assumed to occur instantaneously at the first grid point downstream of the stagnation point. The turbulent Prandtl and Lewis numbers are assumed to be 0.9 and 1.0, respectively.

**Solution Procedure**

The method used to solve the nonequilibrium and equilibrium VSL equations is a spatial-marching, implicit, finite-difference technique,\textsuperscript{8,13} which includes coupling of the global continuity and normal momentum equations and use of the Vigneron pressure condition in the subsonic region (which covers a large part of the forebody of Stardust capsule shown in Fig. 1). The shoulder radius $R_{C1}$ is employed for the reference length in place of the conventionally used nose radius $R_N$. Details of the method of solution are similar to those of Refs. 8 and 13 and, therefore, are not presented here.

**Results and Discussion**

Results are presented for the forebody of the SRC shown in Fig. 1. The overshoot entry trajectory\textsuperscript{2} (which produces maximum heat loads) used in the calculations is given in Fig. 2. The freestream conditions at the calculation points for this trajectory are provided in Table 2. Peak heating and pressure occur at approximately 54 and 66 s, respectively, for this trajectory. The SRC forebody is an

**Table 1 Coefficients for ablation temperature for PICA**

<table>
<thead>
<tr>
<th>Coefficients</th>
<th>1</th>
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<tr>
<td>$\alpha_{1,j}$</td>
<td>3790.0</td>
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<td>$\alpha_{2,j}$</td>
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<td>$\alpha_{3,j}$</td>
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<td>-684.89</td>
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</tbody>
</table>

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**Fig. 1 Geometry of SRC; dimensions in millimeters.**
axisymmetric 60-deg sphere cone with a nose radius $R_{n}$ of approximately 23 cm. The computational domain for the present VSL results extends to just past the highest point C1 on the shoulder, where the flow is predominantly supersonic. The PICA heatshield for the forebody extends beyond this point. A 100 x 125 grid is employed with all of the computations. Variable grid sizes are used both normal to and along the body surface. The minimum distance between normal grid points is $(2 \times 10^{-4}) R_C$. In the direction along the surface, the minimum grid size is $(1 \times 10^{-4}) R_C$ on the shoulder to resolve the flowfield there and is as large as five times this value in the nose region to reduce the computational time. These values of the grid sizes have been established to ensure grid independence of the solution at peak heating condition ($t = 54$ s). Calculations were done by using the Cray C90 computer. The computational time required is about 200 and 600 CPU s per global pass for the nonequilibrium and equilibrium flow calculations around the body (without radiation), respectively. Typically, two global passes are required for convergence of the shock shape and surface heating.

**Nonablating Results Along Trajectory with Nonequilibrium and Equilibrium Chemistry**

Nonablating stagnation-point total heat transfer rate (conduction + diffusion + radiation) is provided at different times along the entry trajectory in Fig. 3. Results are obtained with equilibrium as well as nonequilibrium (finite rate) flowfield chemistry. Three surface recombination boundary conditions, namely, FCW, ECW, NCW are used with the finite rate chemistry calculations. Present results for the FCW case are in good agreement with those obtained by Olynick et al. up to time $t = 60$ s. A maximum value of about 1250 W/cm² is obtained at $t = 54$ s from the present calculations. The corresponding value obtained from Ref. 2 is about 2% higher. However, the difference between these two values increases for $t > 60$ s and may be caused by grid resolution employed in Ref. 2. Further, the present finite rate results with an ECW boundary condition are close to the equilibrium flow results as expected. A maximum value of about 1100 W/cm² is obtained at $t = 54$ s in this case. The NCW predictions are included for reference and give the lowest surface heating with a maximum value of only about 650 W/cm² at peak heating ($t = 54$ s). Figure 4 shows the presently computed stagnation-point equilibrium radiative heat transfer rate as well as the values obtained in Ref. 2 from a nonequilibrium radiation calculation. It is not clear why the nonequilibrium radiative heating is lower at earlier times and higher at later times in the trajectory as compared with the equilibrium calculations. Generally, the nonequilibrium effects (which are likely to be present at earlier times in the trajectory) should increase radiative heating in comparison with the equilibrium value, and it should approach the equilibrium value at later times in the trajectory (with the increasing Reynolds number).

