

NASA
TP
1227
c.1

NASA Technical Paper 1227

LOAN COPY: RETURN TO
AFWL TECHNICAL LIBRARY
KIRTLAND AFB, N. M.



A Numerical Solution
of the Navier-Stokes Equations
for Chemically Nonequilibrium,
Merged Stagnation Shock Layers
on Spheres and Two-Dimensional
Cylinders in Air

Kenneth D. Johnston and William L. Hendricks

MAY 1978

NASA



0134469

NASA Technical Paper 1227

A Numerical Solution of the Navier-Stokes Equations for Chemically Nonequilibrium, Merged Stagnation Shock Layers on Spheres and Two-Dimensional Cylinders in Air

Kenneth D. Johnston

*George C. Marshall Space Flight Center
Marshall Space Flight Center, Alabama*

and

William L. Hendricks

*Lockheed Huntsville Research and Engineering Center
Huntsville, Alabama*



National Aeronautics
and Space Administration

**Scientific and Technical
Information Office**

1978

ACKNOWLEDGMENT

The authors appreciate the assistance of Dr. A. C. Jain who was a Senior Resident Research Associate at Marshall Space Flight Center in 1973-74. Dr. Jain stimulated interest in this problem and provided the computer program which he and his colleagues had formulated at the Indian Institute of Technology. Also, Mr. Larry Donehoo of Marshall Space Flight Center provided an interpolation computer subroutine which was used in the viscosity computations.

TABLE OF CONTENTS

	Page
SUMMARY	1
I. INTRODUCTION.....	1
II. ANALYSIS	2
A. Formulation of the Problem	2
B. Method of Solution.....	26
C. Computer Program	27
III. RESULTS	27
A. Effect of Computer Program Modifications	27
B. Comparison of Data Between Old Program and Modified Program	28
C. Comparison with Experimental Data.....	29
D. General Results	33
APPENDIX A — COMPUTER PROGRAM	59
APPENDIX B — LISTING OF PROGRAM	73
REFERENCES	90

LIST OF ILLUSTRATIONS

Figure	Title	Page
1.	Coordinate systems	4
2.	Comparison of present theory with experimental data for stagnation point heating on a sphere	30
3.	Convergence of temperature profiles for sphere with increasing number of iterations	34
4.	Typical convergence of stagnation point heat transfer coefficient for sphere with increasing number of iterations	35
5.	Flow profiles for sphere with noncatalytic wall at large Reynolds number	36
6.	Flow profiles for sphere with noncatalytic wall at small Reynolds number	38
7.	Flow profiles for sphere with fully catalytic wall	40
8.	Comparison of flow profiles for sphere and cylinder	43
9.	Temperature profiles at various Reynolds numbers for sphere	44
10.	Effect of variation in altitude on dissociation and viscosity in shock layer of sphere	45
11.	Effect of freestream speed on dissociation and viscosity in shock layer of sphere	46
12.	Stagnation point heat transfer coefficients for spheres and cylinders as a function of altitude	47
13.	Stagnation point heat transfer coefficient for sphere as function of freestream speed	49

LIST OF ILLUSTRATIONS (Concluded)

Figure	Title	Page
14.	Stagnation point heat transfer coefficient for sphere as function of wall temperature	50
15.	Stagnation point heat transfer coefficient for sphere as function of K^2	51
16.	Shock layer thickness for sphere and cylinder	53
17.	Slip speed at body surface for sphere and cylinder	54
18.	Temperature of air at wall for sphere and cylinder	55
19.	Space Shuttle external tank stagnation point heating rate	56
A-1.	Location of computation points	71

LIST OF TABLES

Table	Title	Page
1.	Data Comparison	28
2.	Arc-Jet Flow Field Properties	32

DEFINITION OF SYMBOLS

<u>Symbol</u>	<u>Definition</u>
\bar{C}_p	specific heat at constant pressure
\bar{C}_v	specific heat at constant volume
C_H	heat transfer coefficient, $\bar{q}_w / \frac{1}{2} \bar{\rho}_\infty \bar{V}_\infty^3$
D_{ij}	binary diffusion coefficient for species pair i and j
\bar{e}_i	specific internal energy of species i
\vec{f}_i	body force on species i per unit mass of species i
\bar{h}	specific enthalpy of mixture
\bar{h}_i	specific enthalpy of species i
\bar{k}	coefficient of thermal conductivity
k_{fj}	forward reaction rate for jth reaction
k_{rj}	reverse reaction rate for jth reaction
K_{n_∞}	freestream Knudsen number, $\bar{\lambda}_\infty / \bar{r}_b$
K^2	hypersonic similarity parameter [see equation (62)]
Le_{ij}	Lewis number of species pair i and j, $\bar{\rho} D_{ij} \bar{C}_p / \bar{k}$
M_∞	freestream Mach number
M	indicates a catalyst
NS	number of species in gas mixture
n	dimensionless radial distance from the body, $(\bar{r} - \bar{r}_b) / \bar{r}_b$
n_∞	dimensionless radial distance from body to freestream edge (shock layer thickness)

DEFINITION OF SYMBOLS (Continued)

<u>Symbol</u>	<u>Definition</u>
\bar{P}	pressure
Pr	Prandtl number, $\bar{C}_p \bar{\mu}/\bar{k}$
\vec{q}	heat flux vector
\bar{q}_w	heat flux to wall
\bar{r}	radial distance from body center (see Fig. 1)
\bar{r}_b	radius of body
\bar{R}_i	gas constant of species i
\bar{R}	gas constant of mixture
\bar{R}	universal gas constant
$Re_{0\infty}$	freestream stagnation Reynolds number, $\bar{\rho}_{\infty} \bar{V}_{\infty} \bar{r}_b / \bar{\mu}_{0\infty}$
\bar{s}_{ij}	collision cross section for particle i with particle j
Sc_{ij}	Schmidt number for species pair i and j, $\bar{\mu}/\bar{\rho} \bar{D}_{ij}$
\bar{T}	temperature
$\bar{T}_{0\infty}$	freestream stagnation temperature
\bar{t}	time
\bar{u}	velocity component parallel to body surface (see Fig. 1)
\bar{v}	velocity component normal to body surface (see Fig. 1)
\bar{v}	specific volume of gas mixture
\vec{v}	velocity vector

DEFINITION OF SYMBOLS (Continued)

<u>Symbol</u>	<u>Definition</u>
\bar{V}_∞	freestream speed
\vec{v}_i	diffusion velocity vector of species i
w_i	molecular weight of species i
W	equivalent molecular weight of mixture
\dot{w}_i	net mass production rate of species i per unit volume
Y_i	mass fraction of species i
γ	ratio of specific heats, C_p/C_v
γ_i	recombination coefficient for species i
η	dimensionless radial distance from body surface [see Fig. 1 and equation (24)]
Θ_{vi}	characteristic temperature for vibration of diatomic species i
θ	circumferential angle (see Fig. 1)
$\bar{\lambda}_i$	mean free path of species i
$\bar{\lambda}_\infty$	freestream mean free path
$\bar{\mu}$	coefficient of absolute viscosity
$\bar{\mu}_{SU}$	Sutherland coefficient of absolute viscosity
$\bar{\mu}_{0\infty}$	coefficient of absolute viscosity at $\bar{T}_{0\infty}$
$\bar{\rho}$	density of mixture
σ	molecule-surface accommodation coefficient

DEFINITION OF SYMBOLS (Concluded)

<u>Symbol</u>	<u>Definition</u>
τ	viscous stress tensor
Φ	viscous dissipation function

Subscripts

i, j	species indices
0 (zero)	stagnation condition
s	edge of Knudsen layer, slip
sh	conditions behind normal shock
w	wall
∞	freestream property

Superscript

($\bar{}$)	dimensional quantity
--------------	----------------------

TECHNICAL PAPER

A NUMERICAL SOLUTION OF THE NAVIER-STOKES EQUATIONS FOR CHEMICALLY NONEQUILIBRIUM, MERGED STAGNATION SHOCK LAYERS ON SPHERES AND TWO-DIMENSIONAL CYLINDERS IN AIR

SUMMARY

The complete Navier-Stokes equations are solved along the stagnation streamline in merged stagnation shock layers on spheres and two-dimensional cylinders using an iterative finite-difference numerical procedure known as the accelerated successive replacement method. The fluid medium is chemically reacting air consisting of seven species. Velocity components, thermodynamic properties, species mass fractions, and wall heat transfer rates are computed. This report is intended as an explanation of the method and as a user's manual for the computer program.

I. INTRODUCTION

An aerospace vehicle ascending or descending through the Earth's atmosphere traverses several flow regimes from the continuum boundary layer regime at low altitudes, through the transitional regime at intermediate altitudes, to the free molecular regime at very high altitudes. The character of the flow field changes drastically from the boundary layer regime to the free molecular regime, and no single computational approach is valid throughout this range. The broad transitional regime may be divided into several sub-regimes as suggested by Hayes and Probstein [1]. Consider the typical spherical nose of an aerospace vehicle. In the boundary layer regime viscous effects are primarily confined to a thin boundary layer, the bow shock can be treated as a discontinuity, and a region of inviscid flow exists between the shock and the boundary layer. However with increasing altitude, the shock wave and boundary layer thicken and eventually merge into a single viscous layer called the shock layer. The flow regime in which this occurs is called the fully merged shock layer regime which is the condition treated in this report.

At the great speed that a vehicle reenters the atmosphere, the temperature near the body becomes extremely high, especially in the stagnation region. Therefore, the air in the shock layer dissociates and ionizes. For an accurate description of the flow field, one must account for these real gas effects. Also an accurate estimate of the ionization level is needed for radio communication purposes. Therefore, a flow field model including finite rate chemistry is required.

This report describes a method for computing flow properties along the stagnation streamlines of a sphere and a circular cylinder transverse to the flow. Heat transfer rates are computed at the body surface. Although this method is limited to the stagnation region, it still provides valuable design information because maximum heating rates usually occur at the nose. The computational method was developed by Jain and Adimurthy [2] for an ideal gas. The method uses the full Navier-Stokes equations to describe the flow in the entire shock layer from the surface to the freestream. The boundary conditions at the wall are provided by slip velocity and temperature jump equations. Using the concept of local similarity, the governing equations are reduced to a system of nonlinear, coupled ordinary differential equations. Numerical solutions are obtained for points on the stagnation streamline using an iterative finite-difference procedure known as the accelerated successive replacement method. The applicability of this approach and the failure of thin-layer theories for the merged shock layer regime is discussed in Reference 2. Nonequilibrium chemical reactions were included in this method by Kumar and Jain [3] using an air model with seven species and six reactions. Hendricks [4] developed surface slip velocity and temperature jump equations for a multi-component gas, including the effects of wall catalysis, to use with this model. Additional modifications have been made in this report, principally by including the two-dimensional cylindrical geometry and using a multi-component gas model to compute viscosity.

II. ANALYSIS

A. Formulation of the Problem

1. Approach. In the present analysis the full Navier-Stokes equations, with nonequilibrium chemistry, are solved through the merged stagnation shock layer from the freestream to the body. The slip conditions at the gas-wall interface include the effect of wall catalysis and a multicomponent, nonequilibrium gas flow.

The thin shock layer assumption is not made in the present analysis. The full Navier-Stokes equations with chemically-reacting, nonequilibrium air are solved through the merged shock and boundary layer. This allows the shock wave to develop within the computational domain. A seven species air model is used. The species considered are N₂, O₂, NO, N, O, NO⁺, and e⁻. For air dissociation and ionization, the rate expressions recommended by Wray [5] are adopted. Prandtl number, Pr, and Lewis number, Le_{ij}, are taken to be 0.75 and 1.4, respectively, for the cases computed in this report. The viscosity of dissociated and ionized air is obtained from a simple summation formula for a mixture of hard spherical molecules using Hansen's collision cross sections [6].

Solutions are obtained by using the local similarity concept to reduce the governing equations to a set of nonlinear, coupled, ordinary differential equations. This set of equations is integrated using a finite difference method known as the accelerated successive replacement method. It is important to note that the first order local similarity assumption is a good approximation near the stagnation streamline, at least for Re_{sh} ≥ 10 [7]. Reference 7 gives a thorough discussion of local similarity.

2. Governing Equations. The nonlinear, coupled ordinary differential equations governing the flow of a multicomponent gas near the stagnation streamlines of spheres and two-dimensional cylinders are presented. The coordinate system employed is shown in Figure 1a.

a. Basic Equations. The basic conservation equations and the ideal gas equation of state for a multicomponent, reacting gas mixture are as follows [8]:

Global Continuity (of all species):

$$\frac{\partial \bar{\rho}}{\partial \bar{t}} + \vec{\nabla} \cdot (\bar{\rho} \vec{v}) = 0 . \quad (1)$$

Species Continuity:

$$\bar{\rho} \frac{D Y_i}{D t} = \dot{w}_i - \vec{\nabla} \cdot (\bar{\rho} Y_i \vec{V}_i) \quad i = 1, \dots, NS , \quad (2)$$

where NS = number of species in mixture (no summation on repeated indices).

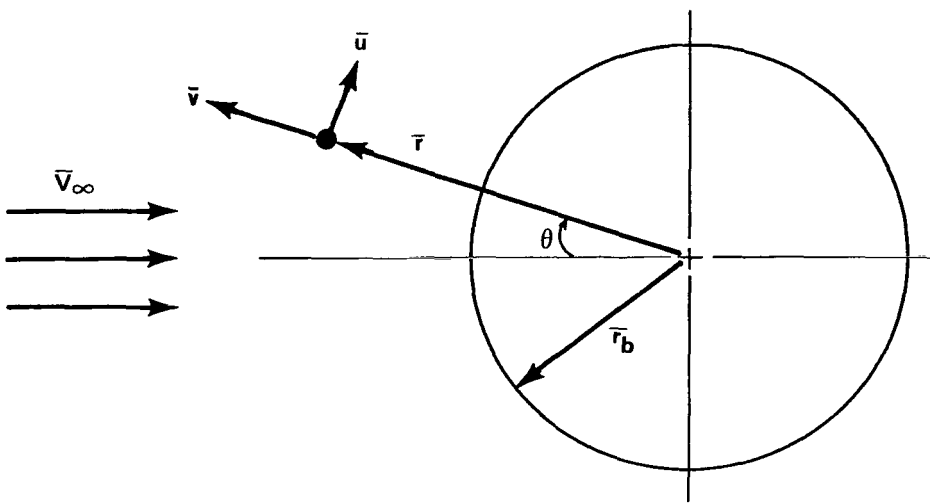


Figure 1a. Dimensional coordinate system.

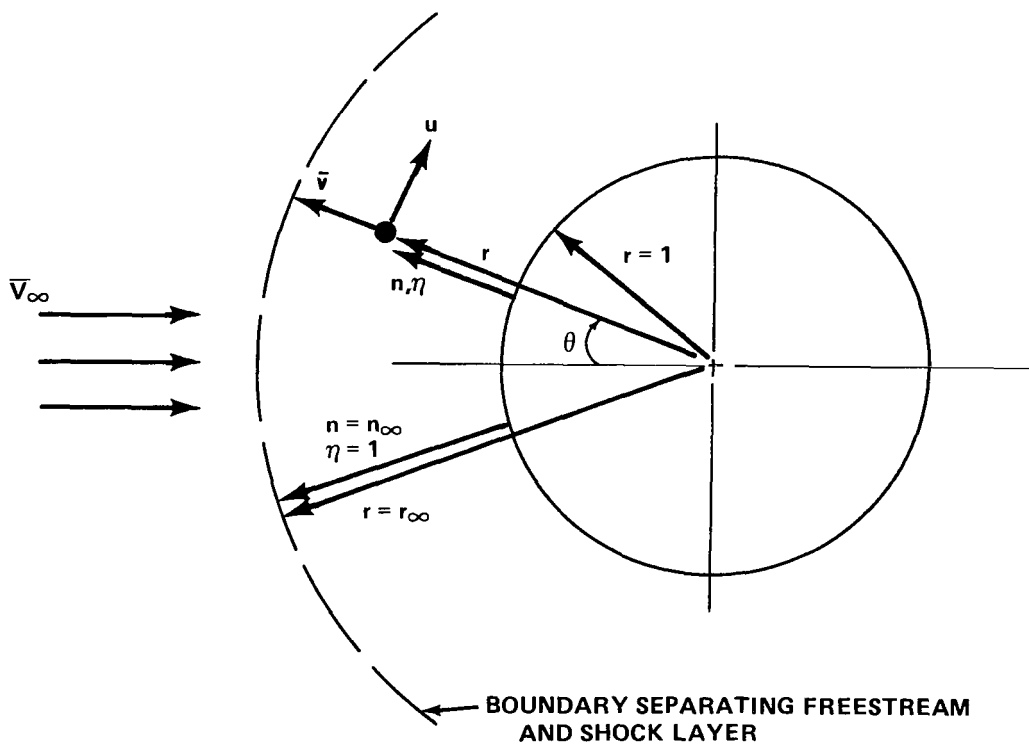


Figure 1b. Dimensionless coordinate system.

Figure 1. Coordinate systems.

Momentum:

$$\bar{\rho} \frac{D\vec{v}}{Dt} = -\vec{\nabla} \bar{P} - \vec{\nabla} \cdot \vec{\tau} + \bar{\rho} \sum_{i=1}^{NS} Y_i \vec{f}_i . \quad (3)$$

The second order tensor, $\vec{\tau}$, is the viscous stress tensor and \vec{f}_i is the body force per unit mass of species i .

Enthalpy:

$$\bar{\rho} \frac{D\vec{h}}{Dt} = \frac{D\bar{P}}{Dt} + \bar{\Phi} - \vec{\nabla} \cdot \vec{q} . \quad (4)$$

The quantity $\bar{\Phi}$ is the viscous dissipation function and \vec{q} is the heat flux vector.

State:

$$\bar{P} = \bar{\rho} \bar{\kappa} \bar{T} \sum_{i=1}^{NS} \frac{Y_i}{W_i} . \quad (5)$$

b. Nondimensional Equations. The basic equations are put in non-dimensional form by introducing dimensionless variables as follows:

$$u = \bar{u}/\bar{V}_{\infty} ,$$

$$v = \bar{v}/\bar{V}_{\infty} ,$$

$$\rho = \bar{\rho}/\bar{\rho}_{\infty} ,$$

$$T = \bar{T}/\bar{T}_{0\infty} ,$$

$$P = \bar{P}/\bar{\rho}_\infty \bar{V}_\infty^2 ,$$

$$h = \bar{h}/\bar{V}_\infty^2 ,$$

$$\mu = \bar{\mu}/\bar{\mu}_{0\infty} ,$$

$$r = \bar{r}/\bar{r}_b ,$$

$$\dot{w}_i = \dot{\bar{w}}_i / (\bar{\rho}_\infty \bar{V}_\infty / \bar{r}_b) ,$$

where \bar{r}_b is the radius of the body, $\bar{T}_{0\infty}$ is the freestream stagnation temperature, and $\bar{\mu}_{0\infty}$ is the coefficient of absolute viscosity evaluated at $\bar{T}_{0\infty}$. The nondimensional similarity parameters

$$Re_{0\infty} = \bar{\rho}_\infty \bar{V}_\infty \bar{r}_b / \bar{\mu}_{0\infty} ,$$

$$Pr = \bar{C}_p \bar{\mu} / \bar{k} ,$$

$$Sc_{ij} = \bar{\mu} / \bar{\rho} \bar{D}_{ij} ,$$

and

$$Le_{ij} = \bar{k} / \bar{\rho} \bar{C}_p \bar{D}_{ij}$$

are also introduced. The basic equations are then simplified by assuming steady flow and Newtonian fluid, neglecting body forces, viscous diffusion stresses, thermal radiation, and thermal diffusion, and using Fick's Law of Diffusion. The equations are presented in cylindrical and spherical coordinates below.

Cylindrical Coordinates:

Global Continuity:

$$\frac{\partial}{\partial r} (\rho rv) + \frac{\partial}{\partial \theta} (\rho u) = 0 \quad . \quad (6)$$

Species Continuity:

$$\rho \left(v \frac{\partial Y_i}{\partial r} + \frac{u}{r} \frac{\partial Y_i}{\partial \theta} \right) = \dot{w}_i + \frac{1}{Re_{0\infty}} \frac{1}{r} \left[\frac{\partial}{\partial r} \left(\frac{\mu r}{Sc} \frac{\partial Y_i}{\partial r} \right) + \frac{\partial}{\partial \theta} \left(\frac{\mu}{Sc r} \frac{\partial Y_i}{\partial \theta} \right) \right] . \\ i = 1, \dots, NS \quad (7)$$

Transverse Momentum (θ Direction):

$$\rho \left(v \frac{\partial u}{\partial r} + \frac{u}{r} \frac{\partial u}{\partial \theta} + \frac{vu}{r} \right) = - \frac{1}{r} \frac{\partial P}{\partial \theta} + \frac{1}{Re_{0\infty}} \left[\frac{1}{r^2} \frac{\partial}{\partial r} \left\{ r^2 \mu \left[r \frac{\partial}{\partial r} \left(\frac{u}{r} \right) + \frac{1}{r} \frac{\partial v}{\partial \theta} \right] \right\} \right. \\ \left. + \frac{1}{r} \frac{\partial}{\partial \theta} \left(\mu \left\{ 2 \left(\frac{1}{r} \frac{\partial u}{\partial \theta} + \frac{v}{r} \right) - \frac{2}{3} \left[\frac{1}{r} \frac{\partial}{\partial r} (rv) + \frac{1}{r} \frac{\partial u}{\partial \theta} \right] \right\} \right) \right] . \quad (8)$$

Radial Momentum (r Direction):

$$\rho \left(v \frac{\partial v}{\partial r} + \frac{u}{r} \frac{\partial v}{\partial \theta} - \frac{u^2}{r} \right) = - \frac{\partial P}{\partial r} + \frac{1}{Re_{0\infty}} \left[\frac{1}{r} \frac{\partial}{\partial r} \left(r \mu \left\{ 2 \frac{\partial v}{\partial r} - \frac{2}{3} \left[\frac{1}{r} \frac{\partial (rv)}{\partial r} + \frac{1}{r} \frac{\partial u}{\partial \theta} \right] \right\} \right) \right. \\ \left. + \frac{1}{r} \frac{\partial}{\partial \theta} \left\{ \mu \left[r \frac{\partial}{\partial r} \left(\frac{u}{r} \right) + \frac{1}{r} \frac{\partial v}{\partial \theta} \right] \right\} - \frac{1}{r} \left(\mu \left\{ 2 \left(\frac{1}{r} \frac{\partial u}{\partial \theta} + \frac{v}{r} \right) \right. \right. \right. \\ \left. \left. \left. - \frac{2}{3} \left[\frac{1}{r} \frac{\partial}{\partial r} (rv) + \frac{1}{r} \frac{\partial u}{\partial \theta} \right] \right\} \right) \right] . \quad (9)$$

Enthalpy:

$$\begin{aligned}
 \rho \left(v \frac{\partial h}{\partial r} + \frac{u}{r} \frac{\partial h}{\partial \theta} \right) = & v \frac{\partial P}{\partial r} + \frac{u}{r} \frac{\partial P}{\partial \theta} + \frac{\mu}{Re_{0\infty}} \left[2 \left\{ \left(\frac{\partial v}{\partial r} \right)^2 + \left(\frac{1}{r} \frac{\partial u}{\partial \theta} + \frac{v}{r} \right)^2 \right\} \right. \\
 & \left. + \left(\frac{1}{r} \frac{\partial v}{\partial \theta} + \frac{\partial u}{\partial r} - \frac{u}{r} \right)^2 \right] - \frac{2}{3} \frac{\mu}{Re_{0\infty}} \left[\frac{\partial v}{\partial r} + \frac{1}{r} \frac{\partial u}{\partial \theta} + \frac{v}{r} \right]^2 \\
 & + \frac{1}{r Pr Re_{0\infty}} \left\{ \frac{\partial}{\partial r} \left(r \mu \frac{\partial h}{\partial r} \right) + \frac{\partial}{\partial \theta} \left(\frac{\mu}{r} \frac{\partial h}{\partial \theta} \right) \right. \\
 & \left. + (Le - 1) \frac{\partial}{\partial r} \left[\sum_{i=1}^{NS} r \mu h_i \frac{\partial Y_i}{\partial r} \right] + (Le - 1) \frac{\partial}{\partial \theta} \left[\sum_{i=1}^{NS} \frac{\mu h_i}{r} \frac{\partial Y_i}{\partial \theta} \right] \right\} .
 \end{aligned} \tag{10}$$

State:

$$P = \rho T \frac{\bar{R} \bar{T}_{0\infty}}{\bar{V}_\infty^2} \sum_{i=1}^{NS} \frac{Y_i}{W_i} . \tag{11}$$

Spherical Coordinates:

Global Continuity:

$$\frac{1}{r} \frac{\partial}{\partial r} (\rho r^2 v) + \frac{1}{\sin \theta} \frac{\partial}{\partial \theta} (\rho u \sin \theta) = 0 . \tag{12}$$

Species Continuity:

$$\rho \left(v \frac{\partial Y_i}{\partial r} + \frac{u}{r} \frac{\partial Y_i}{\partial \theta} \right) = \dot{w}_i + \frac{1}{Re_{0\infty}} \frac{1}{r} \left[\frac{1}{r} \frac{\partial}{\partial r} \left(\frac{r^2 \mu}{Sc} \frac{\partial Y_i}{\partial r} \right) + \frac{1}{\sin \theta} \frac{\partial}{\partial \theta} \left(\frac{\mu \sin \theta}{Sc r} \frac{\partial Y_i}{\partial \theta} \right) \right] \quad i = 1, \dots, NS \quad (13)$$