The radiative equilibrium wall temperatures for the heating calculations of Fig. 3 are shown in Fig. 5. A value of 0.9 is used for char emissivity $ε$ in the present calculations. Similar to the surface heat transfer values, present FCW predictions for surface temperature are in good agreement with those of Olynick et al. Reference 2 employed a value of 1.0 for $ε$, with zero reflectivity assumed. Figure 5 also shows Olynick’s values adjusted for $ε = 0.9$. Differences between the present predictions and those of Ref. 2 increase at a later time in the trajectory for the reasons mentioned earlier. For most of the investigated trajectory, the surface temperatures are greater than 3000 K. Consequently, the FCW boundary condition is physically inappropriate because full recombination of air (for FCW boundary condition) cannot be forced for temperatures greater than about 2000 K. A physically appropriate surface recombination condition for these temperatures is a finite catalytic wall condition, which would be bounded by the ECW (most conservative) and the NCW boundary conditions. A maximum value of about 3800 K is obtained at $t = 54$ s for the present finite rate results with an ECW condition. These results are close to those obtained with the equilibrium flowfield chemistry as expected. The NCW calculations give the lowest surface temperatures as noted with the surface heating results of Fig. 3.
Ablation Results Along Trajectory with Equilibrium Chemistry

Figure 6a shows the equilibrium stagnation total heat transfer rate with and without ablation along the trajectory. Ablation produces a 35% reduction in the heating at peak-heating time of \( t = 54 \) s. The corresponding peak stagnation heat transfer rate of Ref. 2 (without ablation) is about 10% higher than the present value, and their results also show a reduction of about 35% with ablation. Components of the total heat transfer rates of Fig. 6a are shown in Fig. 6b. Reduction in the convective component \( q_c \) [which is given by Eq. (8)] and consists of conduction, diffusion, and convection] by ablation is caused by injection cooling. Ablation injection reduces the surface gradients of temperature and that of various species mass fractions; this causes a decrease in the conductive and diffusive heat fluxes. The radiative component, which is relatively small (less than 11% of the total heating without ablation), is not impacted much by ablation injection. There is a slight increase in radiation with ablation before the peak heating \( (t = 54 \) s). There is a deeper penetration of the shock layer by the ablation species C and CO during earlier times in the trajectory, and the increase in radiation from C line and \( \text{CO}(^4 \text{S}) \) molecular contributions is only partially offset by the absorption of radiation by ablation species during that period.

Surface temperatures used with the equilibrium stagnation heating calculations are given in Fig. 7. Surface heating without ablation is obtained by using the radiative equilibrium wall temperature as already mentioned (see Fig. 5), whereas the ablation temperature from Eq. (3) is used for the ablation injection calculations. The no ablation temperatures are generally higher (because of the higher surface heating) than those with ablation. Present ablation temperature values are close to those obtained by Olynick et al.\(^2\) up to the peak heating time \( (t < 54 \) s) in the trajectory. At later times the present surface temperature values are lower by a maximum of about 18% \( (t = 76 \) s). Also shown in Fig. 7 is the mass fraction of ablation species at surface, with a maximum value of about 0.25 at peak heating \( (t = 54 \) s). This value implies that 75% of the mass at the surface is from the freestream at that time in the trajectory.

The surface ablation injection rate along the trajectory as well as the ratio of injection rate to the freestream mass flux, corresponding to the heat transfer rate of Fig. 6a, are shown in Fig. 8. A maximum value of 3% of the ratio is obtained at time \( t = 34 \) s. The value of this ratio decreases to about 1.5% at peak heating, where the maximum injection rate of about 0.04 kg/m\(^2\)-s is obtained. Even though similar reduction (35%) in heating is obtained with ablation, presently computed values of the injection rate and injection mass flux ratio at peak heating \( (t = 54 \) s) are about one-half of those obtained in Ref. 2. These differences are believed to be caused by the differences in the mass fraction of ablation species and their enthalpies used in the two calculations. As pointed out in the next section, some of the species identified as significant in the present work are not included in the equilibrium surface analysis and flowfield chemical kinetics model of Ref. 2.

**Fig. 5** Nonablating stagnation-point temperature.

**Fig. 6a** Equilibrium stagnation-point total heat transfer rate.

**Fig. 6b** Components of equilibrium stagnation-point heat transfer rate.

**Fig. 7** Stagnation-point surface temperature and ablator mass fraction.
Laminar/Turbulent Ablation Results at Peak Heating Condition with Equilibrium Chemistry

Stagnation Profiles

Figure 9 shows the temperature and ablator mass-fraction profiles at peak heating ($t = 54$ s). The effect of ablation injection is limited to about 20% of the flowfield close to the surface at that time. This effect is also evident from the mass-fraction profiles of the freestream and ablation species shown in Fig. 10. The dominant ablation species not included in Ref. 2 are $C_2H$, $C_3H$, and $C_4H$. Noninclusion of these species can be partly responsible for the differences in present mass loss rate and those of Ref. 2 as mentioned earlier (see Fig. 8).

Forebody Surface Distributions

Surface distributions of the total heat flux for laminar and fully turbulent flow cases over the forebody of the Stardust Capsule are given in Fig. 11. This figure shows the effect of both ablator (PICA) mass injection and turbulence on total heating distributions for the peak-heating time of $t = 54$ s along the trajectory. The coupled PICA mass injection distributions are shown in Fig. 12. The impact of ablation injection on total heating is very pronounced for the laminar flow. In this case the total heating is reduced by about 35% along the forebody (compared with the nonablating surface), essentially through the reduction of the convective component as discussed earlier for the stagnation point. However, for the turbulent solutions, where the flow is assumed to undergo instantaneous transition at $s/R_N$ equal to 0.05, the reduction in total heating is less than 13% as compared with the nonablating laminar flow value on both the conical flank and shoulder. Obviously the benefit of ablation injection in reducing the heating for the laminar flow is partially negated when the flow is assumed to be turbulent. The mass injection rate distributions of Fig. 12, in general, follow the surface heat-flux distributions of Fig. 11. The stagnation nondimensional injection rate $\dot{m}/\rho_\infty V_\infty$ of 0.015 corresponds to a dimensional value of about 0.040 kg/m$^2$s (as noted earlier).

There is no noticeable effect of ablation injection and turbulence on surface pressure distribution as shown in Fig. 13. The stagnation...
These equations are globally iterated with the Vigneron pressure shoulder radius in place of the conventionally employed nose radius. For proper resolution of turbulence.

VSL analysis with and without surface ablation including the effect of the Earth’s atmosphere. Solutions are obtained from an axisymmetric formulation the species boundary condition problem at the surface.

The forebody aeroshell consists of a 60-deg sphere cone with a maximum stagnation heating of about 1250 W/cm² is obtained without ablation injection with nonequilibrium calculations and complete surface recombination (i.e., FCW boundary condition), whereas a value of about 1100 W/cm² is obtained for a more realistic ECW boundary condition with a radiative equilibrium wall temperature of about 3800 K. Stagnation heating similar to the later value is obtained with a fully equilibrium calculation. The maximum value of radiative heating component is about 11% at peak heating. With ablation injection a decrease of about 35% in the total stagnation-point heating (with equilibrium chemistry) is obtained at the peak-heating point in the trajectory. Reduction in heating is slightly less downstream of the stagnation point and along the conical flank, including the shoulder for the laminar case.

For the turbulent solutions where the flow is assumed to undergo instantaneously transition just downstream of the stagnation line, the heating is reduced by only about 13% on the conical flank and shoulder as compared with the nonablating laminar flow. Augmentation of the convective heating by turbulence appears to partially negate the benefit of heating reduction caused by ablation injection in this case.

**Conclusion**

Results are presented for the forebody of the SRC entering the Earth’s atmosphere. Solutions are obtained from an axisymmetric VSL analysis with and without surface ablation including the effect of turbulence.

The forebody aeroshell consists of a 60-deg sphere cone with a shoulder radius one-twelfth that of the nose. For proper resolution of the flowfield over the shoulder, the VSL equations are scaled with the shoulder radius in place of the conventionally employed nose radius. These equations are globally iterated with the Vigneron pressure condition to treat the large embedded subsonic region between the stagnation line and the supersonic outflow at the top of the shoulder.

The no-ablation VSL calculations employ an 11-species nonequilibrium chemistry model. For these calculations an ECW boundary condition to treat the large embedded subsonic region between the stagnation line and the supersonic outflow at the top of the shoulder.

The fully coupled ablation injection calculations are done with a 20-species equilibrium chemistry model. With fully equilibrium calculations the elemental conservation equations are solved iteratively for each element to determine the elemental composition at the surface (adjacent to the ablator) and in the flowfield. In this formulation the species boundary condition problem at the surface encountered with finite rate calculations is avoided for the case when the ablation injection rate is small, and the gas composition at the surface is caused by both the freestream and ablation products. The small injection rates are usually encountered before large-scale oxidation and sublimation drive the species caused by the freestream away from the surface.

The maximum stagnation heating of about 1250 W/cm² is obtained without ablation injection with nonequilibrium calculations and complete surface recombination (i.e., FCW boundary condition), whereas a value of about 1100 W/cm² is obtained for a more realistic ECW boundary condition with a radiative equilibrium wall temperature of about 3800 K. Stagnation heating similar to the later value is obtained with a fully equilibrium calculation. The maximum value of radiative heating component is about 11% at peak heating. With ablation injection a decrease of about 35% in the total stagnation-point heating (with equilibrium chemistry) is obtained at the peak-heating point in the trajectory. Reduction in heating is slightly less downstream of the stagnation point and along the conical flank, including the shoulder for the laminar case.

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**References**


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