Transverse Momentum (θ Direction):

$$\begin{aligned} \rho \left(v \frac{\partial u}{\partial r} + \frac{u}{r} \frac{\partial u}{\partial \theta} + \frac{uv}{r} \right) &= - \frac{1}{r} \frac{\partial P}{\partial \theta} + \frac{1}{Re_{0\infty}} \left\{ \frac{\partial}{\partial r} \left[\mu r \frac{\partial}{\partial r} \left(\frac{u}{r} \right) + \frac{\mu}{r} \frac{\partial v}{\partial \theta} \right] \right. \\ &+ \frac{1}{r^2} \frac{\partial}{\partial \theta} \left[\frac{4}{3} \mu \left(\frac{\partial u}{\partial \theta} + v \right) - \frac{2}{3} \mu \left(r \frac{\partial v}{\partial r} + v + u \cot \theta \right) \right] \\ &\left. + \frac{3\mu}{r} \left[r \frac{\partial}{\partial r} \left(\frac{u}{r} \right) + \frac{1}{r} \frac{\partial v}{\partial \theta} \right] + \frac{2\mu \cot \theta}{r^2} \left[\frac{\partial u}{\partial \theta} - u \cot \theta \right] \right\} . \end{aligned} \quad (14)$$

Radial Momentum (r Direction):

$$\begin{aligned} \rho \left(v \frac{\partial v}{\partial r} + \frac{u}{r} \frac{\partial v}{\partial \theta} - \frac{u^2}{r} \right) &= - \frac{\partial P}{\partial r} + \frac{1}{Re_{0\infty}} \left\{ \frac{\partial}{\partial r} \left[\frac{4}{3} \mu \frac{\partial v}{\partial r} - \frac{2}{3} \frac{\mu}{r} \left(\frac{\partial u}{\partial \theta} + 2v + u \cot \theta \right) \right] \right. \\ &+ \frac{1}{r} \frac{\partial}{\partial \theta} \left[\mu r \frac{\partial}{\partial r} \left(\frac{u}{r} \right) + \frac{\mu}{r} \frac{\partial v}{\partial \theta} \right] + \frac{4\mu}{r} \left(\frac{\partial v}{\partial r} - \frac{v}{r} \right) \\ &\left. - \frac{\mu}{r^2} \left[2 \frac{\partial u}{\partial \theta} + 2u \cot \theta - r^2 \cot \theta \frac{\partial}{\partial r} \left(\frac{u}{r} \right) - \frac{\partial v}{\partial \theta} \cot \theta \right] \right\} . \end{aligned} \quad (15)$$

Enthalpy:

$$\begin{aligned}
 \rho \left(v \frac{\partial h}{\partial r} + \frac{u}{r} \frac{\partial h}{\partial \theta} \right) = & v \frac{\partial P}{\partial r} + \frac{u}{r} \frac{\partial P}{\partial \theta} + \frac{\mu}{Re_{0\infty}} \left[2 \left\{ \left(\frac{\partial v}{\partial r} \right)^2 + \left(\frac{1}{r} \frac{\partial u}{\partial \theta} + \frac{v}{r} \right)^2 \right. \right. \\
 & \left. \left. + \left(\frac{v}{r} + \frac{u \cot \theta}{r} \right)^2 \right\} + \left\{ \frac{1}{r} \frac{\partial v}{\partial \theta} + r \frac{\partial}{\partial r} \left(\frac{u}{r} \right)^2 \right\} - \frac{2}{3} \frac{\mu}{Re_{0\infty}} \left[\frac{\partial v}{\partial r} + \frac{1}{r} \frac{\partial u}{\partial \theta} \right. \right. \\
 & \left. \left. + \frac{2v}{r} + \frac{u \cot \theta}{r} \right]^2 + \frac{1}{Re_{0\infty} Pr} \left\{ \frac{1}{r^2} \frac{\partial}{\partial r} \left(\mu r^2 \frac{\partial h}{\partial r} \right) + \frac{1}{r \sin \theta} \frac{\partial}{\partial \theta} \left(\frac{\mu \sin \theta}{r} \frac{\partial h}{\partial \theta} \right) \right. \right. \\
 & \left. \left. + (Le - 1) \frac{1}{r^2} \frac{\partial}{\partial r} \sum_{i=1}^{NS} \left(\mu r^2 h_i \frac{\partial Y_i}{\partial r} \right) \right. \right. \\
 & \left. \left. + (Le - 1) \frac{1}{r \sin \theta} \frac{\partial}{\partial \theta} \sum_{i=1}^{NS} \left(\frac{\mu h_i \sin \theta}{r} \frac{\partial Y_i}{\partial \theta} \right) \right\} \quad . \right. \quad (16)
 \end{aligned}$$

State:

$$P = \rho T \frac{\bar{R} \bar{T}_{0\infty}}{\bar{V}_\infty^2} \sum_{i=1}^{NS} \frac{Y_i}{W_i} \quad . \quad (17)$$

In the previous equations, Le_{ij} is assumed to be the same for all species pairs i, j and is, therefore, replaced with Le . Similarly, Sc_{ij} is replaced with Sc .

The local similarity approximation for the zone near the axis of symmetry is given by equations (18) through (24). The validity of this approximation was demonstrated by Kao [7].

$$u(r, \theta) = u_1(r) \sin \theta \quad (18)$$

$$v(r, \theta) = v_1(r) \cos \theta \quad (19)$$

$$h(r, \theta) = h_1(r) \quad (20)$$

$$\rho(r, \theta) = \rho_1(r) \quad (21)$$

$$P(r, \theta) = P_1(r) + P_2(r) \sin^2 \theta \quad (22)$$

$$\mu(r, \theta) = \mu_1(r) \quad (23)$$

$$Y_i(r, \theta) = Y_{1i}(r) \quad . \quad (24)$$

Equations (18) through (24) are used to reduce the governing equations to a set of nonlinear, coupled, ordinary differential equations which can be solved rapidly.

A transformation is now made in the normal coordinate by defining

$$\eta \equiv \frac{r - 1}{r_\infty - 1} = \frac{n}{n_\infty} \quad , \quad (25)$$

where r_∞ is the nondimensional distance from the origin to the freestream and n_∞ is the nondimensional distance from the body to the freestream (Fig. 1b). The values of r_∞ and n_∞ are unknown a priori. They are determined as part of the solution. This transformation, equation (25), keeps the body at $\eta = 0$ and the freestream at $\eta = 1$.

By substituting equations (18) through (25) into the governing equations and equating the coefficients of like functions of θ , one obtains the following system of ordinary differential equations:

Cylindrical Coordinates:

$$\frac{\rho'_1}{\rho_1} = - \frac{v'_1}{v_1} - \frac{n_\infty}{1 + \eta n_\infty} \left(1 + \frac{u_1}{v_i} \right) . \quad (26)$$

$$\begin{aligned} \frac{u''_1}{n_\infty^2} &= \frac{Re_{0\infty} \rho_1}{\mu_1} \left(\frac{v_1 u'_1}{n_\infty} + \frac{u_1^2 + u_1 v_1}{1 + \eta n_\infty} \right) + \frac{2 P_2 Re_{0\infty}}{\mu_1 (1 + \eta n_\infty)} \\ &+ \frac{u_1 + v_1}{(1 + \eta n_\infty)} \left[\frac{\mu'_1}{\mu_1 n_\infty} + \frac{7}{3(1 + \eta n_\infty)} \right] - \frac{u'_1}{n_\infty} \left(\frac{2}{1 + \eta n_\infty} + \frac{\mu'_1}{\mu_1 n_\infty} \right) \\ &+ \frac{v'_1}{3n_\infty (1 + \eta n_\infty)} . \end{aligned} \quad (27)$$

$$\begin{aligned} \frac{v''_1}{n_\infty^2} &= \frac{3}{4} \frac{Re_{0\infty}}{n_\infty \mu_1} (P'_1 + \rho_1 v_1 v'_1) - \frac{v'_1}{n_\infty} \left(\frac{\mu'_1}{\mu_1 n_\infty} + \frac{1}{1 + \eta n_\infty} \right) - \frac{u'_1}{4n_\infty (1 + \eta n_\infty)} \\ &+ \frac{(u_1 + v_1)}{1 + \eta n_\infty} \left[\frac{\mu'_1}{2\mu_1 n_\infty} + \frac{7}{4(1 + \eta n_\infty)} \right] . \end{aligned} \quad (28)$$

$$\frac{P'_2}{n_\infty} = - \frac{P'_1}{n_\infty} + \frac{\rho_1 u_1}{1 + \eta n_\infty} (u_1 + v_1) . \quad (29)$$

$$\begin{aligned}
\frac{\rho_1 v_1 h'_1}{n_\infty} \cdot (1 + \eta n_\infty)^2 &= (1 + \eta n_\infty)^2 \left[\frac{v_1 P'_1}{n_\infty} + \frac{2\mu_1}{Re_{0\infty}} \frac{v'_1^2}{n_\infty^2} \right] + \frac{2\mu_1}{Re_{0\infty}} (u_1 + v_1)^2 \\
&\quad - \frac{2}{3} \frac{\mu_1}{Re_{0\infty}} \left[\frac{(1 + \eta n_\infty)}{n_\infty} v'_1 + (u_1 + v_1) \right]^2 \\
&\quad + \frac{(1 + \eta n_\infty)}{Re_{0\infty} Pr n_\infty} \left\{ \mu_1 h'_1 + \frac{(1 + \eta n_\infty)}{n_\infty} (h'_1 \mu'_1 + \mu_1 h'_1) \right. \\
&\quad \left. + (Le - 1) \left[\left(\mu_1 + \frac{\mu'_1 (1 + \eta n_\infty)}{n_\infty} \right) \sum_{i=1}^{NS} h_i Y'_{1i} \right. \right. \\
&\quad \left. \left. + \frac{\mu_1 (1 + \eta n_\infty)}{n_\infty} \sum_{i=1}^{NS} (h'_i Y'_{1i} + h_i Y''_{1i}) \right] \right\} . \tag{30}
\end{aligned}$$

$$\begin{aligned}
\frac{\rho_1 v_1 Y'_{1i}}{n_\infty} &= \dot{w}_i + \frac{1}{(1 + \eta n_\infty) Re_{0\infty} Sc} \left[\frac{\mu_1 Y'_{1i}}{n_\infty} + \frac{\mu'_1 Y'_{1i} (1 + \eta n_\infty)}{n_\infty^2} \right. \\
&\quad \left. + \frac{\mu_1 (1 + \eta n_\infty) Y''_{1i}}{n_\infty^2} \right] \quad i = 1, \dots, NS . \tag{31}
\end{aligned}$$

$$P_1 = \rho_1 T_1 \left(\frac{\bar{R} \bar{T}_{0\infty}}{\bar{V}_\infty^2} \right) \sum_{i=1}^{NS} \left(\frac{Y_{1i}}{W_i} \right) . \tag{32}$$

Here, a prime denotes differentiation with respect to η .

Spherical Coordinates:

$$\frac{\rho'_1}{\rho_1} = -\frac{v'_1}{v_1} - \frac{2n_\infty}{1 + \eta n_\infty} \left(1 + \frac{u_1}{v_1} \right) \quad (33)$$

$$\begin{aligned} \frac{u''_1}{n_\infty^2} &= \frac{Re_{0\infty} \rho_1}{\mu_1} \left(\frac{v'_1 u'_1}{n_\infty} + \frac{u_1^2 + u_1 v_1}{1 + \eta n_\infty} \right) + \frac{2P_2 Re_{0\infty}}{\mu_1 (1 + \eta n_\infty)} \\ &+ \frac{u_1 + v_1}{(1 + \eta n_\infty)} \left[\frac{\mu'_1}{\mu_1 n_\infty} + \frac{8}{3(1 + \eta n_\infty)} \right] - \frac{u'_1}{n_\infty} \left(\frac{2}{1 + \eta n_\infty} + \frac{\mu'_1}{\mu_1 n_\infty} \right) \\ &+ \frac{v'_1}{3n_\infty (1 + \eta n_\infty)} . \end{aligned} \quad (34)$$

$$\begin{aligned} \frac{v''_1}{n_\infty^2} &= \frac{3}{4} \frac{Re_{0\infty}}{n_\infty \mu_1} (P'_1 + \rho_1 v_1 v'_1) - \frac{v'_1}{n_\infty} \left(\frac{\mu'_1}{\mu_1 n_\infty} + \frac{2}{1 + \eta n_\infty} \right) - \frac{u'_1}{2n_\infty (1 + \eta n_\infty)} \\ &+ \frac{(u_1 + v_1)}{(1 + \eta n_\infty)} \left[\frac{\mu'_1}{\mu_1 n_\infty} + \frac{7}{2(1 + \eta n_\infty)} \right] . \end{aligned} \quad (35)$$

$$\frac{P'_2}{n_\infty} = -\frac{P'_1}{n_\infty} + \frac{\rho_1 u_1}{1 + \eta n_\infty} (u_1 + v_1) . \quad (36)$$

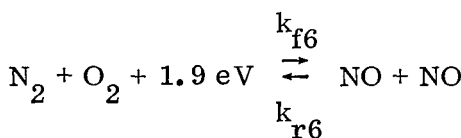
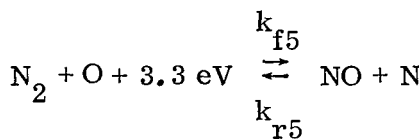
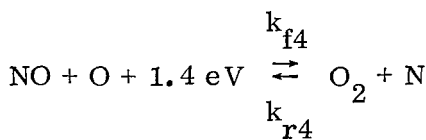
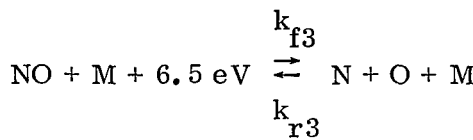
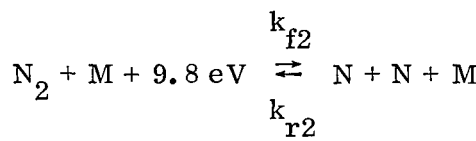
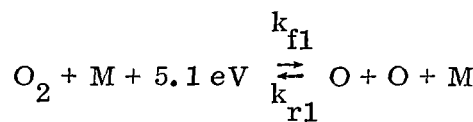
$$\begin{aligned}
& \frac{\rho_1 v_1 h'_1}{n_\infty} (1 + \eta n_\infty)^2 = (1 + \eta n_\infty)^2 \left[\frac{v_1 P'_1}{n_\infty} + \frac{2\mu_1}{Re_{0\infty}} \frac{v'_1^2}{n_\infty^2} \right] + \frac{4\mu_1}{Re_{0\infty}} (u_1 + v_1)^2 \\
& - \frac{2}{3} \frac{\mu_1}{Re_{0\infty}} \left[\frac{(1 + \eta n_\infty)}{n_\infty} v'_1 + 2 (u_1 + v_1) \right]^2 \\
& + \frac{(1 + \eta n_\infty)}{Re_{0\infty} Pr n_\infty} \left\{ 2\mu_1 h'_1 + \frac{(1 + \eta n_\infty)}{n_\infty} (\mu'_1 h'_1 + \mu''_1 h''_1) \right. \\
& \left. + (Le - 1) \left[\left(2\mu_1 + \frac{\mu'_1 (1 + \eta n_\infty)}{n_\infty} \right) \sum_{i=1}^{NS} h_i Y'_{1i} \right. \right. \\
& \left. \left. + \frac{\mu_1 (1 + \eta n_\infty)}{n_\infty^2} \sum_{i=1}^{NS} (h'_i Y'_{1i} + h''_i Y''_{1i}) \right] \right\} . \quad (37)
\end{aligned}$$

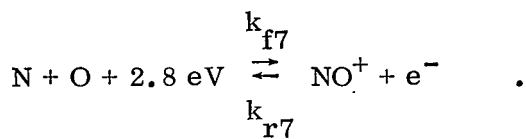
$$\begin{aligned}
& \frac{\rho_1 v_1 Y'_{1i}}{n_\infty} = \dot{w}_i + \frac{1}{(1 + \eta n_\infty) Re_{0\infty} Sc} \left[\frac{2\mu_1 Y'_{1i}}{n_\infty} + \frac{\mu_1 Y'_{1i} (1 + \eta n_\infty)}{n_\infty^2} \right. \\
& \left. + \frac{\mu_1 (1 + \eta n_\infty) Y''_{1i}}{n_\infty^2} \right] \quad i = 1, \dots, NS . \quad (38)
\end{aligned}$$

$$P_1 = \rho_1 T_1 \frac{\bar{R} \bar{T}_{0\infty}}{\bar{V}_\infty^2} \sum_{i=1}^{NS} \left(\frac{Y_{1i}}{W_i} \right) . \quad (39)$$

Equations (26) through (32) and (33) through (39) constitute two sets of non-linear ordinary differential equations for cylindrical and spherical stagnation regions, respectively. Each of these sets contain $6 + NS$ equations and $8 + 2NS$ unknowns ($\mu_1, v_1, P_1, P_2, \rho_1, T_1, h_1, \mu_1, \dot{w}_i$, and Y_{1i}). The required additional information and equations are given below for the mass production rates, \dot{w}_i , enthalphy, h_1 , and viscosity, μ_1 , of a dissociated and ionized air mixture. These equations, like the equation of state above, are independent of the coordinate system.

c. Air Chemistry. The freestream air is assumed to consist of N_2 and O_2 molecules only. The air model used for the shock layer consists of seven species and the seven chemical reactions as follows [5]:





These reactions are written so that the forward reactions are endothermic; the net amounts of energy required to produce the reactions are given on the left side of the equations. The first three reactions are the neutral-particle dissociation-recombination reactions in which the energy of dissociation in the forward reaction is taken primarily from kinetic energy by means of a collision with a "catalytic" molecule M; the chemical energy released in the recombination (reverse reaction) is converted primarily to kinetic energy in a three-body collision involving a catalytic molecule M. The M molecule can be any of the six molecular or atomic species present in the air mixture. The quantities k_{fj} and k_{rj} are the temperature-dependent forward and reverse reaction rates, respectively, for the jth reaction. The experimentally determined reaction rates recommended by Wray [5] are used, together with the concentrations of all constituents, to obtain the mass production rates, \dot{w}_i , of each constituent [9]. The reader is referred to Reference 10 for details.

d. Enthalpy. The specific enthalpy of a mixture of gases is given in dimensional form as follows:

$$\bar{h} = \bar{P} \bar{v} + \sum_{i=1}^{NS} Y_i \bar{e}_i , \quad (40)$$

where

$$\bar{e}_i = \bar{e}_i^0 + \bar{e}_{T_i} + \bar{e}_{R_i} + \bar{e}_{v_i} \quad (41)$$

and

\bar{e}_i^0 = specific energy of formation of species i at the reference temperature (zero absolute)

\bar{e}_{T_i} = specific energy of random translation

\bar{e}_{R_i} = specific energy of rotation (for diatomic molecule)

\bar{e}_{v_i} = specific energy of vibration (for diatomic molecules).

We assume that the energy of electronic excited states is negligible, that the rotational state is fully excited, but that the vibrational state is partially excited. Values for these quantities are given by Vincenti and Kruger [9]:

$$\bar{e}_{T_i} = \frac{3}{2} \bar{R}_i \bar{T} \quad (42)$$

$$\bar{e}_{R_i} = \bar{R}_i \bar{T} \quad (43)$$

$$\bar{e}_{v_i} = \frac{\bar{R}_i \Theta_{v_i}}{e^{\Theta_{v_i}/\bar{T}} - 1}, \quad (44)$$

where

Θ_{v_i} = characteristic temperature for vibration of diatomic species i

$$\Theta_{v_{O_2}} = 2270 \text{ K}$$

$$\Theta_{v_{N_2}} = 3390 \text{ K}$$

$$\Theta_{v_{NO}} = 2740 \text{ K}$$

$$\Theta_{v_{NO^+}} = 2740 \text{ K} .$$

Equation (40) now becomes

$$\bar{h} = \frac{5}{2} \bar{\rho} \bar{T} \sum_{i=1}^{NS} \frac{Y_i}{W_i} + \bar{\rho} \sum_i \frac{Y_i}{W_i} \left(\bar{T} + \frac{\bar{\theta}_v^i}{\bar{\theta}_v^i / \bar{T} - 1} \right) + \sum_{i=1}^{NS} \bar{e}_i^0 . \quad (45)$$

(for
diatomic
molecules)

Nondimensionalizing equation (45), we obtain

$$h = \frac{5}{2} \frac{\bar{\rho} \bar{T} \bar{T}_{0\infty}}{\bar{V}_\infty^2} \sum_{i=1}^{NS} \frac{Y_i}{W_i} + \frac{\bar{\rho} \bar{T}_{0\infty}}{\bar{V}_\infty^2} \sum_i \frac{Y_i}{W_i} \left(T + \frac{\bar{\theta}_v^i}{\bar{\theta}_v^i / T - 1} \right) + \frac{1}{\bar{V}_\infty^2} \sum_{i=1}^{NS} Y_i \bar{e}_i^0 . \quad (46)$$

(for
diatomic
molecules)

e. Viscosity. The Sutherland formula for the viscosity of air gives acceptable results at moderate temperatures, but it fails at the extremely high temperatures encountered in hypersonic flight. The viscosity begins deviating from the Sutherland value due to the onset of dissociation at approximately 3000 K. The viscosity near the wall is also affected by the extremely low pressures encountered at high altitude [11]. This effect is due to velocity slip at the wall (see next section). The Sutherland formula, equation (47), is recommended for temperatures less than 3000 K.

$$\bar{\mu}_{su} = \frac{1.458 \times 10^{-5} \bar{T}^{3/2}}{110.4 + \bar{T}} \quad [\text{gm/cm sec}] . \quad (47)$$

This formula is also used in the computer program (Appendix A) to calculate $\bar{\mu}_{0\infty}$ using $\bar{T}_{0\infty}$ computed from the adiabatic relation for temperature,

although it is realized that neither $\bar{\mu}_{su}$ nor $\bar{T}_{0\infty}$ has valid physical meaning at extremely high stagnation enthalpy. This use is justified because $\bar{\mu}_{0\infty}$ is used only for nondimensionalizing $\bar{\mu}$ and for computing $Re_{0\infty}$.

The viscosity of dissociated and ionized air is approximated in Reference 6 using a simple summation formula for a mixture of hard spherical molecules:

$$\frac{\bar{\mu}}{\bar{\mu}_{su}} = \sum_{i=1}^{NS} Y_i \sqrt{\frac{W_R}{W_i}} \frac{\bar{\lambda}_i}{\bar{\lambda}_R}, \quad (48)$$

where

$\bar{\mu}_{su}$ = viscosity at same temperature from Sutherland formula

W_R = equivalent molecular weight of undissociated air

$\bar{\lambda}_i$ = mean free path of species i

$\bar{\lambda}_R$ = mean free path of undissociated air molecules.

The ratio of mean free paths in equation (48) is given by

$$\frac{\bar{\lambda}_R}{\bar{\lambda}_i} = \sum_{j=1}^{NS} \left[Y_j \frac{W_R}{W_j} \frac{\bar{s}_{ij}}{\bar{s}_R} \left(\frac{1 + \frac{W_i}{W_j}}{2} \right)^{1/2} \right], \quad (49)$$

where

\bar{s}_{ij} = collision cross section for particle i with particle j

\bar{s}_R = collision cross section for undissociated air molecules.

The collision cross sections are tabulated as a function of temperature in Reference 6.

The viscosity given by equation (48) is a function of the composition of the gas mixture and also of the temperature via the dependence of \bar{S}_R and \bar{S}_{ij} on temperature. Since in the present analysis the air is not in chemical equilibrium, the viscosity obtained from the computer program at a given temperature differs greatly from that shown in Reference 6 for equilibrium conditions. As the mass fractions of the components of air approach their undissociated values, the numerical value of the viscosity ratio in equation (48) approaches 1.0, i.e., $\bar{\mu}$ approaches the Sutherland value as expected.

3. Boundary Conditions. To solve the governing equations given in the previous section, freestream and wall boundary conditions are required.

Freestream ($\eta = 1$):

The air at the freestream boundary is in its undisturbed state.

$$u_1 = 1 ,$$

$$v_1 = -1 ,$$

$$\rho_1 = 1 ,$$

$$T_1 = T_\infty ,$$

$$P_1 = P_\infty ,$$

$$P_2 = 0 ,$$

$$Y_{1N_2} = 0.767 ,$$

$$Y_{1O_2} = 0.233 ,$$

$$Y_{1NO} = Y_{1N} = Y_{1O} = Y_{1NO^+} = Y_{1e^-} = 0 .$$

Wall ($\eta = 0$):

In high-altitude, low-Reynolds number flight, the continuum model of the gas breaks down in regions of large gradients of the physical properties near the wall. Hence, the Navier-Stokes description is invalid for the gas layer near the wall (Knudsen layer) with thickness on the order of the mean free path [11]. Also, the familiar continuum zero-velocity and zero-temperature-jump wall boundary conditions are not applicable. Although the Navier-Stokes equations are invalid near the wall, they can still be used, down to quite low Reynolds numbers, to describe the outer flow field if the proper boundary conditions are used at the outer edge of the Knudsen layer. These boundary conditions, known as slip conditions, are the mean velocity, temperature, and species mass fractions. To compute these slip conditions, a kinetic theory approach must be used for the Knudsen layer. The boundary conditions for the Knudsen layer are the mean slip conditions at the outer edge and the kinetic gas-surface conditions at the wall.

The Boltzmann equation is the governing equation for the kinetic theory description of a flow field. However, due to the difficulty in solving the Boltzmann equation for the Knudsen layer, we resorted to an approximate kinetic theory slip model. Reference 4 gives the details of the derivation of the slip conditions for a multicomponent reacting gas. By matching the species, momenta, and energy fluxes at the outer edge of the Knudsen layer to the difference between the incident and reflected fluxes at the wall, the jump in the desired properties across the Knudsen layer is obtained. The fluxes are calculated by taking moments of the velocity distribution function which is approximated by using a Chapman-Enskog expansion for a multicomponent mixture. The species flux is greatly affected by the catalytic nature of the wall. The wall is assumed to be catalytic with respect to recombination of dissociated molecules. Equations are obtained for a partially catalytic wall. The extremes of noncatalytic and fully catalytic walls are easily obtained from the equations for a partially catalytic wall. The resulting nondimensional equations for slip velocity, temperature, and species are as follows:

$$u_s = \frac{\mu_s}{P_w R e_\infty M_\infty} \left[\sqrt{\pi} \left(\frac{2 - \sigma}{2\sigma} \right) \sqrt{\frac{2T_s}{\gamma T_\infty}} \left(\frac{\partial u}{\partial r} + \frac{1}{r} \frac{\partial v}{\partial \theta} \right)_s \right. \\ + \frac{1}{5(\gamma - 1) M_\infty \Pr T_\infty} \left(\frac{1}{r} \frac{\partial T}{\partial \theta} \right)_s \sum_i \sqrt{\frac{w_i}{w}} \\ \left. - \frac{P_w M_\infty L_e}{\rho_w \Pr} \sum_j \left(\frac{1}{r} \frac{\partial Y_j}{\partial \theta} \right)_s \sum_i \sqrt{\frac{w_i}{w}} \right] . \quad (50)$$

$$\begin{aligned}
T_s &= \left(T_w \sum_i \left\{ \frac{\rho Y_i}{\left[\frac{W_i}{W_j} \sum_j \frac{Y_j}{W_j} \right]^{3/2}} \left[1 + \frac{\mu}{3P_i Re_{0\infty}} \left(\frac{1}{r} \frac{\partial u}{\partial \theta} - 2 \frac{\partial v}{\partial r} \right) \right. \right. \right. \\
&\quad \left. \left. \left. + \frac{\sqrt{\pi}}{Y_i} \left(\frac{2-\sigma}{\sigma} \right) \frac{M_\infty}{Re_{0\infty}} \frac{Le}{Pr} \left[W_i \sum_j \frac{Y_j}{W_j} \right] \frac{\mu}{\rho} \sqrt{\frac{\gamma W_i T_\infty}{2WT}} \sum_j \frac{\partial Y_j}{\partial r} \right] \right\}_s \right. \\
&\quad \left. + \left\{ \frac{2-\sigma}{\sigma} \sqrt{\pi} \left[\frac{M_\infty}{Pr Re_{0\infty}} \frac{\gamma}{\gamma-1} \frac{\mu}{2} \sqrt{\frac{\gamma T_\infty}{2T}} \frac{\partial T}{\partial r} \right. \right. \right. \\
&\quad \left. \left. \left. - \frac{5P}{2\rho} \frac{\mu}{Re_{0\infty}} M_\infty^3 \gamma T_\infty \sqrt{\frac{\gamma T_\infty}{2T}} \frac{Le}{Pr} \sum_j \frac{\partial Y_j}{\partial r} \right] \right\}_s \right) \\
&\quad \left\{ \sum_i \frac{\rho Y_i}{\left[\frac{W_i}{W_j} \sum_j \left(\frac{Y_j}{W_j} \right) \right]^{3/2}} \left[1 + \frac{\mu}{2P_i Re_{0\infty}} \left(\frac{1}{r} \frac{\partial u}{\partial \theta} - 2 \frac{\partial v}{\partial r} \right) \right] \right\}_s . \quad (51)
\end{aligned}$$

$$Y_i^s = \frac{\frac{2-\gamma}{2\gamma_i} \frac{M_\infty}{Re_{0\infty}} \frac{Le}{Pr} \frac{\mu_s}{\rho_s} \sqrt{\frac{2\pi\gamma W_i T_\infty}{WT_s}} \sum_j \left(\frac{\partial Y_j}{\partial r} \right)_s}{1 + \frac{M_\infty^2}{Re_{0\infty}} \left(\frac{\mu}{3\rho T Y_i} \right)_s \frac{\gamma W_i T_\infty}{W} \left(\frac{1}{r} \frac{\partial u}{\partial \theta} - 2 \frac{\partial v}{\partial r} \right)_s} . \quad (52)$$

The superscript, s, on Y_i^s denotes the value of Y_i at the outer edge of the Knudsen layer.

The similarity equations (18) through (24) and the coordinate transformation equation (25) are used in equations (50) through (52) to yield the following slip equations:

$$u_{1s} = \frac{\mu_{1s}}{P_{1w} Re_{0\infty} M_\infty} \left[\sqrt{\pi} \left(\frac{2-\sigma}{2\sigma} \right) \sqrt{\frac{2T_1}{\gamma T_\infty}} \left(\frac{u'_1}{\eta_\infty} \right) \right]_s , \quad (53)$$

$$\begin{aligned} T_{1s} = & \left(T_w \sum_i \left\{ \frac{\rho_1 Y_{1i}}{\left[w_i \sum_j \frac{Y_{1j}}{w_j} \right]^{3/2}} \left[1 + \frac{\mu_1}{3P_{1i} Re_{0\infty}} \left(\frac{u_1}{1 + \eta n_\infty} - \frac{2v'_1}{n_\infty} \right) \right. \right. \right. \right. \\ & + \frac{\sqrt{\pi}}{Y_{1i}} \left(\frac{2-\sigma}{\sigma} \right) \frac{M_\infty}{Re_{0\infty}} \frac{Le}{Pr} \left[w_i \sum_j \frac{Y_{1j}}{w_j} \right] \frac{\mu_1}{\rho_1} \sqrt{\frac{\gamma w_i T_\infty}{2w T_1}} \sum_j \frac{Y'_{1j}}{n_\infty} \left. \left. \left. \left. \right\} s \right. \right. \\ & + \frac{2-\sigma}{\sigma} \sqrt{\pi} \left[\frac{M_\infty}{Pr Re_{0\infty}} \frac{\gamma}{\gamma-1} \frac{\mu_1}{2} \sqrt{\frac{\gamma T_\infty}{2T_1}} \times \frac{T'_1}{n_\infty} \right. \\ & \left. \left. \left. \left. - \frac{5P_1}{2\rho_1} \frac{\mu_1}{Re_{0\infty}} M_\infty^3 \gamma T_\infty \sqrt{\frac{\gamma T_\infty}{2T_1}} \frac{Le}{Pr} \sum_j \frac{Y'_{1j}}{n_\infty} \right] \right\} s \right. \\ & \div \left\{ \sum_i \frac{\rho_1 Y_{1i}}{\left[w_i \sum_j \frac{Y_{1j}}{w_j} \right]^{3/2}} \left[1 + \frac{\mu_1}{2P_{1i} Re_{0\infty}} \left(\frac{u_1}{1 + \eta n_\infty} - \frac{2v'_1}{n_\infty} \right) \right] \right\}_s , \end{aligned} \quad (54)$$

and

$$Y_{1i}^S = \frac{\frac{2 - \gamma_i}{2\gamma_i} \frac{M_\infty}{Re_{0\infty}} \frac{Le}{Pr} \frac{\mu_{1s}}{\rho_{1s}} \sqrt{\frac{2\pi\gamma W_i T_\infty}{WT_{1s}}} \sum_j \left(\frac{Y'_{1j}}{n_\infty} \right)_s}{1 + \frac{M_\infty^2}{Re_{0\infty}} \left(\frac{\mu_1}{3\rho_1 T_1 Y_{1i}} \right)_s \frac{\gamma W_i T_\infty}{W} \left(\frac{u_1}{1 + \eta n_\infty} - \frac{2v'_1}{n_\infty} \right)_s} . \quad (55)$$

The quantity γ_i in equation (55) is the recombination coefficient for the i species; it is a measure of the catalyticity of the wall. The value of γ_i varies from 0 (noncatalytic wall) to 1 (fully catalytic wall).

The computer program used in this report has the fully catalytic wall and noncatalytic wall options, but it does not, in the present form, have a partially catalytic wall option [for a given catalyticity γ_i , equation (55) would give Y_i^S]. In the fully catalytic wall option, the wall is assumed to be catalytic only with respect to recombination of neutral atomic species; it is assumed to be non-catalytic with respect to recombination of the charged particles NO^+ and e^- . The wall boundary conditions on Y_i^S for these special cases are given as follows:

Noncatalytic Wall:

For a noncatalytic surface ($\gamma_i = 0$), equation (55) reduces to

$$\sum_j (Y'_{1j})_s = 0 . \quad (56)$$

A sufficient condition for equation (56) to be satisfied is that

$$(Y'_{ij})_s = 0 \quad j = 1, \dots, NS . \quad (57)$$

Equation (57) is also the most physically plausible means by which equation (56) can be satisfied; therefore, equation (57) is taken as the noncatalytic boundary condition for species mass fractions.

Fully Catalytic Wall:

For the fully catalytic wall, the surface is assumed to be fully catalytic with respect to recombination of neutral atomic species, but it is assumed to be noncatalytic with respect to recombination of the charged particles NO^+ and e^- . Therefore, the effect of the wall is to drive the gas towards its freestream composition, except for the charged particles NO^+ and e^- . The following boundary conditions are then obtained:

$$\left(Y_{N_2} \right)_s = 0.767 \quad (58)$$

$$\left(Y_{O_2} \right)_s = 0.233 \quad (59)$$

$$(Y_{NO})_s = (Y_N)_s = (Y_O)_s = 0 \quad (60)$$

$$(Y'_{NO^+})_s = (Y'_{e^-})_s = 0 \quad . \quad (61)$$

Equations (58), (59), and (61) produce the impossible result that

$$\sum_{i=1}^{NS} Y_i > 1 ;$$

however, because $(Y_{NO^+})_s$ and $(Y_{e^-})_s$ are usually very small, this error is tolerable.

B. Method of Solution

Equations (26) through (32) and (33) through (39) together with equations (46) and (48) constitute two sets of nonlinear, coupled ordinary differential equations with boundary conditions previously given. The first order equations are solved by direct numerical quadrature, and the second order equations are

integrated by a finite difference method known as the successive accelerated replacement method which is an iterative scheme that starts from a guessed solution. The salient feature of this method, proposed by Lieberstein [13], is that the successive corrections applied to the flow profiles in each iteration are controlled by acceleration factors which are used to increase the rate of convergence of the computed quantities. Thus, this method can be successfully applied even if the initial, guessed profiles do not approximate the converged solutions very well. For the present application, this statement holds true in the midrange of the flow regime for which our analysis is applicable. However, divergence problems are encountered at the continuum end of the regime (Appendix A).

C. Computer Program

Details of the computer program are found in Reference 10, while Appendix A is a current users manual.

III. RESULTS

Results from the computer program, with the modifications introduced in this report, are first compared with the 'old' program. Then, the program output is compared with some available experimental data. Finally, some general results from the program and input data for running the program are presented.

A. Effect of Computer Program Modifications

The program described as the old program includes all the analysis in this report except the constitutive equations (46) and (48) for enthalpy and viscosity, respectively. The old enthalpy equation assumed a fully excited vibrational state for diatomic molecules. This assumption is always violated near the freestream and near the wall because temperatures in these regions are less than Θ_v . The old viscosity equation (Sutherland) is restricted to non-dissociated air or to temperatures less than approximately 3000 K at equilibrium conditions. This temperature is greatly exceeded in the central part of the shock layer at reentry speeds.

B. Comparison of Data Between Old Program and Modified Program

Computed values of shock layer thickness, n_{∞} ; wall heat transfer coefficient at the stagnation point, C_H ; and maximum ratio of viscosity to Sutherland viscosity on the stagnation streamline, μ/μ_{sv} , are given in Table 1 for a 30.5-cm radius sphere using (1) the "old" program, (2) the program with enthalpy modification only, and (3) the program with enthalpy and viscosity modifications. The conditions for which the runs were made are listed in Table 1. The thermodynamic properties associated with altitude are obtained throughout this report from Reference 14.

TABLE 1. DATA COMPARISON

$$\text{alt} = 86 \text{ km}$$

$$\bar{r}_b = 30.5 \text{ cm}$$

$$\bar{V}_{\infty} = 793\,000 \text{ cm/sec}$$

Program Identification	Shock Layer Thickness n_{∞}	Wall Heat Transfer Coefficient C_H	$\left(\frac{\mu}{\mu_{sv}}\right)_{\max}$
1. Old Program	0.1361	0.147	1.0
2. Modified Program (a) New enthalpy computation, equation (46) (b) Sutherland viscosity	0.1409	0.154	1.0
3. Modified Program (a) New enthalpy computation, equation (46) (b) New viscosity, equation (48)	0.1411	0.172	1.37

For these runs, conditions were chosen at which the modifications in viscosity and enthalpy computation would have their greatest effect. These conditions are the ones which yield a high degree of dissociation and ionization, equations (46) and (48), i.e. high speed and high freestream density. Therefore, an altitude was chosen (86 km) where freestream density is near the maximum for which the program will run (see General Results section). The table shows that the enthalpy and viscosity modifications had a significant effect on heat transfer rate and shock layer thickness. The heat transfer rate was increased 17 percent due to the combined effect of both modifications.

C. Comparison with Experimental Data

Very little experimental data exist for the low density, hypersonic, high stagnation enthalpy flow regime for which this computer program was designed. Therefore, we do not have the detailed comparison with experimental data which is desired to validate the present analysis. However, a limited amount of heat transfer and pressure data for spheres are available from an arc-jet facility at moderate stagnation enthalpy [15], and a larger body of stagnation heat transfer data exists for relatively low enthalpy, and hypersonic flow [16].

Figure 2 compares computed stagnation point heat transfer coefficients, C_H , for a sphere with experimental data from References 15 and 16. The data in this figure represent several levels of stagnation enthalpy and, hence, several levels of real gas effects, so good agreement among all the data is not expected. The purpose here is to compare the computed results directly with each set of experimental data and to observe the trend in C_H with changing stagnation enthalpy. A best fit curve through the relatively low enthalpy data of Vidal and Wittliff [16] is shown by the solid line, and the square symbols give the computed values for the same flow conditions. Although the computer program was designed for real gas, chemically reacting flows, it can be run at the low enthalpy conditions of the Vidal and Wittliff data, in which case the computed dissociation levels are very low. The data are plotted as a function of the similarity parameter, K^2 , used by Vidal and Wittliff, and defined as follows:

$$K^2 = \frac{Re_\infty}{\gamma_\infty M_\infty^2 C^*} , \quad (62)$$

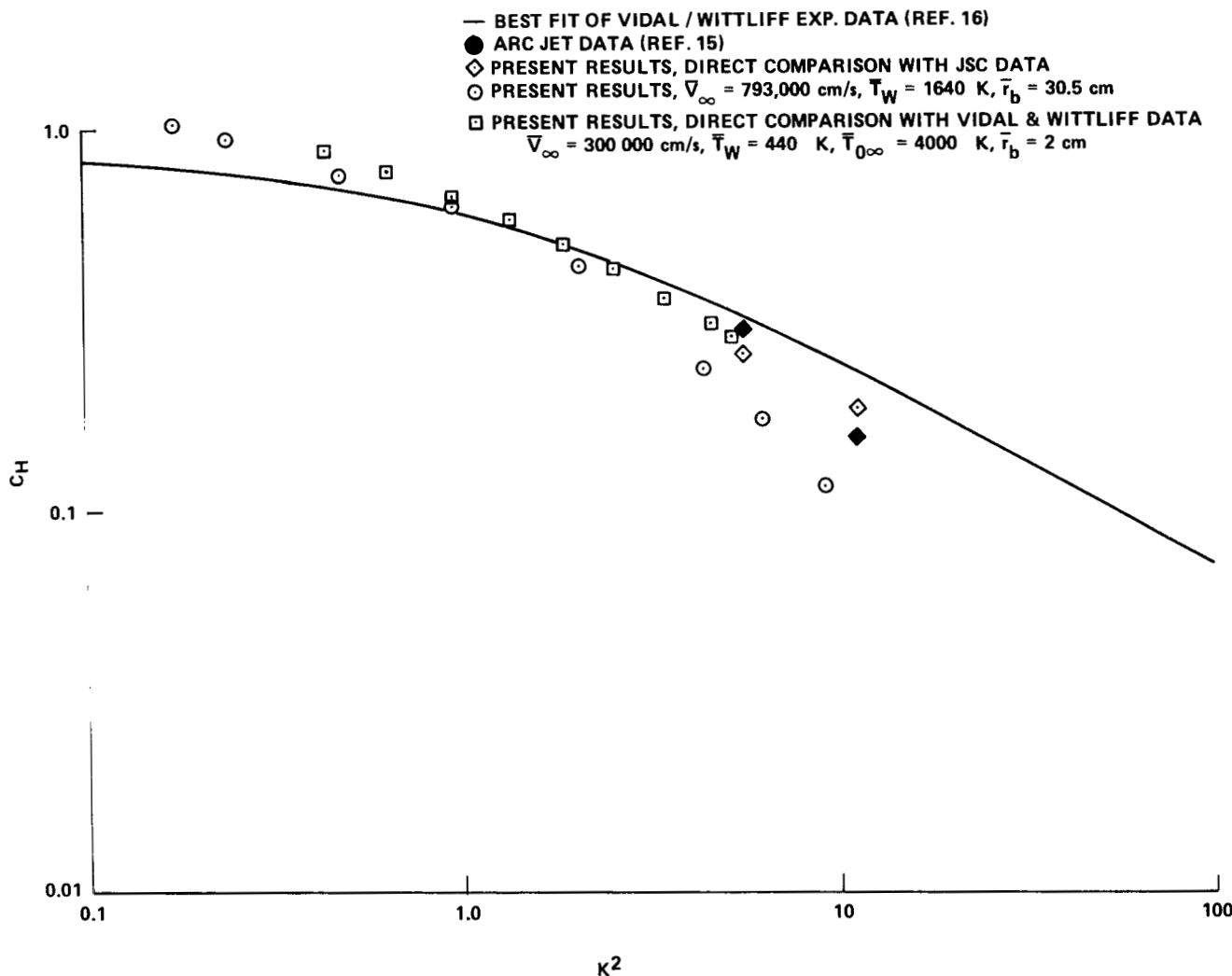


Figure 2. Comparison of present theory with experimental data for stagnation point heating on a sphere.

where

$$C^* = \frac{\bar{\mu}^*}{\bar{\mu}_\infty} \frac{\bar{T}_\infty}{\bar{T}^*} \quad (63)$$

and

$$\bar{T}^* = \frac{1}{2} (\bar{T}_{0\infty} + \bar{T}_w) \quad , \quad (64)$$

where

$\bar{\mu}^*$ = viscosity computed by Sutherland's formula using \bar{T}^*

$\bar{\mu}_\infty$ = viscosity computed by Sutherland's formula using \bar{T}_∞ .

The computed data do not agree with the mean of the Vidal and Wittliff data as well as we would like, but the computed data points fall within the scatter of the experimental points [16]. The computed values tend to overestimate C_H at rarefied conditions and underestimate it for denser flow. It is important to note that at the higher enthalpy cases, the wall catalysis can affect the value of C_H by a factor of 3 or more [4]. Hence, any comparison of experimental and calculated heating must be for similar wall catalysis.

The arc-jet data of Scott [15] are shown by the filled diamond symbols, and computed data for the same conditions are shown by the empty diamond symbols. The flow conditions for these tests are given in Table 2.

The freestream boundary conditions in the computer program were modified to match the dissociated arc-jet freestream. Also, the spheres in the arc-jet tests were coated with teflon to produce a noncatalytic wall and, therefore, the noncatalytic wall option in the computer program was used for comparison with these data points. The noncatalytic wall option was also used for the remainder of the computer data in this figure and the entire report unless otherwise noted.

The comparison of the computed data with Scott's experimental data is considered acceptable.

TABLE 2. ARC-JET FLOW FIELD PROPERTIES

	Case 1	Case 2
Mach number, M_∞	10	8
Reynolds number, Re_∞	380	553
Freestream speed, \bar{V}_∞ (cm/sec)	423 000	412 000
Freestream temperature, \bar{T}_∞ (K)	270	414
Freestream density, $\bar{\rho}_\infty$ (gm/cm ³)	3.28×10^{-8}	6.41×10^{-8}
Wall temperature, \bar{T}_w (K)	450	450
Radius of sphere, \bar{r}_b (cm)	5.08	5.08
Species mass fractions in freestream:		
N ₂	0.4138	0.6124
O ₂	0.6519×10^{-5}	0.1084×10^{-4}
N	0.3514	0.1522
O	0.2348	0.2347
NO	0.5380×10^{-5}	0.3761×10^{-4}
NO ⁺	0.2221×10^{-5}	0.6849×10^{-5}
e ⁻	0.3850×10^{-10}	0.1187×10^{-9}

Some high-enthalpy computed data, for which the dissociation and ionization levels are very high, are shown in Figure 2 by the circular symbols. These data points fall considerably below the lower enthalpy data at the continuum end of the flow regime (at large values of K^2) where there is a consistent trend of decreasing C_H with increasing enthalpy for a given K^2 . The parameter K^2 does not account for real gas effects and, therefore, should not be expected to correlate data with widely different enthalpy levels.

Acceptable comparisons have also been obtained with other theoretical predictions by Kumar and Jain [3] who used an analysis identical to the present one except for slip, enthalpy, and viscosity computations.

D. General Results

Figures 3 and 4 illustrate typical convergence behavior of the solutions with increasing number of iterations. Figure 3 shows the successive computed nondimensional temperature profiles, including the initial guess, along the stagnation streamline of a sphere with noncatalytic wall. The temperature profiles did not quite converge by the final iteration (2000), so it was necessary to make a rerun using the output of run No. 1 as input data for run No. 2. The figure illustrates that extremely accurate initial guesses are not generally required to achieve convergence; however, the more accurate the initial guess, the faster the solutions will converge. At very large Reynolds numbers, where gradients in flow properties are large, more accurate initial guesses are required to avoid divergence. Figure 4 shows the convergence behavior of wall heat transfer coefficient, C_H , for the same run. The converged value of C_H , obtained in run No. 2, is shown at the right margin.

Figure 5a gives the computed velocity components, and the pressure, temperature, and density profiles along the stagnation streamline of a sphere with noncatalytic wall at relatively high freestream Reynolds number (1315). Figure 5b gives the mass fraction profiles for the same run. This run is approaching the maximum Reynolds number for which the program will converge easily without extremely accurate input data or modifications to the program such as spacing the computation increments closer together. The meaning of the quantities in Figure 5a are given in equations (18) through (24); e.g. P_1 is the pressure on the stagnation streamline, and P_2 gives the correction in pressure for small angles away from the stagnation streamline. A thick "shock wave" is indicated by the steep gradients in the region $1.06 < r < 1.12$.

The same quantities presented in Figure 5 are plotted in Figure 6 for a smaller Reynolds number (185). Figure 6a reveals that all evidence of a shock wave has disappeared at this low Reynolds number. Comparison of Figures 5b and 6b shows that the degree of dissociation and ionization decreases drastically in lower density flow.

Figure 7 gives the flow profiles along the stagnation streamline of a sphere with fully catalytic wall at a medium Reynolds number (458). The Sutherland viscosity formula was used because this run was made before the

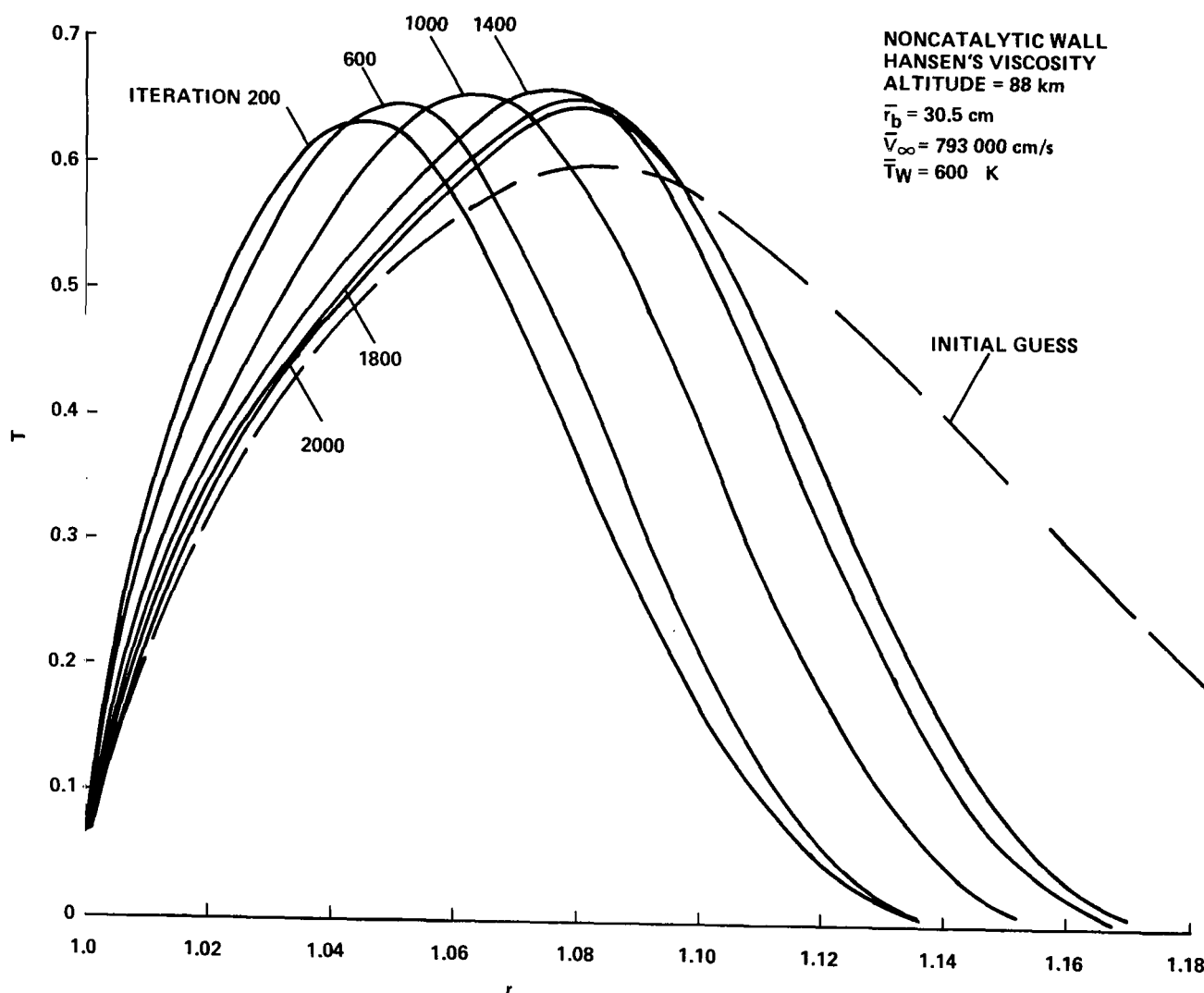


Figure 3. Convergence of temperature profiles for sphere with increasing number of iterations.

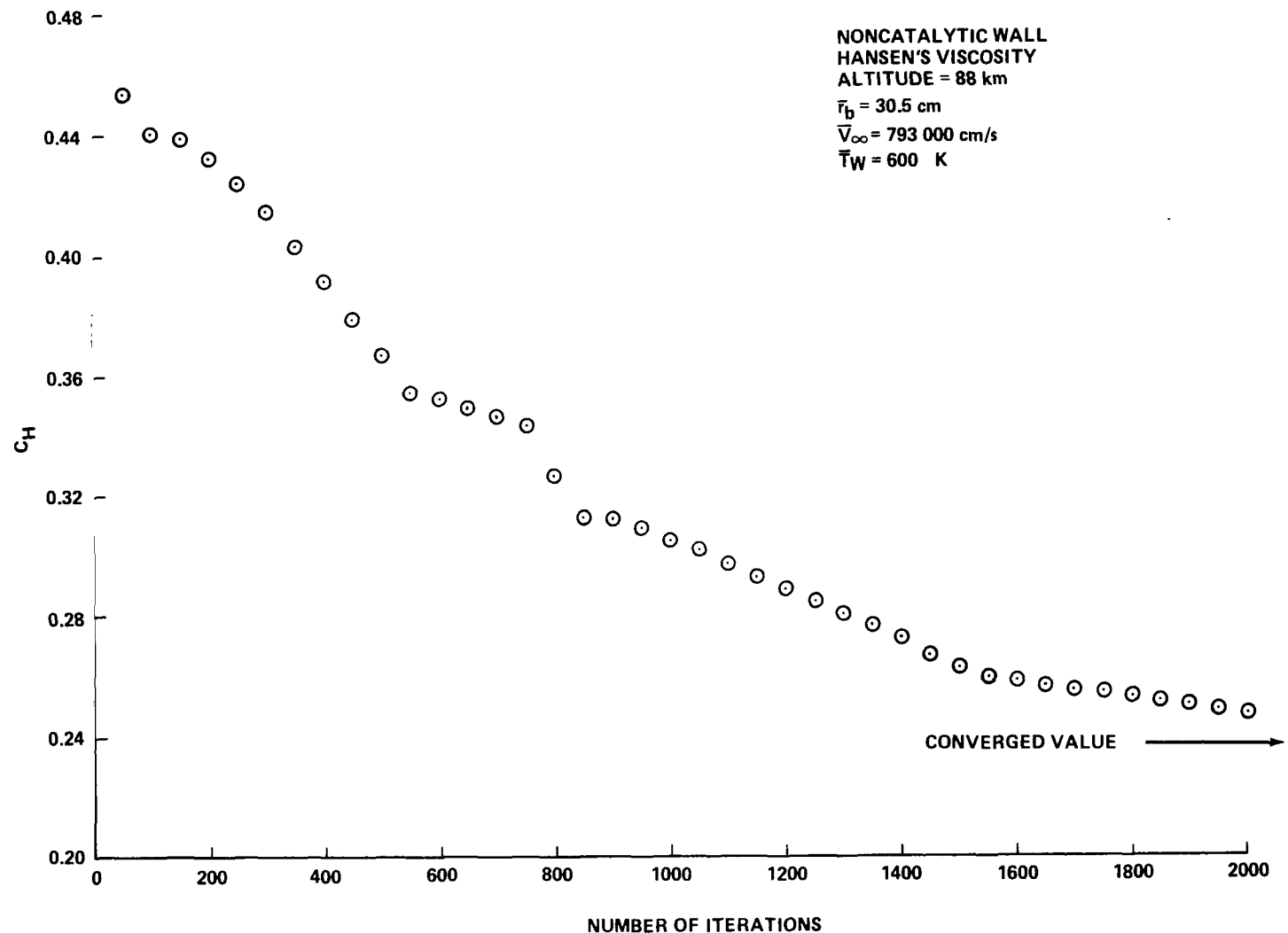
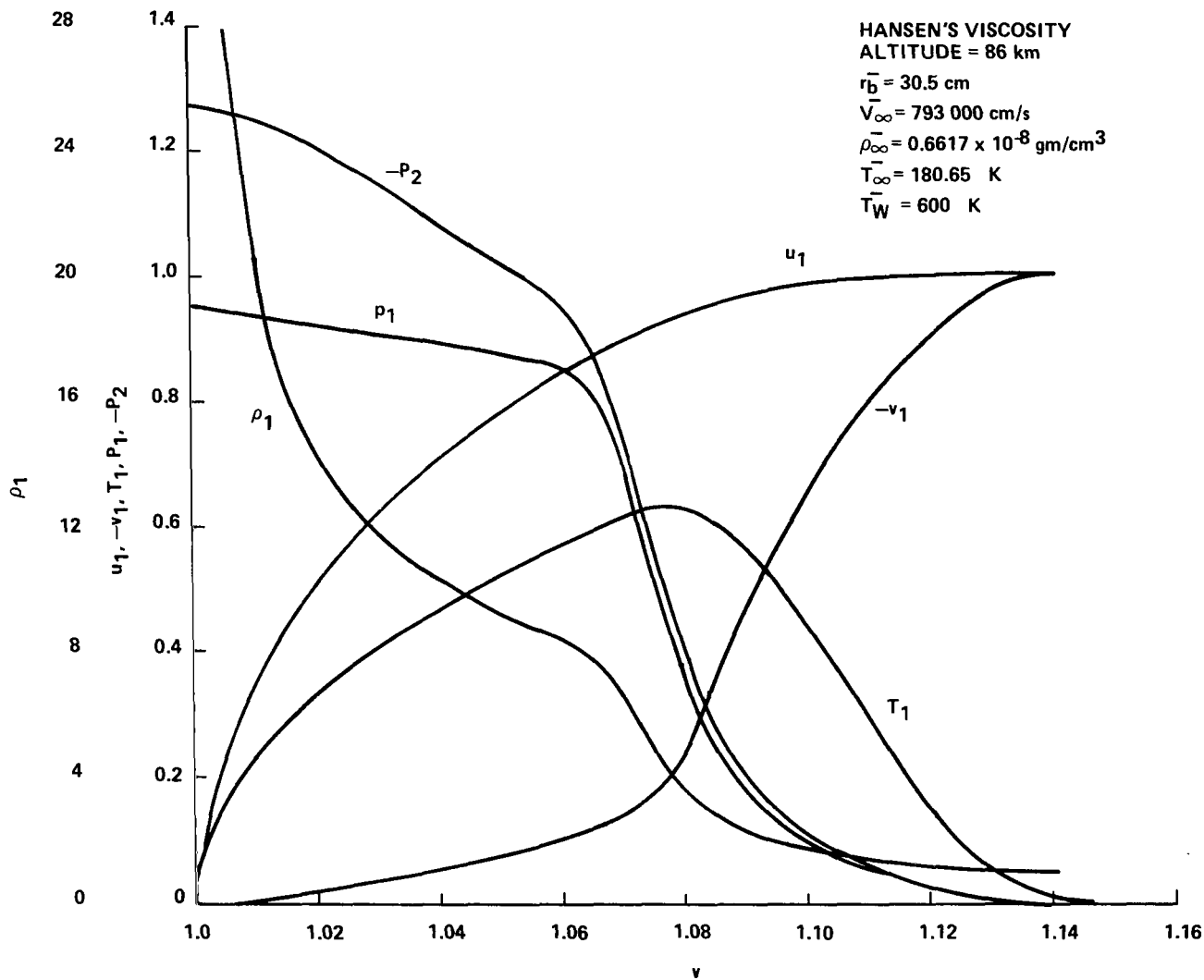
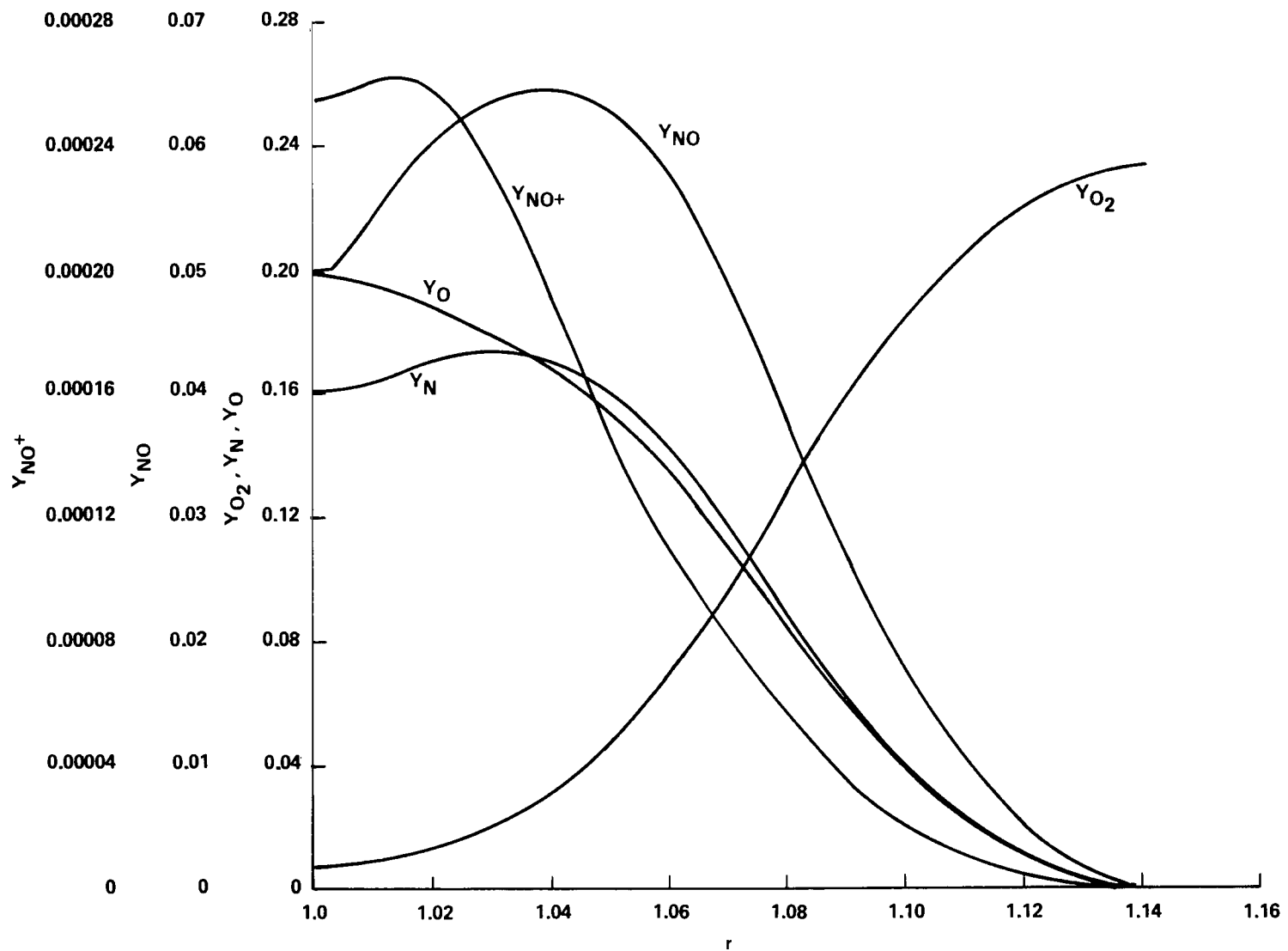


Figure 4. Typical convergence of stagnation point heat transfer coefficient for sphere with increasing number of iterations.



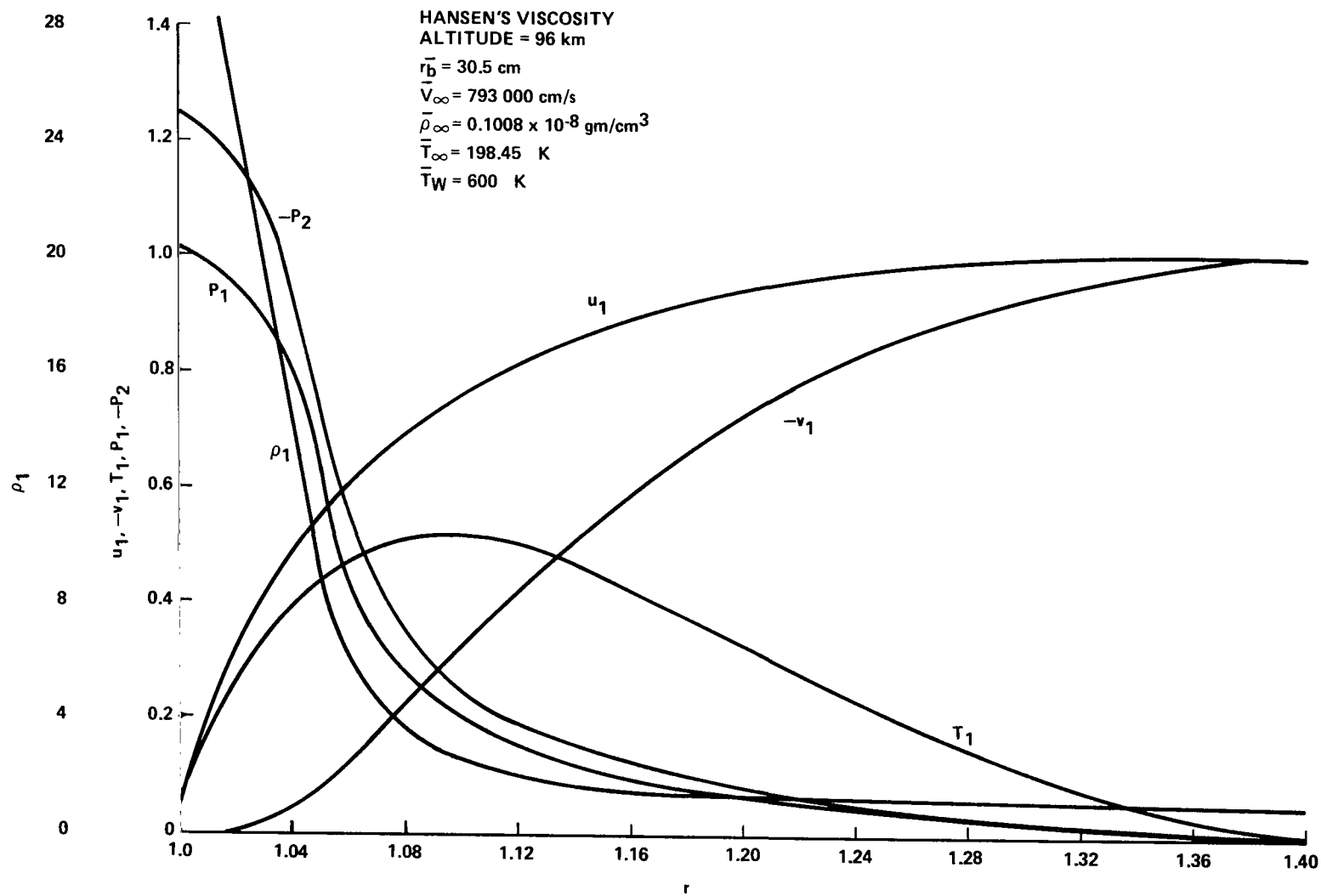
a. Velocity and thermodynamic property profiles.

Figure 5. Flow profiles for sphere with noncatalytic wall at large Reynolds number ($Re_\infty = 1315$).



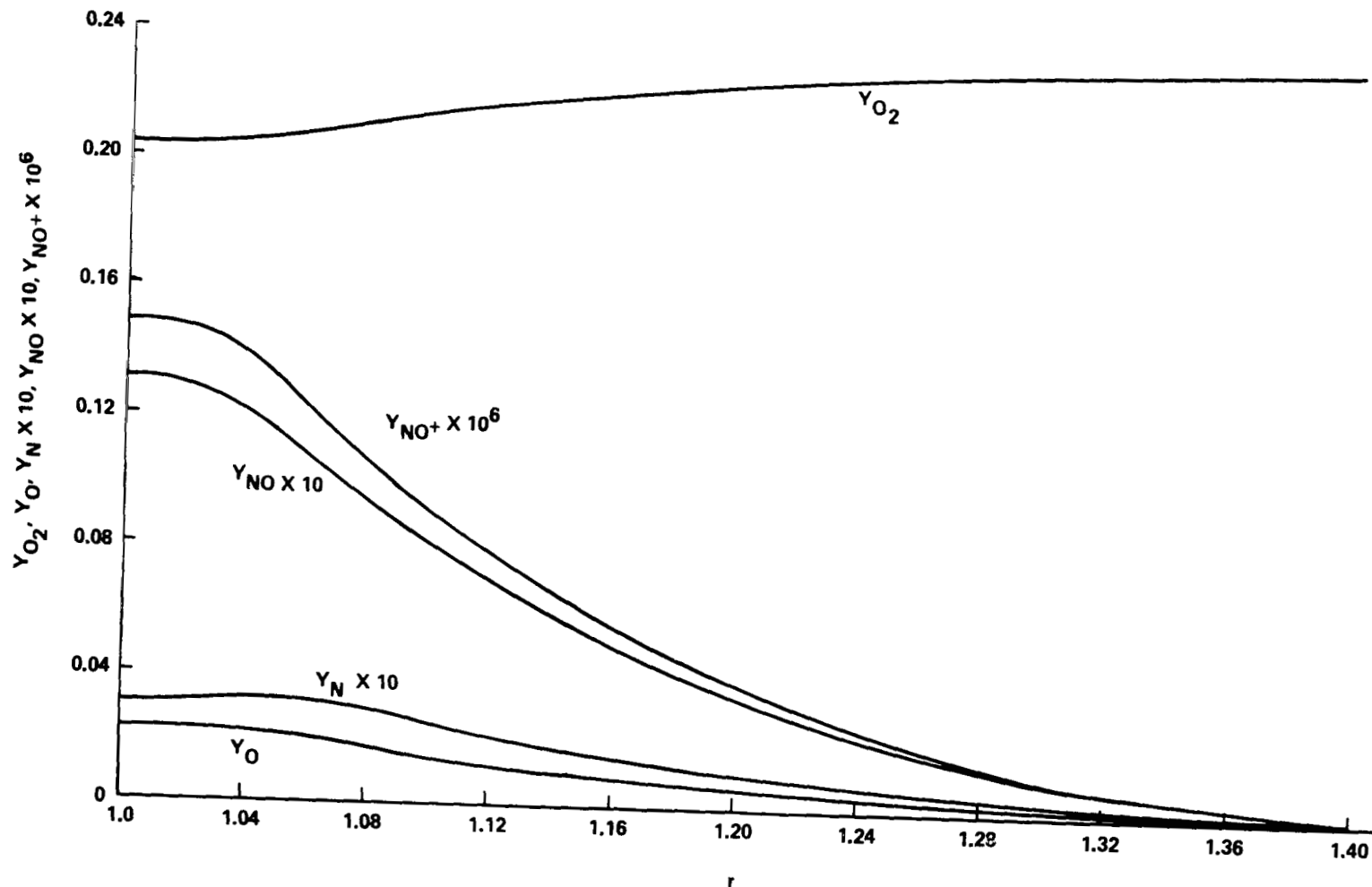
b. Mass fraction profiles.

Figure 5. (Concluded)



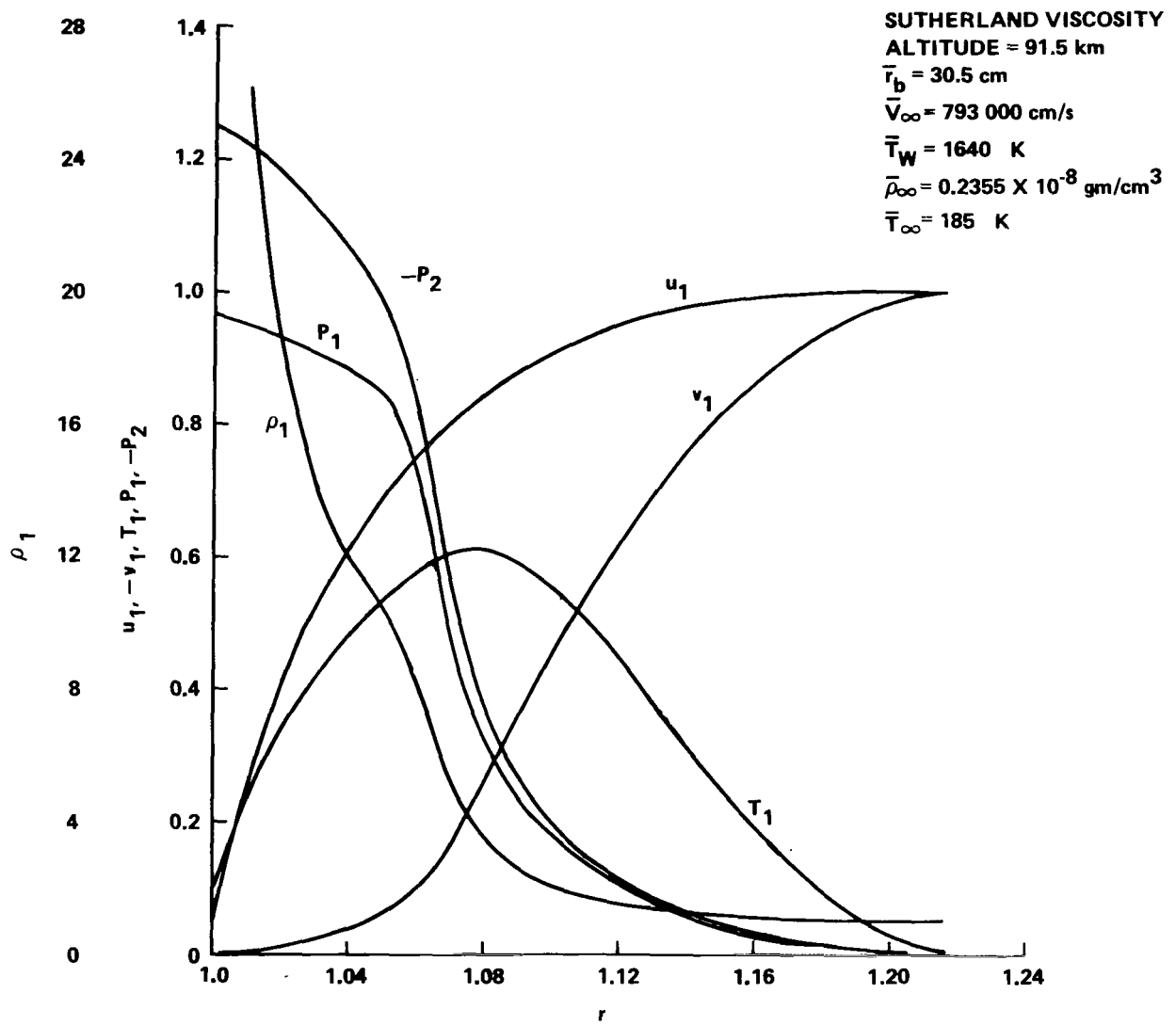
a. Velocity and thermodynamic property profiles.

Figure 6. Flow profiles for sphere with noncatalytic wall at small Reynolds number ($Re_\infty = 185$).



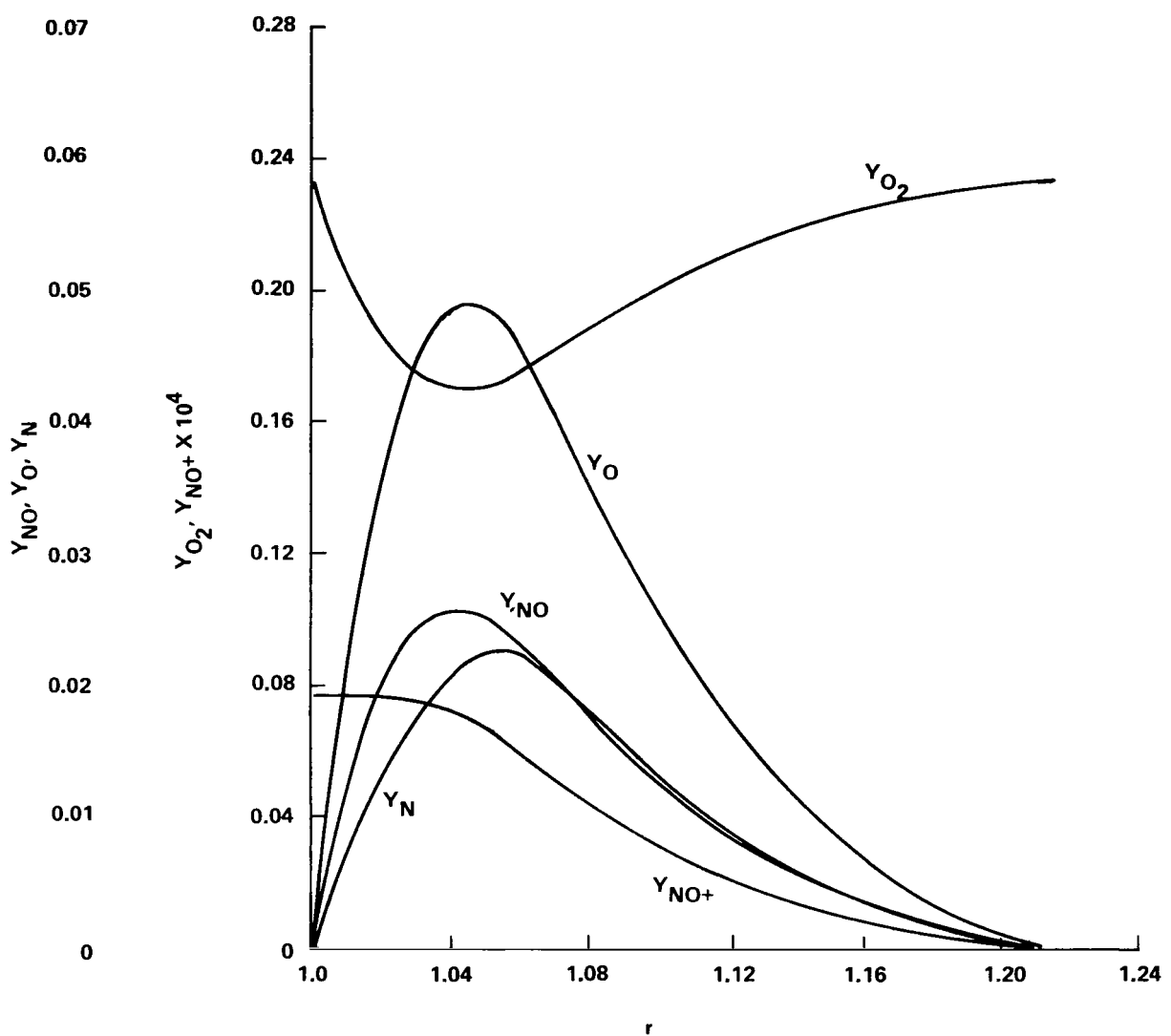
b. Mass fraction profiles.

Figure 6. (Concluded)



a. Velocity and thermodynamic property profiles.

Figure 7. Flow profiles for sphere with fully catalytic wall ($Re_\infty = 458$).



b. Mass fraction profiles.

Figure 7. (Concluded)

viscosity modification to the program. The density profile gives a hint of a shock wave similar to that shown in Figure 5a. Comparison of Figures 5b, 6b, and 7b illustrate the difference in species mass fraction profiles between a noncatalytic wall and a fully catalytic wall. The species mass fractions approach their freestream values at the catalytic wall except for the ions NO^+ and e^- . The wall is assumed noncatalytic with respect to ion recombination in this program.

Figures 5 through 7 provide useful starting data for running the program at conditions near the ones in these figures.

Figure 8 compares temperature, density, and velocity profiles for a sphere and cylinder at the same flow conditions as in Figure 5. The profiles are remarkably similar. Therefore, the profiles for a sphere, Figures 5, 6, and 7, are adequate to use as starting data for the cylindrical option of the program.

The change in the stagnation line temperature profile with freestream density, or Re_∞ , is illustrated in Figure 9. The peak value in nondimensional temperature decreases and the shock layer thickness increases with increasing altitude or decreasing density.

The freestream density and speed greatly affect the degree of dissociation of the air molecules and, hence, the viscosity of the gas mixture. Figure 10 gives the peak values from the viscosity and mass fraction profiles for N, O, and NO as a function of altitude for a fixed large value of freestream speed. As before, the freestream thermodynamic properties associated with altitude are obtained from Reference 14. Although the stagnation enthalpy is large, the figure shows that dissociation becomes negligible and viscosity approaches the Sutherland value at altitudes greater than approximately 100 km. This is due to the decreased reaction rates at the lower temperatures which occur in the shock layer at higher altitudes (Fig. 9). Figure 11 illustrates that dissociation and the viscosity ratio increase with increasing freestream speed at the fixed altitude of 96 km.

A parametric study was made to determine the effect of altitude (or density), freestream speed, and wall temperature on the stagnation point heat transfer coefficient, C_H . Figure 12 gives C_H for a sphere and cylinder as a function of altitude with fixed freestream speed and wall temperature. Hansen's viscosity, equation (48), was used for the sphere, but Sutherland's viscosity was used for the cylinder because those runs were made before the program was modified to compute Hansen's viscosity. The figure shows a steady increase

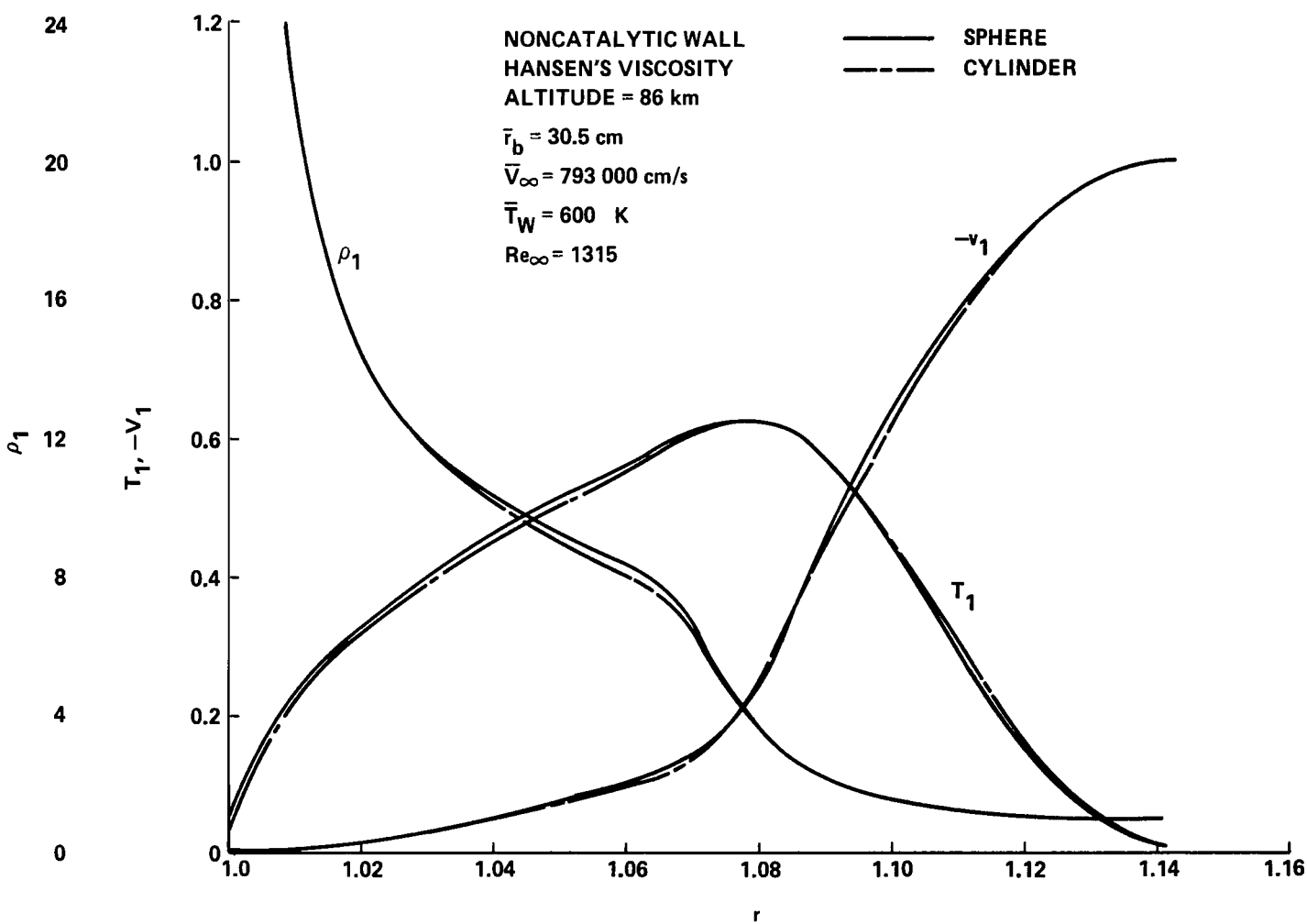


Figure 8. Comparison of flow profiles for sphere and cylinder.

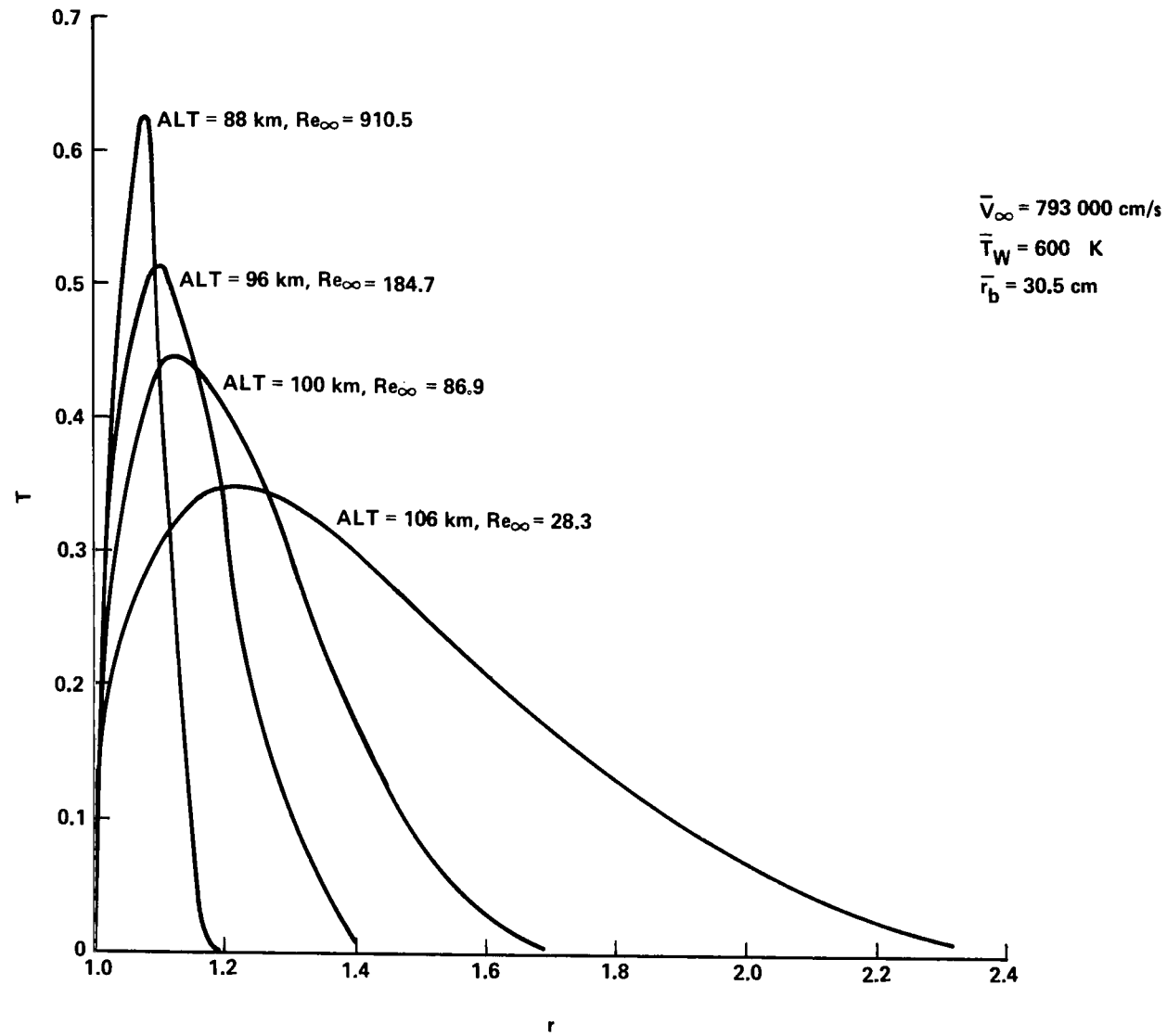


Figure 9. Temperature profiles at various Reynolds numbers for sphere.

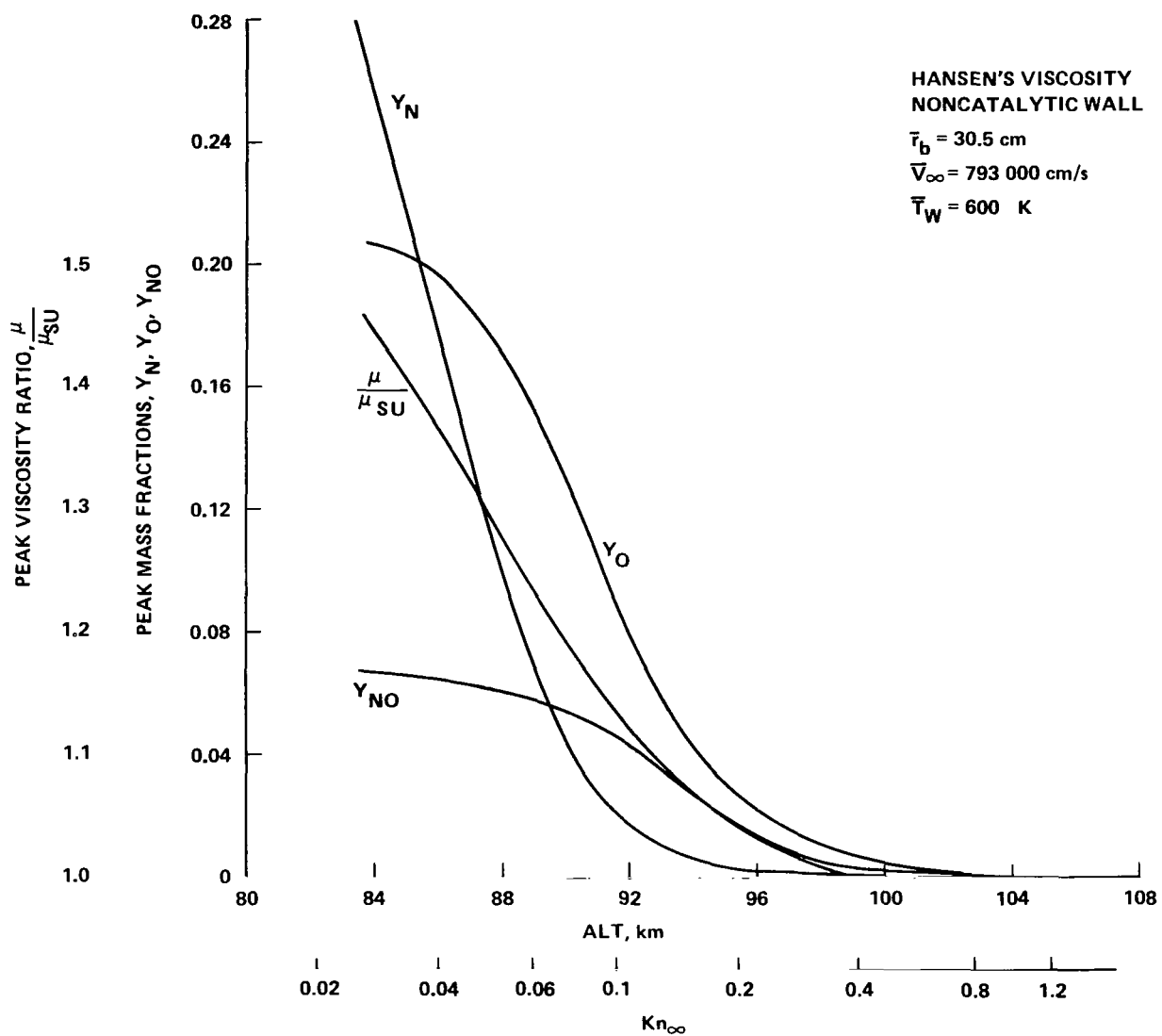


Figure 10. Effect of variation in altitude on dissociation and viscosity in shock layer of sphere.

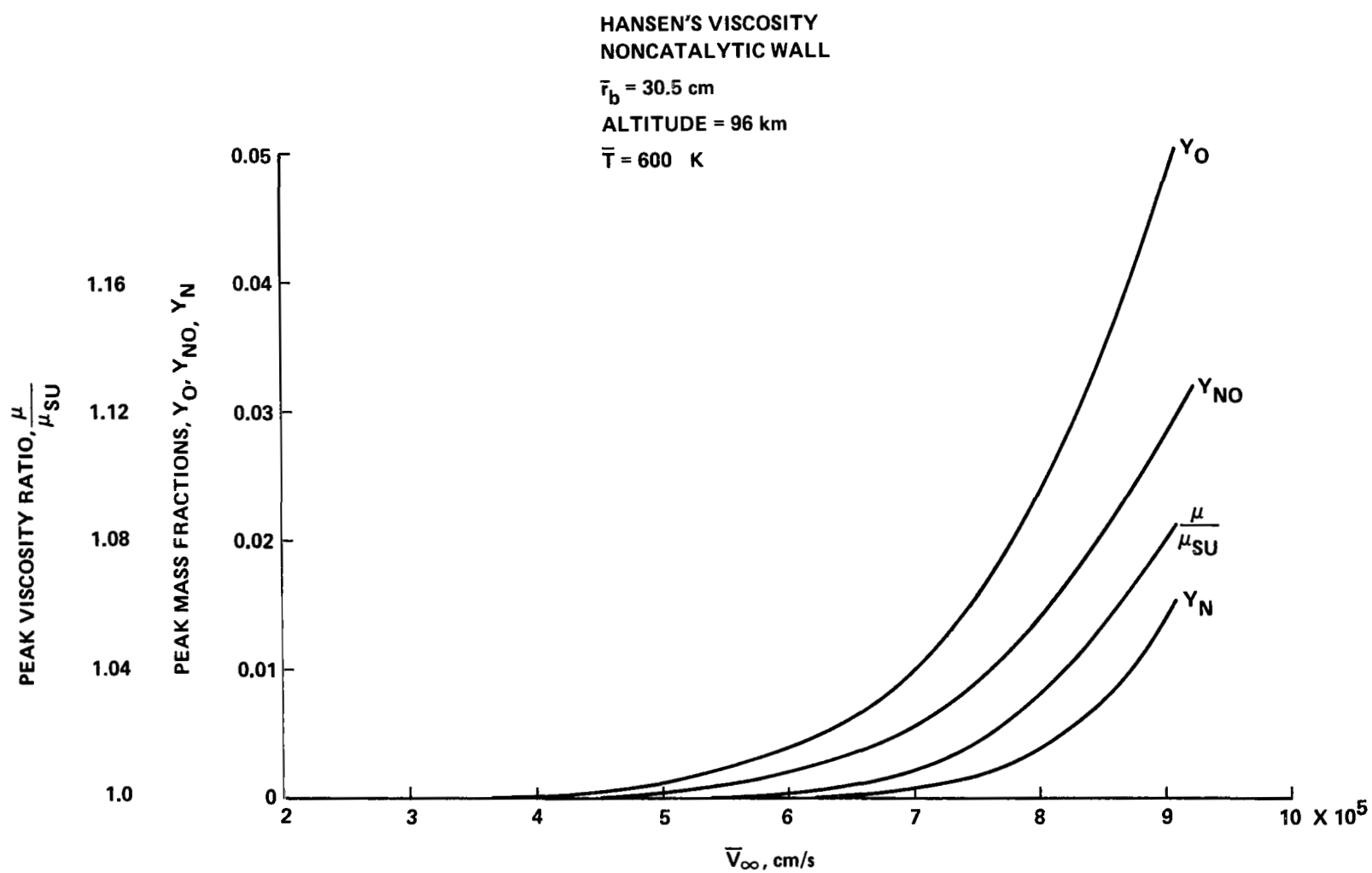


Figure 11. Effect of freestream speed on dissociation and viscosity in shock layer of sphere.

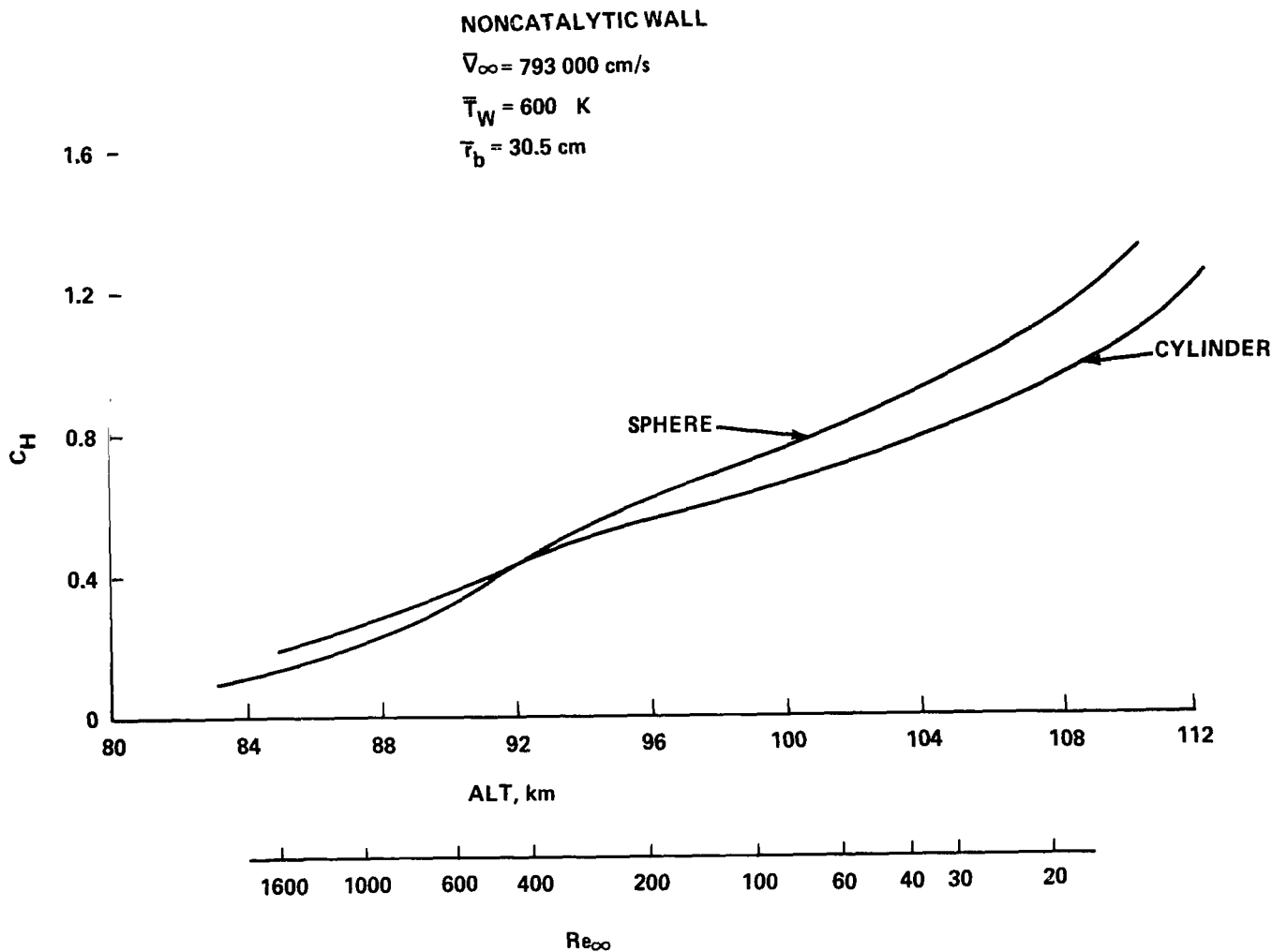


Figure 12. Stagnation point heat transfer coefficients for spheres and cylinders as a function of altitude.

in C_H with increasing altitude. The theoretical free molecular value ($C_H = 1.0$) is exceeded for the sphere at approximately 106 km and for the cylinder at approximately 109 km. The curves continue to climb with increasing slope at higher altitudes. This unrealistic behavior for a fixed wall temperature might be attributed to a breakdown in the continuum approach (Navier-Stokes equations with slip boundary conditions) at extremely high altitudes and low densities. In particular, the assumption of a thin Knudsen layer, which is implicit in the present analysis, becomes invalid at very low densities. One must also consider the practicality of a specified T_w in rarefied flow. This breakdown is gradual, and one cannot pinpoint a sharp boundary beyond which the program gives invalid results. However, the results should be treated with increasing skepticism, for this body size and flow conditions, at altitudes greater than approximately 104 km, or $Re_\infty < 40$.

An interesting feature of Figure 12 is the crossover of the curves for the sphere and cylinder. This crossover should not be attributed to the differences in viscosity computation because the effects of Hansen's viscosity (used for the sphere but not for the cylinder) is to increase C_H at low altitudes but not to affect C_H at high altitudes where dissociation is negligible. Therefore, if Hansen's viscosity were also used for the cylinder, the crossover of the two curves should be expected at a higher altitude. The explanation for this crossover is a result of the stronger merged shock layer effect on the sphere.

The variation of C_H with freestream speed for a sphere at fixed altitude (96 km) and wall temperature (600 K) is shown in Figure 13. The effect of increasing freestream speed, or stagnation enthalpy, is to decrease C_H .

Figure 14 gives the variation in C_H with wall temperature for a sphere at fixed altitude (96 km) and speed (793 000 cm/sec). An increase in wall temperature produces an increase in C_H .

The data in Figures 12 through 14 are replotted in Figure 15 as a function of K^2 ; the similarity parameter is given by equation (62). This parameter includes all the independent variables in Figures 12, 13, and 14, but the effect of freestream density is obviously dominant in determining the value of K^2 and C_H . The data were also plotted as a function of other similarity parameters (not shown), but K^2 correlated the data as well as, or better than, any of the other parameters. This is not the case if wall catalysis is varied. Therefore, K^2 was selected as the similarity parameter to use in presenting the remainder of the data.

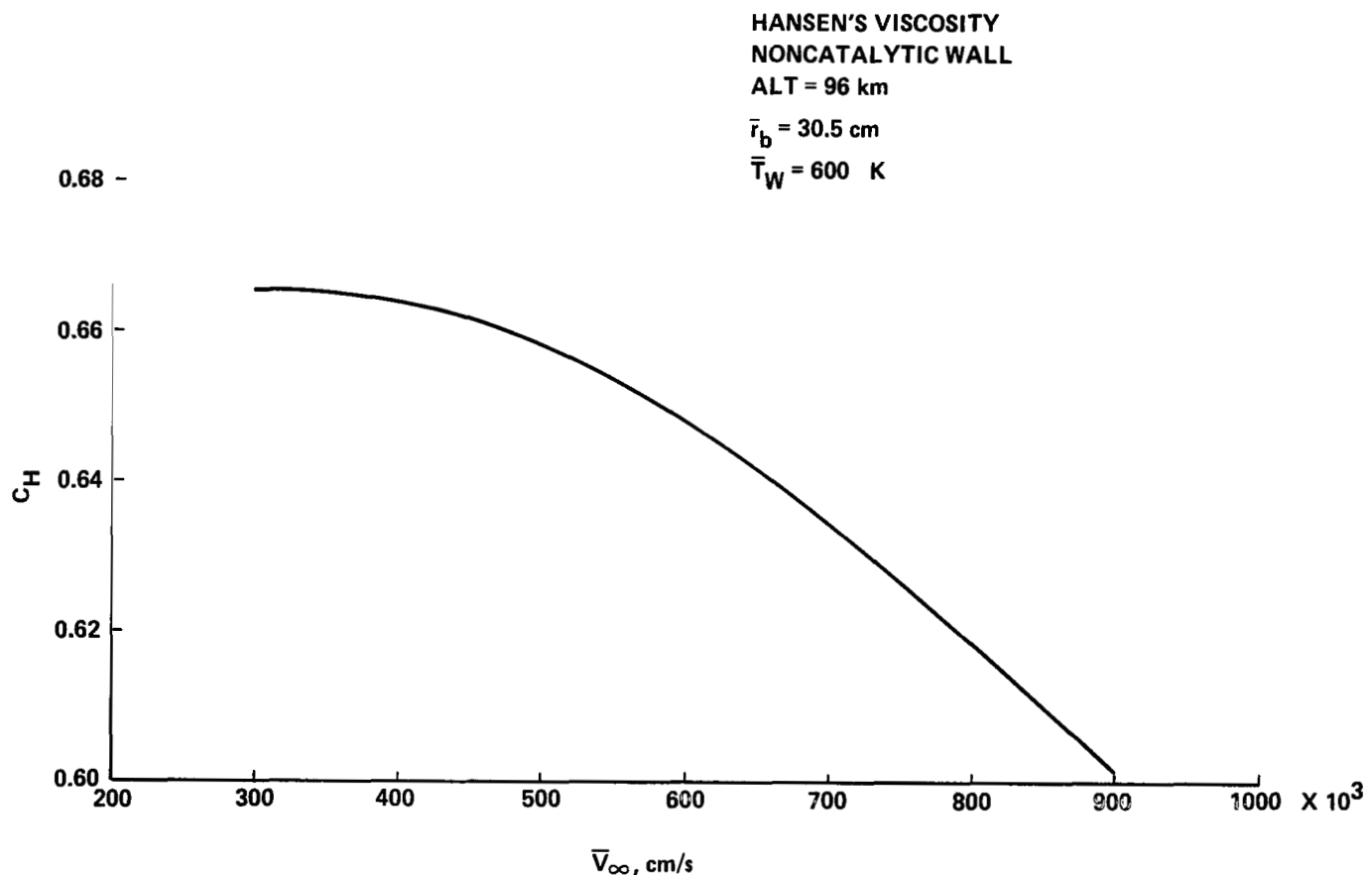


Figure 13. Stagnation point heat transfer coefficient for sphere as function of freestream speed.

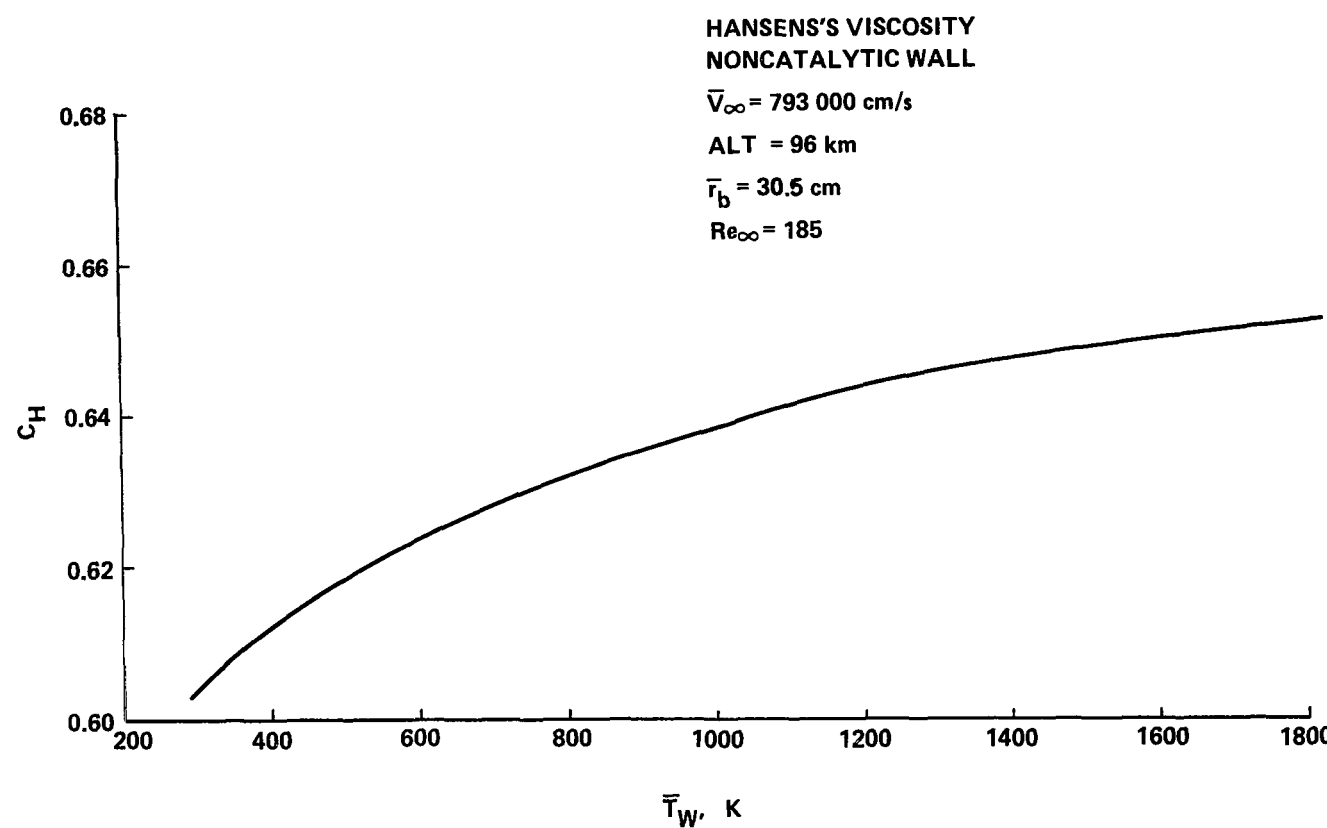


Figure 14. Stagnation point heat transfer coefficient for sphere as function of wall temperature.

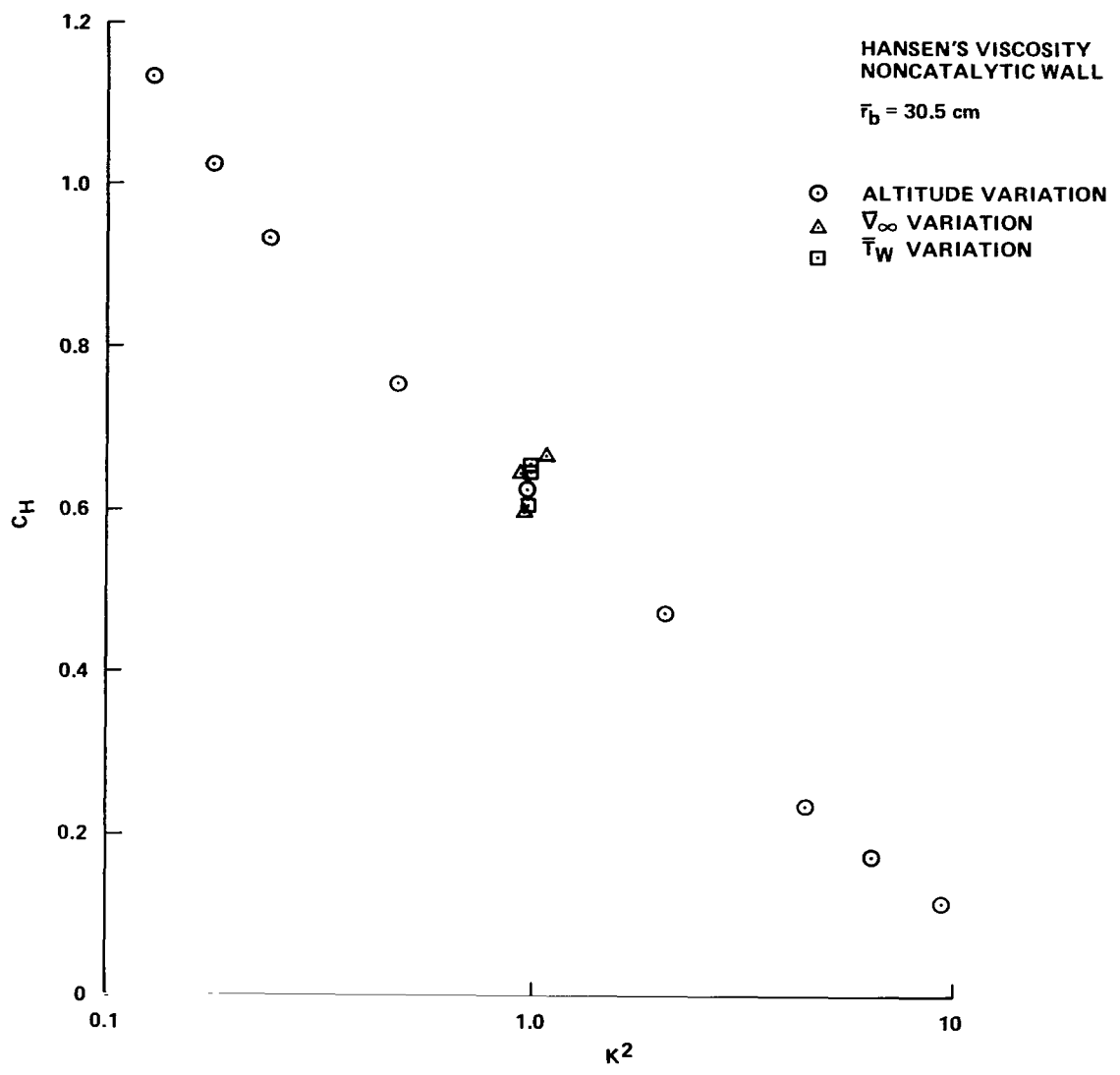


Figure 15. Stagnation point heat transfer coefficient for sphere as function of K^2 .

Figure 16 gives the nondimensional shock layer thickness, n_∞ , for spheres and cylinders as a function of K^2 . The shock layer is thin at the continuum end (large K^2) of the flow regime and becomes very thick at highly rarefied conditions (small K^2).

Figure 17 presents the computed slip speed for a sphere and cylinder as a function of K^2 . The slip speed is computed at the outer edge of the Knudsen layer, but it is applied at the wall in the outer flow solution, i.e., a thin Knudsen layer is assumed. This is a source of error at high altitudes where the Knudsen layer thickness becomes significant. The slip speed is greatly affected by variation in freestream speed and wall temperature. However, these effects are not correlated well by the parameter K^2 as shown for the sphere by the partially filled symbols. To estimate the slip speed for given conditions, one can extrapolate from the solid curves using the partially filled symbols as a guide in correcting for freestream speed and wall temperature. Once again it appears that the proper T_W must be used for rarefied flow calculations. The decrease in slip speed with increasing altitude (decreasing K^2) at high altitude (small K^2) seems unrealistic and has been criticized. However, decreasing slip speed decreases the C_H computed by this program. Since the computed C_H is too large at high altitudes for a fixed T_W (Figs. 12 and 15), the decrease in slip speed at high altitudes tends to compensate for the overprediction of C_H and is, from a practical standpoint, a favorable phenomenon.

The temperature at the edge of the Knudsen layer is given as a function of K^2 in Figure 18. This temperature also depends strongly on freestream speed and wall temperature.

The information in Figures 15 through 18 is useful as input data in running the computer program (Appendix A).

An example of the practical use of this computer program is given in Figure 19. Stagnation point heating rates for the spherical nose of the Space Shuttle External Tank are given as a function of trajectory time. The solid curve is the heating rate predicted by the MSFC Thermal Environment Branch (ED33) and is based on theoretical continuum methods at low altitudes and experimental data at high altitudes. The filled diamond symbols were computed using the present program. The trajectory time interval from approximately 275 to 475 sec corresponds to an altitude interval which is too high for this program, i.e., the predicted values of C_H were unrealistically large. For a

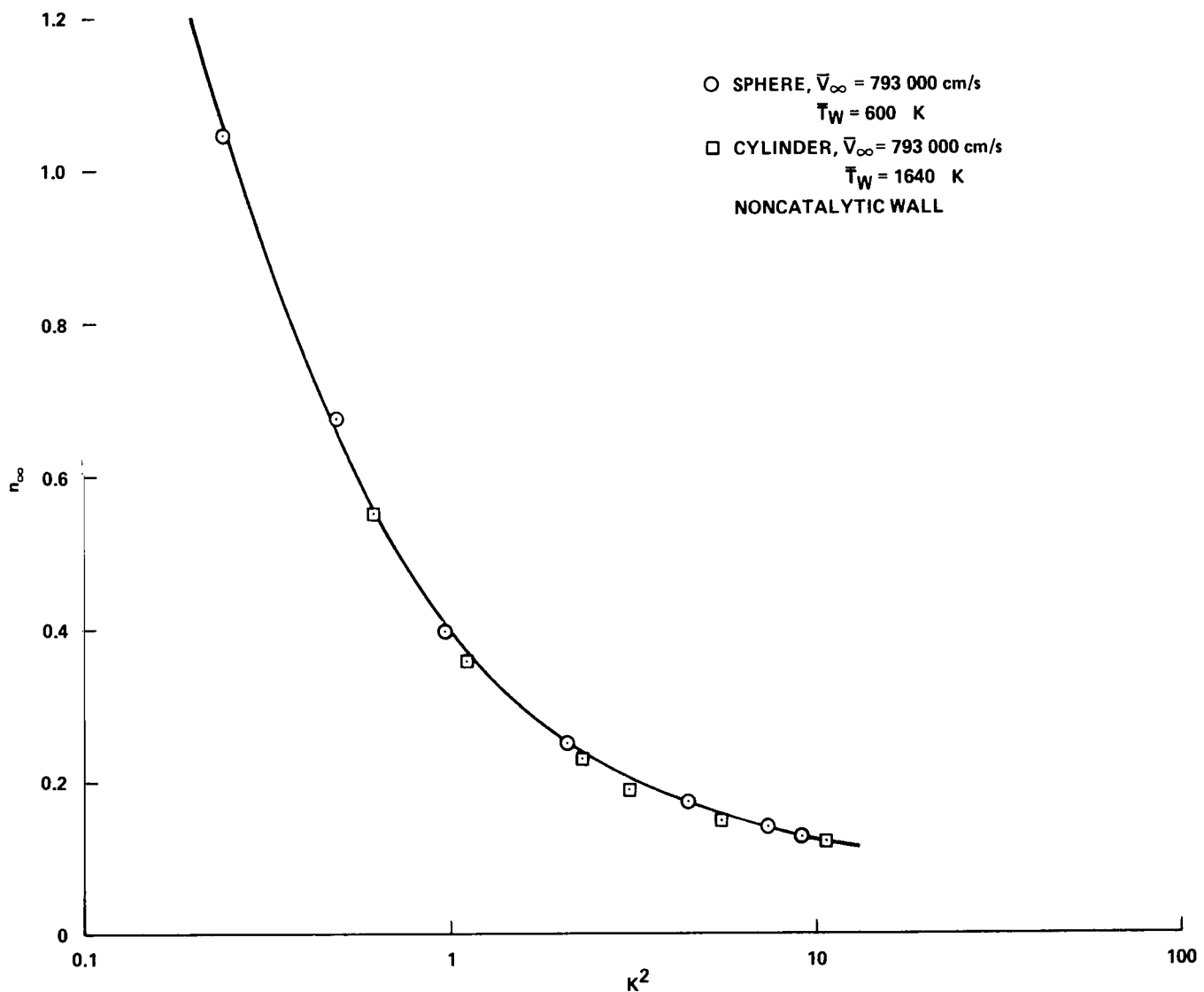


Figure 16. Shock layer thickness for sphere and cylinder.

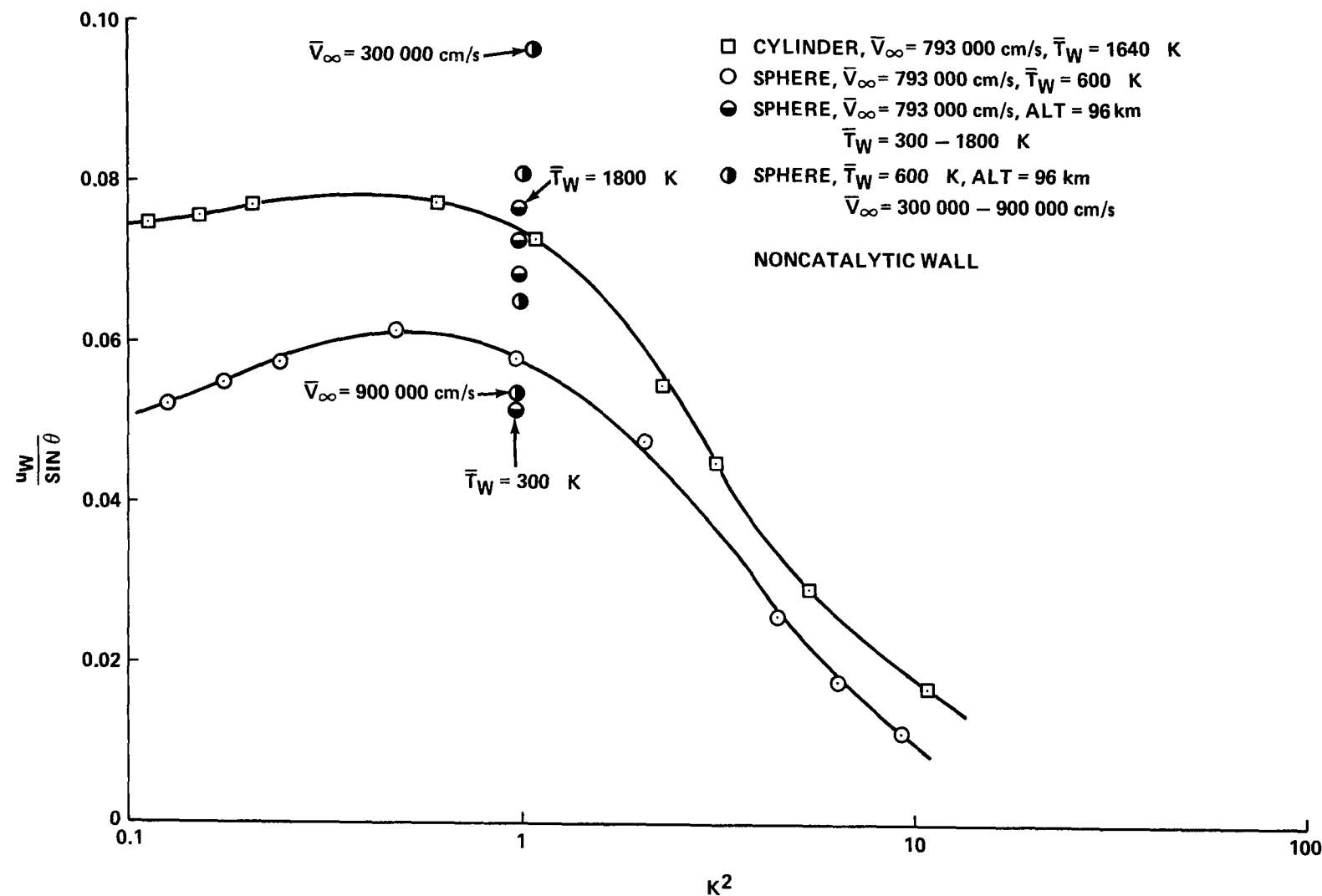


Figure 17. Slip speed at body surface for sphere and cylinder.

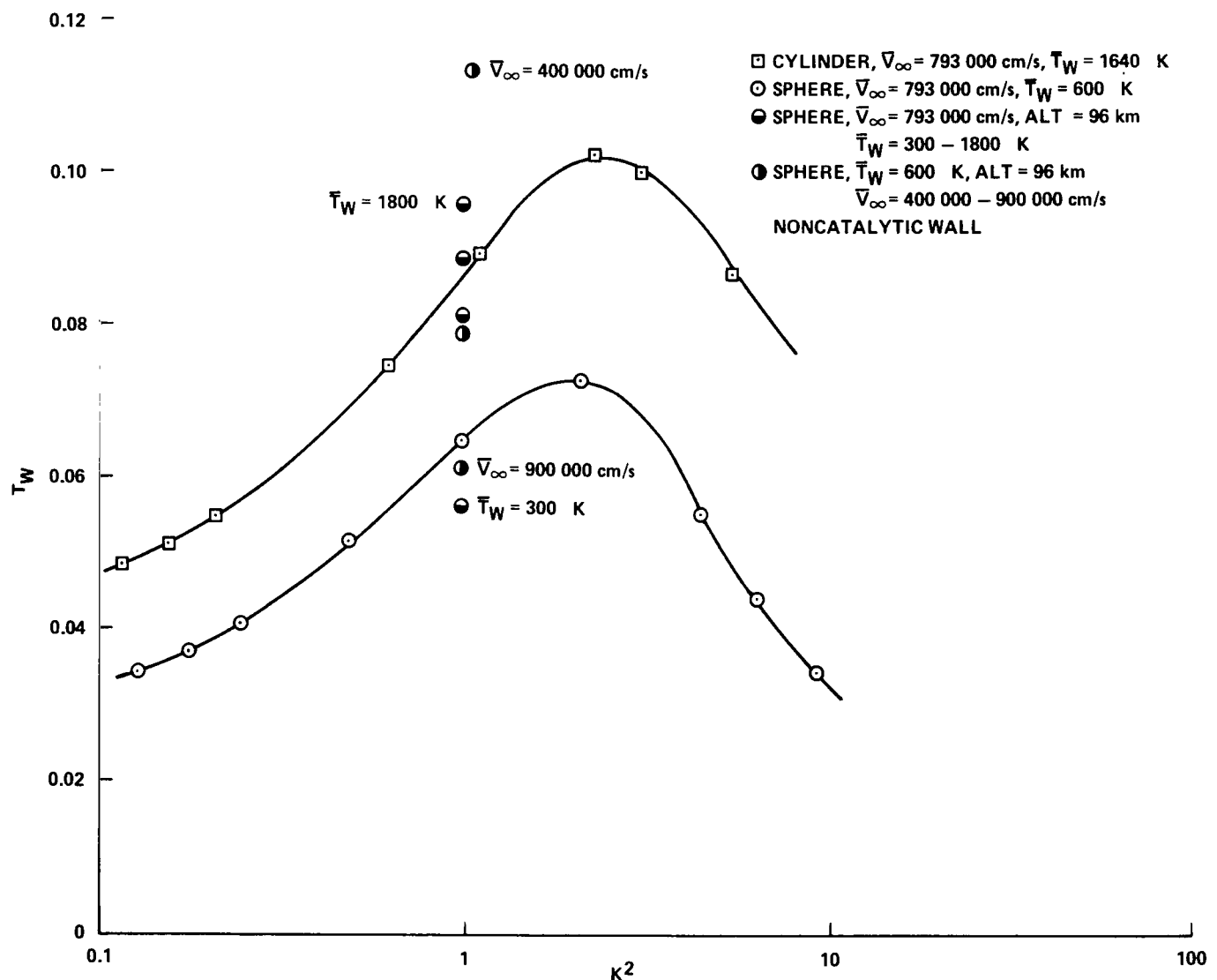


Figure 18. Temperature of air at wall for sphere and cylinder.

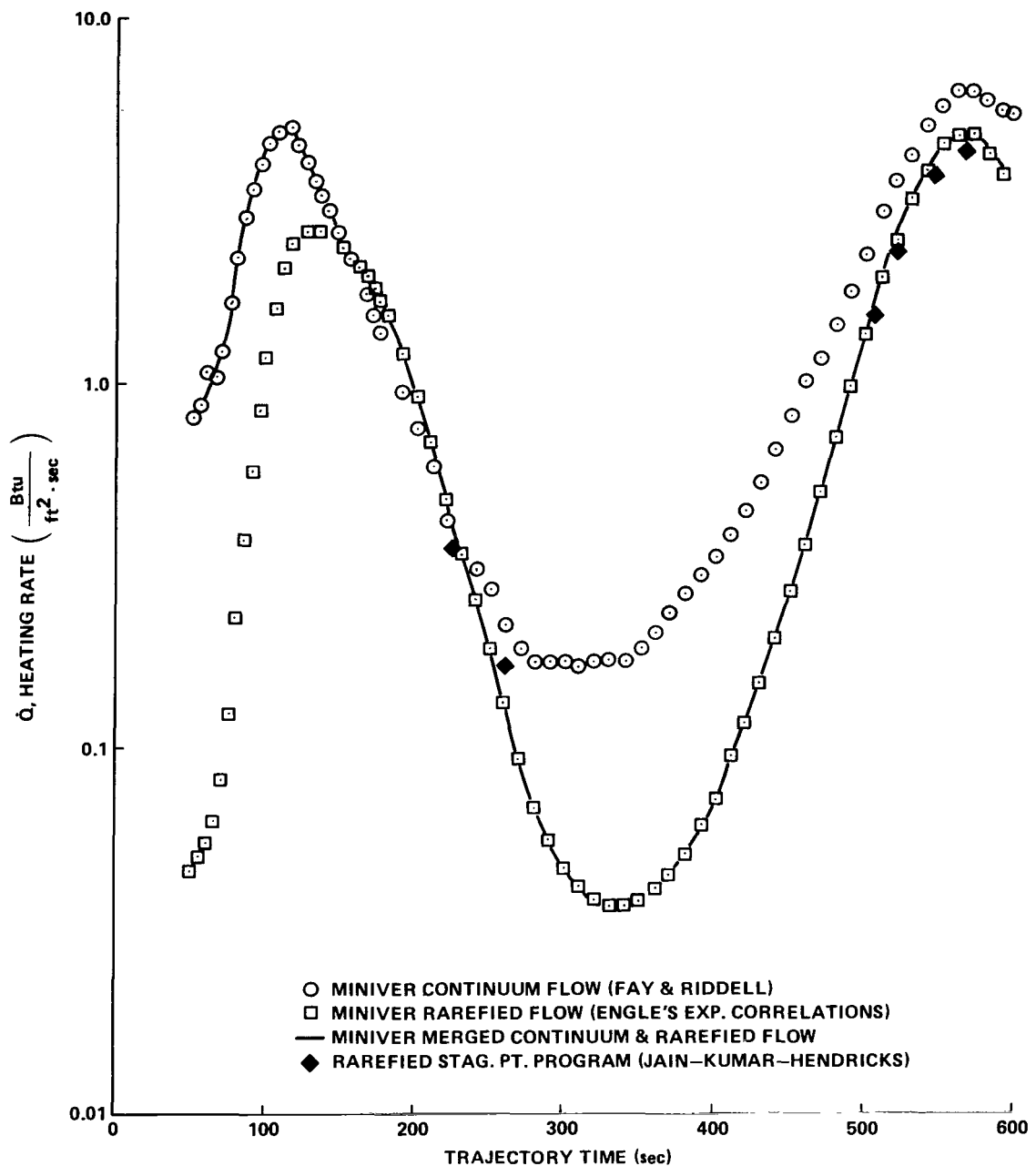


Figure 19. Space Shuttle external tank stagnation point heating rate.

body of this size ($\bar{r}_b = 30.5$ cm), the altitude interval for which this program is applicable is approximately 86 to 105 km. For this altitude interval, the predicted values from this program agree quite well with the predictions from experimental data for a known T_w .

APPENDIX A. COMPUTER PROGRAM

APPENDIX A. COMPUTER PROGRAM

General Information

The computer program is written in Fortran IV language, and it is run on the Univac 1108 computer at MSFC. As currently written, the program is limited to a maximum of 2000 iterations. If convergence is not achieved in 2000 iterations, the program is rerun using the output from the first run as starting data for the second run.

The program has three options:

- (1) Body: sphere or two-dimensional circular cylinder
- (2) Wall catalyticity: noncatalytic wall or fully catalytic wall
- (3) Viscosity: Sutherland or Hansen's high temperature model.

The choices made in options (1) and (2) make very little difference in the computer time used. However, the Hansen viscosity option takes significantly more computer time than the Sutherland option; e.g., if the full 2000 iterations are used, the Sutherland option takes approximately 8 min and the Hansen option takes about 10 min.

Program Input

The quantities needed for input to the program are defined and their functions described as follows:

Computer Program Variable	Symbol Used in Report	Dimensions	Description	Suggested Source or Range of Values
EFFR1	n_∞	Dimensionless	Initial guess for nondimensional shock layer thickness	Figure 16
TOL		Dimensionless	Computation is stopped when C_H converges within a prescribed tolerance. If $ C_{H_N} - C_{H_{N-1}} < TOL$, stop program and print results	0.00001 - 0.0001
EPSI		Dimensionless	Maximum change allowed in all computed dimensional quantities from one iteration to the next	0.001 - 0.005

Computer Program Variable	Symbol Used in Report	Dimensions	Description	Suggested Source or Range of Values
ALT		km	Altitude corresponding to the freestream thermodynamic properties (used for identification only)	
RADB	r_b	cm	Radius of body	
UFS	\bar{V}_∞	cm/s	Freestream speed	
TFS	\bar{T}_∞	K	Freestream temperature	
TWK	\bar{T}_W	K	Temperature of body surface	
ROFS	$\bar{\rho}_\infty$	gm/cm ³	Freestream density	
U(1)	$u_{1w} = u_W / \sin\theta$	Dimensionless	Initial guess for $(\text{slip speed}) / \sin\theta$	Figure 17
T(1)	T_{1w}	Dimensionless	Initial guess for nondimensional temperature of gas at outer edge of Knudsen layer	Figure 18
U(I)	u_1	Dimensionless	Initial guess for u_1 (see equation 18) at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
V(I)	v_1	Dimensionless	Initial guess for v_1 (see equation 19) at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
T(1)	T_1	Dimensionless	Initial guess for T_1 (see equation 20) at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
RO(I)	ρ_1	Dimensionless	Initial guess for ρ_1 (see equation 21) at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
P2(I)	P_2	Dimensionless	Initial guess for P_2 (see equation 22) at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
CO2(I)	y_{O_2}	Dimensionless	Initial guess for mass fraction of O_2 at location I ($I = 2, 3, \dots, 50$)	Figures 5-7

Computer Program Variable	Symbol Used in Report	Dimensions	Description	Suggested Source or Range of Values
CNOI(I)	Y_{NO^+}	Dimensionless	Initial guess for mass fraction of NO^+ at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
CNO(I)	Y_{NO}	Dimensionless	Initial guess for mass fraction of NO at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
CN(I)	Y_N	Dimensionless	Initial guess for mass fraction of N at location I ($I = 2, 3, \dots, 50$)	Figures 5-7
CO(I)	Y_O	Dimensionless	Initial guess for mass fraction of O at location I ($I = 2, 3, \dots, 50$)	Figures 5-7

The quantities $U(I)$ through $CO(I)$ must be input for each of the 49 equally-spaced points in the shock layer along the stagnation streamline starting with $I = 2$ near the wall and ending with $I = 50$ near the freestream (see Fig. A-1). The wall ($I = 1$) and freestream ($I = 51$) boundary conditions are given by the quantities UFS through T(1).

The format for the input data is as follows:

Card	Format	Variable Name(s)	Location and Description
1	12A6	AB1, ..., AB/2	Columns 1-72 contain a comment for project identification
2	3F10.0	XBOD	Columns 1-10 contain the body option (sphere = 1.0, cylinder = 2.0)
		XCAT	Columns 11-20 contain the wall catalyticity option (noncatalytic wall = 1.0, fully catalytic wall)
		XVIS	Columns 21-30 contain the viscosity option (Sutherland viscosity = 1.0, Hansen viscosity = 2.0)
3	2F10.0	CASE	Columns 1-10 contain the run number. This identification is useful to distinguish between runs whenever a rerun is necessary

<u>Card</u>	<u>Format</u>	<u>Variable Name(s)</u>	<u>Location and Description</u>
		EFFR1	Columns 11-20
4	2F10.0	TOL	Columns 1-10
		EPSI	Columns 11-20
5	5F10.0	ALT	Columns 1-10
		RADB	Columns 11-20
		UFS	Columns 21-30
		TFS	Columns 31-40
		TWK	Columns 41-50
6	E13.8	ROFS	Columns 1-13
7	2F10.0	U(1)	Columns 1-10
		T(1)	Columns 11-20
8-56	5E16.9	U(I)	Columns 1-16
		V(I)	Columns 17-32
		T(I)	Columns 33-48
		RO(I)	Columns 49-64
		P2(I)	Columns 65-80
57-105	5E16.9	CO2(I)	Columns 1-16
		CNOI(i)	Columns 17-32
		CNO(I)	Columns 33-48
		CN(I)	Columns 49-64
		CO(I)	Columns 65-80

PROGRAM OUTPUT

The program printout is described in this section. Figure 5 of the text was obtained from this printout. Explanation of the printout follows.

Title Page

Options used — The options which were selected for body, wall catalyticity, and viscosity are printed out here.

Input data — The input data are printed out for record.

Computed freestream data — Some freestream quantities which are used for nondimensionalizing and for computation of various similarity parameters are printed here. The equations used are as follows:

$$VSOUND = \sqrt{\gamma \bar{R} \bar{T}_{\infty}} \quad [\text{cm/sec}] \quad , \quad (\text{A-1})$$

where

$$\gamma = 1.4$$

$$\bar{R} = 2.8708 \times 10^6 \text{ cm}^2/\text{sec}^2\text{K} \quad .$$

$$M_{\infty} = AMACH = \frac{UFS}{VSOUND} \quad (\text{A-2})$$

$$\bar{T}_{0\infty} = STAGFS = \bar{T}_{\infty} \left(1 + \frac{\gamma - 1}{2} M_{\infty}^2 \right) \quad [{}^{\circ}\text{K}] \quad (\text{A-3})$$

$$\bar{\mu}_{\infty} = RTFS = \frac{1.458 \times 10^{-5} \bar{T}_{\infty}}{\bar{T}_{\infty} + 110.4} \quad [\text{gm/cm}\cdot\text{sec}] \quad (\text{A-4})$$

$$\bar{\mu}_{0\infty} = RTSTAG = \frac{1.458 \times 10^{-5} \bar{T}_{0\infty}}{\bar{T}_{0\infty} + 110.4} \quad [\text{gm/cm}\cdot\text{sec}] . \quad (\text{A-5})$$

It is recognized that the stagnation temperature, equation (A-3), has little physical meaning at the high stagnation enthalpies for which this computer program is intended. This is because equation (A-3) is based on the assumption of constant specific heat for temperatures up to $\bar{T}_{0\infty}$; the assumption of constant specific heat is violated at moderate temperatures. However, $\bar{T}_{0\infty}$ is used only for nondimensionalizing and for computing $\bar{\mu}_{0\infty}$. It does not influence the computation of the flow field in any way. Likewise, $\bar{\mu}_{0\infty}$ is a fictitious viscosity because $\bar{T}_{0\infty}$ is fictitious and also because the Sutherland formula is not applicable at high temperatures.

Similarity Parameters — Some similarity parameters are printed out here. Several of these have been used, with some success, in correlating hypersonic or low density data. The similarity parameters are defined as follows:

$$Re_{\infty} = REYF = \frac{\bar{\rho}_{\infty} \bar{V}_{\infty} \bar{r}_b}{\bar{\mu}_{\infty}} , \quad (A-6)$$

$$Re_{0\infty} = REYN = \frac{\bar{\rho}_{\infty} \bar{V}_{\infty} \bar{r}_b}{\bar{\mu}_{0\infty}} , \quad (A-7)$$

$$Re_w = REWALL = \frac{\bar{\rho}_{\infty} \bar{V}_{\infty} \bar{r}_b}{\bar{\mu}_w} , \quad (A-8)$$

where $\bar{\mu}_w$ is computed by the Sutherland formula using \bar{T}_w .

$$K_{n_{\infty}} = XKNFS = \frac{\bar{\lambda}_{\infty}}{\bar{r}_b} , \quad (A-9)$$

$$\bar{V} = V_{\text{BAR}} = M_{\infty} \sqrt{\frac{C_{\infty}}{Re_{\infty}}} , \quad (A-10)$$

where

$$C_{\infty} = \frac{\bar{\mu}_w}{\bar{\mu}_{\infty}} \frac{\bar{T}_{\infty}}{\bar{T}_w} .$$

$$K^2 = XKSQ = \frac{Re_{\infty}}{\gamma_{\infty} M_{\infty}^2 C_*} , \quad (A-11)$$

where

$$C_* = \frac{\bar{\mu}_*}{\bar{\mu}_{\infty}} \frac{\bar{T}_{\infty}}{\bar{T}_*}$$

and

$$T_* = \frac{1}{2} (\bar{T}_{0\infty} + \bar{T}_w)$$

$\bar{\mu}_*$ = Sutherland viscosity computed at \bar{T}_* .

The similarity parameter ϕ is given as the final item in the printout. It is placed in this location because it uses the computed enthalpy at the wall. This parameter was suggested by Potter (Rarefied Gas Dynamics, Supplement 5, Vol. I, 1969).

$$\phi = PHI = Re_w \left(\frac{2\bar{h}_w}{\bar{V}_{\infty}^2} \right)^{0.6} . \quad (A-12)$$

Data Pages

A set of computed data is printed out after every 200 iterations. Each set of data consists of three pages.

1. First Page. The computed shock layer thickness (EFFR) and wall heat transfer coefficient (C_H) are printed at the top of the page after every 50 iterations. The control factors which are used to update the computed quantities to obtain input variables for the next iteration are also printed here.

The columns on this page give nondimensional values of (1) radial position from the center of the body to the equally-spaced points at which the independent variables are computed, (2) the velocity component u_1 [see equation (18)], (3) the velocity component v_1 [see equation (19)], (4) specific enthalpy, (5) temperature, (6) viscosity ratio, local ratio of Hansen viscosity to Sutherland viscosity, (7) pressure P_1 [see equation (22)], (8) pressure correction P_2 [see equation (22)], and (9) density.

Note that VFS is the same as UFS elsewhere in the program. It is the freestream speed \bar{V}_∞ .

2. Second Page. This page gives the reaction rates for each of the seven species.

3. Third Page. This page gives the mass fractions of the chemical species which compose the air. Note that mass fraction was denoted by "Y" in the text rather than by "C." The electron number density is also given.

When C_H converges within the prescribed tolerance, or when the iterations reach 2000, the iterations are stopped and the final data set is printed out. At this point a set of 98 cards is punched to record the velocity and thermodynamic data given on page 1 and the mass fraction data on page 3 of the data printout. This set of cards can then be used as starting data for a new run.

To determine whether or not the program has converged sufficiently and whether the results are reasonable, the following checks are suggested.

(1) Check EFFR for convergence

(2) Check C_H for convergence

(3) Check the slip conditions for convergence, i.e. compare U_1 and T at $RAD/RADB = 1.0$ for the last few sets of data

(4) Scan the velocity component and thermodynamic property columns to determine if the profiles are smooth and the values are reasonable

(5) Scan the mass fraction columns to check for reasonable behavior. Particularly check the N_2 column near the wall to determine if these mass fractions exceed the freestream value.

SOME DIFFICULTIES ENCOUNTERED IN RUNNING PROGRAM

The program generally runs without much difficulty if the starting data are within reasonable bounds; however, problems do occur, and some of the most common problems are discussed as follows.

1. Failure of EFFR to Converge. The shock layer thickness, EFFR, usually is the first quantity to converge; however, it sometimes diverges. The most common cause of this is using starting data from a run with a much different stagnation enthalpy, or $\bar{T}_{0\infty}$, from the case being run. This causes a discontinuity in the nondimensional temperature profile between the points $I = 50$ and $I = 51$ because the point $I = 50$ comes from the starting data from a previous run and point $I = 51$ is the freestream boundary condition. The decision on whether to decrease, increase, or not change EFFR is made on the basis of the temperature gradient near the freestream (see Appendix B). Therefore, if a sizable discontinuity in temperature exists at this location, EFFR will diverge.

One solution to this problem is to modify the last few input cards to make a smooth transition in the temperature profile to the freestream temperature. Probably a better solution would be to change the nondimensionalizing temperature from $T_{0\infty}$ to \bar{T}_∞ . This would guarantee a smooth transition, in the starting data, to the freestream temperature. To prevent divergence of EFFR, the program has been modified to restrict EFFR within the range

$$\frac{1}{2} \text{EFFR1} \leq \text{EFFR} \leq \frac{3}{2} \text{EFFR1} .$$

If EFFR1 is not guessed closely enough, this restriction will necessitate a rerun using a different EFFR1.

2. Unrealistic Chemical Composition. If the species mass fractions in the starting data are greatly different from the true values for the case being run, unrealistic computed mass fractions can result. This can lead to unrealistic computed values for all other quantities. For example, this might occur for a rather low speed, high density case in which dissociation should be negligible (Fig. 11). However, since starting data from a low speed, high density run were not available, data were used from a high speed, high density run in which the degree of dissociation was high. The program could not handle the grossly incorrect starting data, and the computed results became more and more unrealistic from one iteration to the next. The unrealistic data included mass fractions for N₂ much larger than the freestream value, negative C_H (heat transferred from the body to the gas) and a monotonically increasing EFFR beyond reasonable bounds (Fig. 16). Note that C_H was called "STANC" in the older printout.

SOME SUGGESTIONS FOR RUNNING PROGRAM

To minimize the previously discussed problems and to facilitate operation of the program, the following suggestions on input data are made.

1. EFFR1, U(1), T(1), and TOL. Figures 16, 17, and 18 can be used for obtaining initial guesses for EFFR1, U(1), and T(1), respectively. The value for TOL probably should be set smaller than the desired tolerance on C_H to assure that the other quantities converge also. A value of 0.00001 for TOL was used in most of the runs made for this report.

2. Velocity, Thermodynamic Property, and Mass Fraction Profiles. For best results, the starting data profiles for the quantities U(I) through CO(I) should be obtained from a run which matches the required freestream conditions of the new run as closely as possible. Close matching is especially important at low altitude, high density conditions for which gradients in the flow properties are large. The question arises as to what constitutes a

sufficiently close "match" and which matching, or similarity, parameter should be used. If every freestream and body condition is matched fairly closely, it is unnecessary to use a similarity parameter. From experience, it is found that, for a given body, changes in freestream density, $\bar{\rho}_\infty$, and speed, \bar{V}_∞ , in that order, have the greatest effect on the outcome of the run. Therefore, for bodies of the same size, one should try to match $\bar{\rho}_\infty$ and \bar{V}_∞ . A good rule of thumb is that $\bar{\rho}_\infty$ for the new run should not differ from $\bar{\rho}_\infty$ for the starting data run by more than the equivalent of 3 km altitude in the standard atmosphere. Near the low altitude extreme of the program's capability, this difference should not be more than approximately 1 km. A difference in \bar{V}_∞ of approximately 2000 m/sec can usually be tolerated. For bodies of greatly different size, one should probably use a similarity parameter, such as K^2 , to match starting data to the new run.

Figures 5, 6, and 7 can be used for starting data. If the desired free-stream properties for the new run are farther removed from those of Figures 5, 6, and 7 than the previously suggested increments, it is advisable to reach the desired conditions in two or more runs. If severe difficulties like those previously discussed are encountered, it is probably best to go back to a "good," or converged, set of starting data and use smaller increments in freestream properties rather than to use the output of the "bad" run as starting data for a rerun. If the program is used frequently, a library of starting data card sets can be built up to cover the range of possible freestream conditions for which the program is applicable.

3. Hansen-Sutherland Viscosity Option. If the computed ratio of Hansen's viscosity to Sutherland's viscosity is approximately 1.0 for a given set of freestream conditions, one can use the Sutherland viscosity option for nearby freestream conditions to reduce computer time.

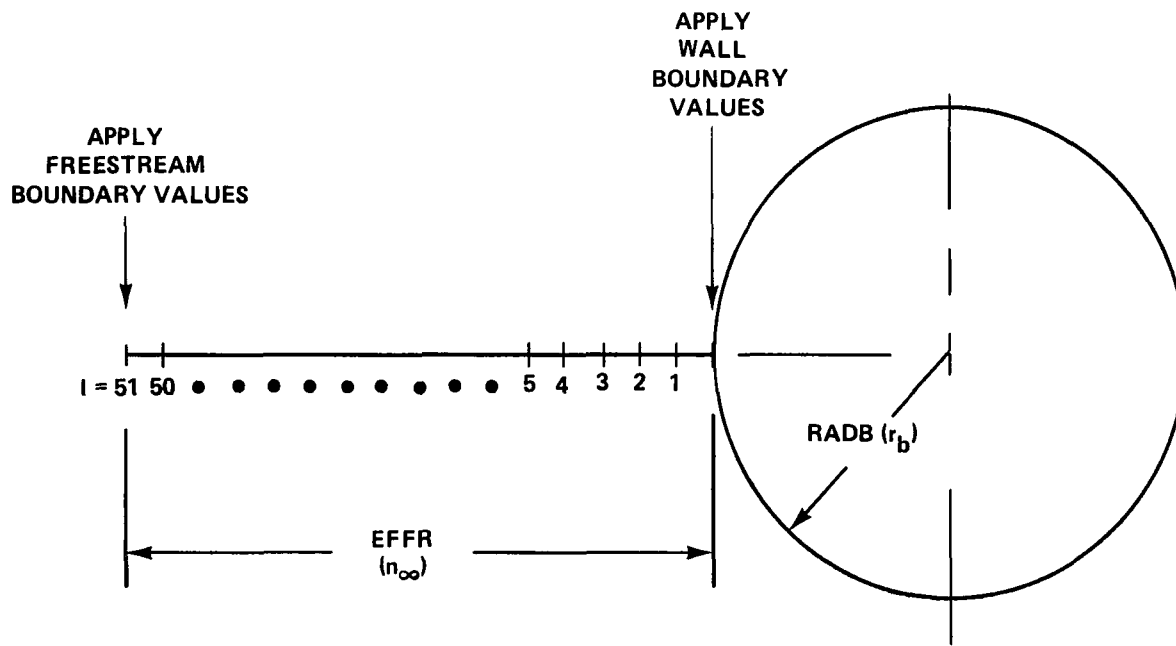


Figure A-1. Location of computation points.

|||

APPENDIX B. LISTING OF PROGRAM

MAIN

DATE 120877

```

00100   1*   C   SUCCESSIVE ACCELERATED REPLACEMENT METHOD          000000
00100   2*   C   NONEQUILIBRIUM FLOW IN THE MERGED SHOCK LAYER OF A BLUNT BODY 000000
00100   3*   C   NEAR THE STAGNATION REGION USING FULL N.S. EQUATIONS 00000C
00100   4*   C   SLIP CONDITIONS AT THE WALL 000000
00100   5*   C   CATALYTIC OR NONCATALYTIC WALL 000000
00100   6*   C   ***** 000000
00100   7*   C   000000
00101   8*   C   DIMENSION U(100),V(100),T(100),H(100),P(100),P2(100),RT(100), 000000
00101   9*   C   1D1(100),RAD(100),X(100),C02(100),CN2(100),CN(100),C0(100), 000000
00101  10*   C   1CNO(100),CCL(100),R0(100) 000000
00103  11*   C   DIMENSION W02(100),WN2(100),WN0(100),WN(100),W0(100),WN0I(100), 000001
00103  12*   C   1WEL(100),PCL(100) 000001
00104  13*   C   DIMENSION HN2(100),H02(100),H0(100),HN(100),HN0(100),HNOI(100) 000001
00105  14*   C   DIMENSION HEL(100) 000001
00106  15*   C   DIMENSION C1(10),C2(10),C3(10),PP1(10),W(10),EL(10) 000001
00106  16*   C   000001
00106  17*   C   INPUT HANSEN'S VISCOSITY DATA 000001
00107  18*   C   DIMENSION XMUSU(100),TAB5(100),XMURAT(100),XMU(100) 000001
00110  19*   C   DIMENSION TEMP(37),TAB1(37),TAB2(37),TAB3(37),TAB4(37) 000001
00111  20*   C   DATA TEMP /2000.0,2500.0,3000.0,3500.0,4000.0,4500.0,5000.0, 000001
00111  21*   C   1.5E0C.0,6000.0,6500.0,7000.0,7500.0,8000.0,8500.C,9000C.0,9500.0, 000001
00111  22*   C   2.10000.C,10500.0,11000.0,11500.0,12000.0,12500.0,13000.0,13500.0, 000001
00111  23*   C   3.14000.C,14500.0,15000.C,15500.0,16000.C,16500.0,17000.C,0,17500.0, 000001
00111  24*   C   4.18000.0,18500.0,19000.0,19500.0,20000.0/ 000001
00113  25*   C   DATA TAB1 /C.886,C.846,C.830,C.815,C.803,C.792,C.782,C.773,C.764, 000001
00113  26*   C   1.0,757,C.750,C.743,C.737,C.731,C.725,C.720,C.715,C.710,C.706, 000001
00113  27*   C   2.0,701,C.697,C.693,C.689,C.685,C.681,C.677,C.672,C.668,C.664, 000001
00113  28*   C   3.0,660,C.656,C.651,C.647,C.643,C.639,C.635,C.630/ 000001
00115  29*   C   DATA TAB2 /C.742,C.742,C.705,C.676,C.650,C.626,C.608,C.591,C.575,C.561, 000001
00115  30*   C   1.0,548,C.536,C.524,C.514,C.504,C.495,C.486,C.478,C.470,C.463, 000001
00115  31*   C   2.0,456,C.448,C.443,C.437,C.431,C.426,C.420,C.415,C.407,C.401, 000001
00115  32*   C   3.0,395,C.389,C.384,C.376,C.371,C.365,C.359,C.352/ 000001
00117  33*   C   DATA TAB3 /C.468,C.457,C.445,C.434,C.423,C.412,C.401,C.397,C.380, 000001
00117  34*   C   1.0,366,C.353,C.342,C.331,C.321,C.313,C.304,C.297,C.290,C.283, 000001
00117  35*   C   2.0,281,C.270,C.266,C.261,C.256,C.252,C.247,C.243,C.236,C.230, 000001
00117  36*   C   3.0,224,C.218,C.213,C.206,C.201,C.196,C.190,C.185/ 000001
00121  37*   C   DATA TAB4 /123.30,122.20,120.80,118.10,114.80,109.70,102.00, 000001
00121  38*   C   1.89,90.75,60.64,50.55,70.48,60.42,80.37,90.33,80.30,40.27,40, 000001
00121  39*   C   2.24,90.22,70.20,80.19,C9.17,60.16,27,15.10,14.04,13.09,12.24, 000001
00121  40*   C   3.11,25,10.38,10.48,10.01,9.50,9.01,8.75,8.01,7.51,7.68/ 000001
00121  41*   C   000001
00121  42*   C   ***** 000001
00121  43*   C   000001
00123  44*   C   READ(5,702)AB1,AB2,AB3,AB4,AB5,AB6,AB7,AB8,AB9,AB10,AB11,AB12 000001
00141  45*   C   702 FORMAT(12A6) 000022
00142  46*   C   READ(5,700) XBOD,XCAT,XVIS 000022
00147  47*   C   READ(5,700)CASE,EFFR1 000032
00153  48*   C   READ(5,700)TOL,EPST 000041
00157  49*   C   READ(5,700)ALT,RADB,UFS,TFS,TWK 000050
00166  50*   C   700 FORMAT(6F10.0) 000062
00167  51*   C   READ(5,701)ROFS 000062
00172  52*   C   701 FORMAT(E13.8) 000070
00173  53*   C   EFFR=EFFR1 000070
00174  54*   C   ICASE=CASE 000072
00174  55*   C   000072

```

MAIN	DATE 120877
00174 56*	000072
00174 57*	000072
00174 58*	000072
00175 59*	000101
00201 60*	000110
00201 61*	000110
00201 62*	000110
00201 63*	000110
00202 54*	000114
00205 65*	000114
00207 66*	000116
0021C 67*	000117
00211 68*	000120
CC212 69*	000122
00213 70*	000123
CC214 71*	000125
00214 72*	000125
00214 73*	000125
00214 74*	000125
00214 75*	000125
00215 76*	000130
00217 77*	000134
00217 78*	000134
00220 79*	000134
00222 80*	000140
00223 81*	000140
00225 82*	000144
00226 83*	000144
00230 84*	000150
00230 85*	000150
00231 86*	000150
00233 87*	000154
00233 88*	000154
00234 89*	000154
00236 90*	000160
00237 91*	000160
00241 92*	000164
00242 93*	000164
00244 94*	000170
00244 95*	000170
00245 96*	000170
00247 97*	000174
00247 98*	000174
00250 99*	000174
00252 100*	000200
00253 101*	000200
00255 102*	000204
00255 103*	000204
00255 104*	000204
00255 105*	000204
00256 106*	000204
00260 107*	000210
00260 108*	000210
00260 109*	000210
00261 110*	000210
00263 111*	000214

***** BOUNDARY CONDITIONS AT THE BODY *****
 READ(5,700)U(1),T(1)
 V(1)=0.

EL REPRESENTS LEWIS NUMBER AND A IS AVAGADRO NUMBER
 DO 1 K=1,7
 1 CL(K)=1.4
 CL1=1.4
 CL2=1.4
 PR=0.75
 GAMMA=1.4
 SC1=PR/CL1
 SC2=PR/CL2

***** PRINT PROGRAM TITLE AND INPUT DATA *****
 PRINT 10
 1C FORMAT(1H1,///25X,"HYPERSONIC, MERGED SHOCK LAYER, STAGNATION LINE")
 1 FLOW FIELD FOR SPHERE OR CIRCULAR CYLINDER)
 PRINT 11
 11 FORMAT(/SOX,"JAIN-KUMAR-HENDRICKS PROGRAM")
 PRINT 12
 12 FORMAT(45X,"(OBTAINED FROM BILL HENDRICKS, JULY 1975)")
 PRINT 13
 13 FORMAT(///40X,"FEATURES: SIMILARITY SOLUTION OF NAVIER-STOKES EQUATIONS")
 PRINT 14
 14 FORMAT(5DX,"NONEQUILIBRIUM CHEMISTRY")
 1 /50X,"SLIP BOUNDARY CONDITIONS")
 PRINT 19
 19 FORMAT(//31X,"AVAILABLE OPTIONS:")
 PRINT 21
 21 FORMAT(50X,35H. BODY: (A) SPHERE OR (B) CYLINDER)
 PRINT 22
 22 FORMAT(50X, "2. WALL CATALYTICITY: (A) NONCATALYTIC WALL OR (B) FULLY CATALYTIC WALL")
 PRINT 23
 23 FORMAT(50X, "3. VISCOSITY: (A) SUTHERLAND OR (B) HANSEN'S HIGH TEMP MODEL")
 1 MP. MODEL(NACA TR R-50)")
 PRINT 24
 24 FORMAT(//35X,"MODIFICATIONS:")
 PRINT 25
 25 FORMAT(50X,"1. 9/2/76 (KEN JOHNSTON) ENTHALPY COMPUTATION MODIFIED
 1 TO ALLOW FOR PARTIALLY EXCITED VIBRATIONAL STATE (VINCENTI AND KRUGER, P.135)
 2 STATEMENTS 30,130, " /53X, "PREVIOUS ASSUMPTION
 3: FULLY EXCITED VIBRATIONAL STATE")
 PRINT 26
 26 FORMAT(//5DX,"2. 9/2/76 (KEN JOHNSTON) OPTION ADDED TO COMPUTE VISCOSITY BY HANSEN'S HIGH TEMP.")
 108* 109* "BECOMES INACCURATE AT HIGH TEMP.")
 PRINT 350
 350 FORMAT(50X,"3. 9/9/76 (KEN JOHNSTON) CATALYTIC AND NONCATALYTIC P

MAIN

DATE 120877

00263	112*	PROGRAMS COMBINED."//55X,"(OPTION AVAILABLE TO CHOOSE DESIRED WALL C	000214
00263	113*	CATALYTIC CONDITION")	000214
00264	114*	PRINT 351	000214
00266	115*	351 FORMAT(/50X,"4. 9/9/76 (KEN JOHNSTON) SPHERE AND CYLINDER PROGRAMS	000220
00266	116*	1 COMBINED."//55X,"(OPTION AVAILABLE TO CHOOSE DESIRED BODY")	000220
00267	117*	PRINT 18,AB1,AB2,AC3,AB4,AB5,AB6,AB7,AB8,AB9,AB10,AB11,AB12	000220
00305	118*	18 FORMAT(//3X, "PROJECT: ",12A6)	000240
00306	119*	PRINT 27	000240
00310	120*	27 FORMAT(13X,"OPTIONS USED:")	000244
00311	121*	IF(XBOD=0.9 .GT. 0.2) GO TO 29	000244
00313	122*	PRINT 28	000251
00315	123*	28 FORMAT(27X,"SPHERE")	000255
00316	124*	GO TO 41	000255
00317	125*	29 PRINT 40	000257
00321	126*	40 FORMAT(27X,"CIRCULAR CYLINDER")	000263
00322	127*	41 IF(XCAT=0.9 .GT. 0.2) GO TO 43	000263
00324	128*	PRINT 42	000267
00326	129*	42 FORMAT(27X,"NONCATALYTIC WALL")	000273
00327	130*	GO TO 45	000273
00330	131*	43 PRINT 44	000275
00332	132*	44 FORMAT(27X,"FULLY CATALYTIC WALL")	000301
00333	133*	45 IF(XVIS=0.9 .GT. 0.9) GO TO 47	000301
00335	134*	PRINT 46	000305
00337	135*	46 FORMAT(27X,"SUTHERLAND VISCOSITY FORMULA")	000311
00340	136*	GO TO 49	000311
00341	137*	47 PRINT 48	000313
00343	139*	48 FORMAT(27X,"HANSEN'S HIGH TEMP VISCOSITY FORMULA")	000317
00344	139*	49 PRINT 15,ICASE,EFFR1,U(1),T(1),TOL,EPSI	000317
00354	140*	15 FORMAT(//15X,"RUN NO.:",I4,	000330
00354	141*	1 //15X,"INPUT DATA"	000330
00354	142*	1 //18X,"EFFECTIVE RADIUS (EFFR) = ",F7.4," (STANDOFF DISTANCE/BO	000330
00354	143*	1DY RADIUS")	000330
00354	144*	2 /36X,"U1(1) = ",F9.5," (U1BAR(1)/FREESTREAM SPEED)"	000330
00354	145*	3 /35X,"T1(1) = ",F8.5," (T1BAR(1)/FREESTREAM STAG. TEMP.)"	000330
00354	146*	4 /4X,"HEAT TRANSFER COEFF. TOLERANCE (TOL) = "F8.6,	000330
00354	147*	5 /37X,"EPSI = ",F8.6,")	000330
00355	149*	PRINT 16,ALT,RADB,UFS,TFS	000330
00363	149*	16 FORMAT(27X,17HALITUDE (ALT) = ,F9.4,4H KM/23X,Z1HBODY RADIUS (R	000340
00363	150*	1ADB) = ,F9.4, 4H CM/19X,25HFREESTREAM SPEED (UFS) = ,F11.3, 8H C	000340
00363	151*	2M/SEC/13X,31HFREESTREAM TEMPERATURE (TFST) = ,F9.4,12H DEG KELVIN)	000340
00364	152*	PRINT 17,ROFS,TWK	000340
00370	153*	17 FORMAT(I6X,"FREESTREAM DENSITY (ROFS) = ",E10.5," GM/CM**3"	000346
00370	154*	1 /19X,"WALL TEMPERATURE (TWK) = ",F9.4," DEG KELVIN")	000346
00370	155*	C *****	000346
00370	156*	C *****	000346
00370	157*	C *****	000346
00370	158*	C COMPUTE AND PRINT FREESTREAM CONDITIONS AND SIMILARITY PARAMETERS	000346
00371	159*	VSCUNDE=100.0*((1.4*287.08*TFST)**0.5	000346
00372	160*	RTFS=(1.458E-05*TFST**1.5)/(TFST+110.4)	000357
00373	161*	AMACH=UFS/VSCOUND	000371
00374	162*	STAGFS=TFST*(1.0+((GAMMA-1.0)/2.0)*AMACH**2)	000374
00375	163*	RTSTAG=(1.458E-05*STAGFS**1.5)/(STAGFS+110.4)	000404
00376	164*	RTWALL=(1.458E-05*TWK**1.5)/(TWK+110.4)	000416
00377	165*	REYF=UFS*ROFS*RADB/RTFS	000430
00400	166*	REYN=REYF*RTFS/RTSTAG	000435
00401	167*	REWALL=REYF*RTFS/RTWALL	000441

MAIN

DATE 120877

```

00402    163*          XLAMDA=(1.2562*RTFS)/(R0FS*(2.8702E+06*TFS)**0.5)      000444
00403    169*          XK1FS=XLAMDA/RAD3                                         000460
00404    173*          CF02= RTWALL*TFS/(RTFS*TWK)                                000462
00405    171*          VEAR= AMACH*(CFS/REYF)**0.5                           000470
00406    172*          TSTAR=C.5*(STAGFS+TWK)                               000500
00407    173*          RTSTAR=(1.4E8E-05*TSTAR**1.5)/(TSTAR+110.4)           000504
00408    174*          CSTAR=RTSTAR*TFS/(RTFS*TSTAR)                            000515
00411    175*          XKSG= REYF/(CAMMA*AMACH**2*CSTAR)                         000522
00412    176*          PRINT 400                                         000530
00414    177*          400 FCFMAT///*COMPUTED FREESTREAM DATA:*/
00415    179*          PRINT 401*VSOUND,AMACH,STAGFS                           000534
00420    179*          401 FCPMAT(7X,*FREESTREAM SPEED OF SOUND (VSCUND) = ',F10.3,' CM/SEC' 000543
00422    180*          1 /11X,*FREESTREAM MACH NUMBER (AMACH) = ',F6.3,/12X,*FREESTREAM S 000543
00422    181*          2TAC TEMP (STAGFS) = ',F9.3.' DEG KELVIN'                 000543
00423    182*          PRINT 402*RTFS,RTSTAC                                000543
00427    187*          402 FORMAT(14X,*FREESTREAM VISCOSITY (RTFS) = ',E10.5,' CM/CM-SEC' 000551
00427    188*          1 /7X,*FREESTREAM STAC VISCOSITY (RTSTAG) = ',E10.5,' GM/CM-SEC') 000551
00432    185*          PRINT 403                                         000551
00432    186*          403 FORMAT(4X,*SIMILARITY PARAMETERS:*)                   000555
00433    187*          PRINT 404,REYF,REYN,REWALL,XKNFS,VCAR,XKSG                000555
00443    188*          404 FORMAT(3X,*FREESTREAM REYNOLDS NUMBER (REYN) = ',F9.2, 000567
00443    189*          1 /EX,*STACNATION REYNOLDS NUMBER (REYN) = ',F9.2,        000567
00443    190*          2 /12X,*WALL REYNOLDS NUMBER (REWALL) = ',F9.2,        000567
00443    191*          3 /8X,*FREESTREAM KNUDSEN NUMBER (XKNFS) = ',F9.4,        000567
00443    192*          4 /37X,*VCAR = ',F9.4,                                000567
00443    193*          5 /25X,*K SQUARED (XKSG) = ',F9.4,//)                  000567
00443    194*          C NOTE THE SUBSTITUTION REYN=REYF FOR REMAINDER OF PROGRAM 000567
00443    195*          REYN=REYF                                         000567
00444    196*          PRINT 180                                         000571
00445    197*          000571
00445    198*          000571
00445    199*          C ****
00445    200*          000571
00447    201*          C
00450    202*          UNIR=1.087                                         000575
00451    203*          A=6.0222*10.*.*23                                         000577
00452    204*          WTA2=28.0                                         000601
00453    205*          WTC2=32.0                                         000603
00453    206*          WTN=14.0                                         000605
00454    206*          WTO=16.0                                         000607
00455    207*          WTN0=30.0                                         000611
00456    208*          WTN0I=30.0                                         000613
00457    209*          WTGL=1./1820.                                         000614
00460    210*          NDIV=50                                         000616
00461    211*          DIVN=NDIV                                         000620
00462    212*          DCL TA=1./DIVN                                000623
00463    213*          NDIV1=NCIV+1                                000626
00464    214*          HALT=25.0                                         000633
00464    215*          C ****
00464    216*          C ****                                         000633
00464    217*          C ****                                         000633
00465    218*          READ 300,(U(I),V(I),T(I),R0(I),P2(I),I=2,NDIV)           000635
00477    219*          READ 300,(C02(I),CNO1(I),CNO1(I),CN(I),C0(I),I=2,NDIV) 000653
00477    220*          C ****                                         000653
00477    221*          C ****                                         000653
00477    222*          C ****                                         000653
00477    223*          C BOUNDARY CONDITIONS AT FREESTREAM                         000653

```

MAIN

DATE 12C877

00511	224*	U(NDIV1)=1.	000670
00512	225*	V(NDIV1)=-1.	000672
00513	226*	T(NDIV1)=TFS/STAGFS	000673
00514	227*	RC(NDIV1)=1.	000676
00515	228*	RT(NDIV1)=1.0	000677
00516	229*	P2(NDIV1)=0.	000700
00517	230*	CN2(NDIV1)=0.767	000701
00520	231*	C02(NDIV1)=0.233	000703
00521	232*	CNC(NDIV1)=0.	000705
00522	233*	CO(NDIV1)=0.	000706
00523	234*	CN(NDIV1)=0.	000707
00524	235*	CNCI(NDIV1)=0.	000710
00525	236*	CEL(NDIV1)=0.	000711
00525	237*	C *****	000711
00525	238*	*****	000711
00525	239*	*****	000711
00525	240*	C CATALYTIC BOUNDARY CONDITIONS	000711
00525	241*	*****	000711
00525	242*	IF(XCAT-C.9 .LT. 0.2)GO TO 450	000711
00526	247*	CN2(1)=0.767	000712
00530	244*	C02(1)=0.233	000717
00531	245*	CNO(1)=0.	000720
00532	246*	CN(1)=0.0	000721
00533	247*	CO(1)=0.0	000722
00534	248*	CNCI(1)=CNOI(2)	000723
00535	249*	CEL(1)=CNOI(2)*WTEL/WTNOI	000724
00536	250*	PEL(1)=A*RO(1)*R0FS*CNOI(1)/WTNOI	000726
00537	251*	GO TO 451	000732
00540	252*	*****	000740
00540	253*	*****	000740
00540	254*	*****	000740
00540	255*	C NONCATALYTIC WALL BOUNDARY CONDITIONS (FOR FIRST ITERATION)	000740
00540	256*	*****	000740
00540	257*	*****	000740
00541	253*	450 CN2(2)=1.0-C02(2)-CN0(2)-C(2)-CNOI(2)*(1.0*WTEL/WTNOI)	000742
00542	259*	CN2(1)=CN2(2)	000754
00543	260*	C02(1)=C02(2)	000755
00544	261*	CNC(1)=CN0(2)	000757
00545	262*	CN(1)=CN(2)	000761
00546	263*	CO(1)=C(2)	000763
00547	264*	CNOI(1)=CNOI(2)	000765
00550	265*	CEL(1)=CNOI(2)*WTEL/WTNOI	000767
00551	266*	PEL(1)=0.	000773
00551	267*	C *****	000773
00551	268*	*****	000773
00551	269*	*****	000773
00551	270*	C INITIAL GUESS FOR CNOI, CEL AND H	000773
00552	271*	451 HIN2=0.0	000775
00553	272*	HIO2=0.	000775
00554	273*	HIN0=3050.*10.**7	000776
00555	274*	HIN=33620.*10.**7	001000
00556	275*	HIC=15430.*10.**7	001002
00557	276*	HINOI=9000.*10.**7	001004
00560	277*	HICL=0.	001006
00561	278*	W1=8.3143*10.**7*STAGFS/UFS**2	001007
00562	279*	DO 30 I=1,NDIV1	001200

MAIN

DATE 120877

00565	280*	CN2(I)=1.-C02(I)-CNO(I)-CN(I)-C0(I)-CNO(I)*(1.+WTNL/WTNOI)	001216
00566	281*	CEL(I)=CNO(I)*WTNL/WTNOI	001226
00567	292*	W31=T(I)*STAGFS	001232
00570	287*	VNC=3390./W31	001235
00571	294*	V02=2270./W31	001240
00572	285*	VNC=2740./W31	001243
00573	296*	VNOI=2740./W31	001246
00574	287*	IF(VN2 .GT. 80.C) GO TO 455	001247
00576	293*	VI0=(CN2(I)/28.)*VN2/(EXP(VN2)-1.)+(C02(I)/32.)*V02/(EXP(V02)-1.)	001252
00576	289*	1*(CNO(I)/30.)*VNO/(EXP(VNO)-1.)+(CNO(I)/30.)*VNOI/(EXP(VNOI)-1.)	001252
00577	290*	GO TO 456	001317
00600	291*	455 VIS=0.C	001321
00601	292*	456 CONTINUE	001322
00602	297*	H(I)=3.5*(CN2(I)/28.+C02(I)/32.+CNO(I)/30.+CNO(I)/30.)*VIS	001322
00602	294*	1+2.5*(CN(I)/14.+C0(I)/16.+CNO(I)/30.)	001322
00603	295*	H(I)=W1*T(I)*H(I)+(CNO(I)*HINO*CN(I)*HIN+C0(I)*HIO+CNO(I)*HINC1)	001347
00603	296*	1/U*5**2	001347
00603	297*	C	001347
00603	298*	C *****	001347
00603	299*	C *****	001347
00605	300*	DO 500 K=1,2000	001374
00610	301*	Q=CFPR*CELT	001377
00610	302*	C TO FIND P AMD MUE AT ALL POINTS	001377
00611	303*	DO 50 I=1,NCIV1	001406
00614	304*	TA5(I)=T(I)*STAGFS	001411
00615	305*	IF(I .EQ. 1) GO TO 31	001414
00617	306*	P(I)=R0(I)*T(I)*W1*(CN2(I)/WTN2+C02(I)/WT02+CNO(I)/WTN0+CN(I)/WTN	001414
00617	307*	1+C(I)/WT02+.*CNO(I)/WTNOI)	001414
00620	308*	31 XMUSU(I) = 1.453E-35*TABS(I)**1.5/(TABS(I) + 119.4)	001444
00621	309*	IF(XVIS-C.9 .LT. 0.2) GO TO 33	001457
00623	310*	IF(TABS(I) .GE. 2000.0) GO TO 32	001461
00625	311*	33 XMURAT(I)=1.0	001466
00626	312*	GO TO 37	001467
00627	313*	32 TA5=TABS(I)	001471
00630	314*	CALL INTERP(1,4,1,37,TEMP,TABS)	001472
00631	315*	CALL INTERP(2,4,1,37,TAB1,SN2N)	001502
00632	316*	CALL INTERP(2,4,1,37,TAB2,SNN)	001512
00633	317*	CALL INTERP(2,4,1,37,TAB3,SNE)	001522
00634	318*	CALL INTERP(2,4,1,37,TAB4,SEE)	001532
00635	319*	XLMN2=1.03329*CN2(I)+.87542*C02(I)+2.5310*CN(I)*SN2N	001542
00635	320*	I +2.12036*C01(I)*SN2N+.94819*CNO(I)+.94819*CNO(I)	001542
00636	321*	XLMC2=1.069E*CN2(I)+.90415*C02(I)*Z.6488*CN(I)*SN2N	001556
00636	322*	I +2.2146*C0(I)*SN2N+.9903*CN0(I)+.9803*CNO(I)	001566
00637	323*	XLMN=.8949*CN2(I)*SN2N+.7665*C02(I)*SN2N+2.0666*CN(I)*SNN	001612
00637	324*	I +1.7508*C0(I)*SNN+.8259*CN0(I)*SN2N+.8259*CNO(I)*SN2N	001612
00640	325*	XLMC=.9159*CN2(I)*SN2N+.7830*C02(I)*SN2N+2.1391*CN(I)*SNN	001642
00640	326*	I +1.8083*C0(I)*SNN+.8444*CN0(I)*SN2N+.8444*CNO(I)*SN2N	001642
00641	327*	XLMN0=1.0516*CN2(I)+.9859*C02(I)*Z.5906*CN(I)*SN2N	001672
00641	328*	I +2.1680*C0(I)*SN2N+.9644*CN0(I)+.9644*CNO(I)	001672
00642	329*	XLMN0I=1.0516*CN2(I)+.8759*C02(I)+3.2475*CN(I)*SN2N	001716
00642	330*	I +2.1680*C0(I)*SN2N+.9644*CN0(I)+.9644*CNO(I)	001716
00643	331*	XLMEL=-.7307*CN2(I)*SNN+.6393*C02(I)*SNN+1.4613*CN(I)*SNE	001731
00643	332*	I +2.2786*C0(I)*SNE+.6819*CN0(I)*SNN+.6819*CNO(I)*SEE	001731
00643	333*	2 +52656.24*CEL(I)*SEE	001731
00644	334*	XMRAT(I)=1.0165*CN2(I)/XLMN2+.9509*C02(I)/XLM02	001765
00644	335*	1 +1.4376*CN(I)/XLMN+1.8083*C0(I)/XLM0+.9820*CNO(I)/XLMN0	001765

MAIN

DATE 120877

```

00644 335*      3    +0.3820*CNOI(I)/XLMN01 + 229.469*CEL(I)/XLMEL          001765
00645 337*      37 XMUI(I) = XMURAT(I)*XMUSU(I)                                002022
00646 333*      50 RT(I) = XMU(I)/RTFS                                         002024
00646 339*      C   TO FIND PRESSURE ON THE SURFACE USING MOMENTUM EQUATION
00650 340*      U1C=(4.*U(2)-U(3)-3.*U(1))/DELTA/2.                            002030
00651 341*      V1C=(4.*V(2)-V(3)-3.*V(1))/DELTA/2.                            002D41
00652 342*      RT1C=(4.*RT(2)-RT(3)-3.*RT(1))/DELTA/2.                         002051
00653 343*      V2C=-V(4)+4.*V(3)-5.*V(2)+2.*V(1)                             002C62
00654 344*      DEL4V=V(5)-4.*V(4)+6.*V(3)-4.*V(2)+V(1)                         002074
00655 345*      V2C=V2C+11./12.*DEL4V                                         002105
00656 346*      V2C=V2C/DELTA**2                                         002110
00656 347*      C   SPHERE/CYLINDER OPTION FOLLOWS
00657 348*      IF(XSOD-0.9 .GT. 0.2) GO TO 51                               002112
00657 349*      C   SPHERE
00661 350*      P1C=V2C*RT(1)/CFFR+V10*(2.*RT(1)+RT1C/CFFR)+U10*RT(1)/2.-U(1)*( 002114
00661 351*      1RT1C+3.5*RT(1)*CFFR)                                         002114
00662 352*      GO TO 52                                              002137
00662 353*      C   CYLINDER
00663 354*      51 P1C=V2C*RT(1)/CFFR+V10*(RT(1)+RT1C/CFFR)+U10*RT(1)/4.-C-U(1)* 002141
00663 355*      1*(FT1C/2.C+7.C/4.C*RT(1)*EFFR)                           002141
00664 356*      52 P1C=P10*4.C/3.C/RCPN                                         002166
00665 357*      P(1)=P(2)-DELTA*P(1)                                         002172
00666 358*      R0(1)=P(1)/H1/T(1)/(CN2(1)/WTN2+C02(1)/WT02+CNOI(1)/WTN0+CN(1)/WTN 002175
00666 359*      +CC(1)/WT0+2.*CNOI(1)/WTNOI)                                002175
00667 360*      DO 60 N=2,NDIV                                         002252
00672 361*      EN=N                                              002267
00673 362*      53 X(N)=1.+(EN-1.)*Q                                     002272
00673 363*      C
00673 364*      C   SLIP CONDITIONS
00673 365*      C
00675 366*      AC=1.0                                         002301
00676 367*      W(1)=WTN2                                         002303
00677 368*      W(2)=WT02                                         002305
00700 369*      W(3)=WTN                                           002307
007C1 370*      W(4)=WT0                                           002311
007C2 371*      W(5)=WTNO                                         002313
00703 372*      W(6)=WTNOI                                         002315
00704 373*      W(7)=KTEL                                         002317
00705 374*      C1(1)=CN2(1)                                         002321
00706 375*      C1(2)=C02(1)                                         002323
00707 376*      C1(3)=CN(1)                                         002325
00710 377*      C1(4)=C0(1)                                         002327
00711 378*      C1(5)=CNO(1)                                         002331
00712 379*      C1(6)=CNOI(1)                                         002333
00713 380*      C1(7)=CEL(1)                                         002335
00714 381*      C2(1)=CN2(2)                                         002337
00715 382*      C2(2)=C02(2)                                         002341
00716 383*      C2(3)=CN(2)                                         002343
00717 394*      C2(4)=C0(2)                                         002345
00720 385*      C2(5)=CNO(2)                                         002347
00721 396*      C2(6)=CNOI(2)                                         002351
00722 387*      C2(7)=CEL(2)                                         002353
00723 389*      C3(1)=CN2(3)                                         002355
00724 389*      C3(2)=C02(3)                                         002357
00725 390*      C3(3)=CN(3)                                         002361
00726 391*      C3(4)=C0(3)                                         002363

```

EAT N

```

0C727   397*          C3(5)=CNO(3)
0C730   393*          C3(6)=CNOI(3)
0C731   394*          C3(7)=ECEL(3)
0C732   395*          TW=TWK/STAGFS
0C733   396*          SUM1=0.3
0C734   397*          DO 61 J=1,7
0C737   393*          61 SUM1=SUM1+C1(J)/W(J)
0C741   399*          DO 62 J=1,7
0C744   400*          C2 PP1(J)=(C1(J)/W(J))+P(1)/SUM1
0C746   401*          SUM2=C.C
0C747   402*          DC 63 J=1,7
0C752   403*          F3 SUM2=SUM2+(C2(J)-C1(J))/0
0C754   404*          SUM3=1.
0C755   405*          SUM4=C.C
0C756   406*          DO 65 J=1,7
0C761   407*          G5 SUM4=SUM4+(TW*RC(1)*C1(J)/(W(J)*SUM1)+*1.5)*(1.+RT(1)/B./PP1(J)/
0C761   408*          1*REYN*(U(1)-4.*V(2)/0+V(3)/0)+((3.1416*GAMMA*W(J)*SUM1*TF5/2./T(1)/
0C761   409*          2*STAGFS)*C.51/C1(J)*(2.-AC1)/AC*AMACH/REYN*EL(J)/PR*(W(J)*SUM1)*
0C761   410*          3RT(1)*RO(1)*SUM2)
0C763   411*          SUM5=C.C
0C764   412*          DC 66 J=1,7
0C767   413*          E6 SUM5=SUM5+C1(J)*RO(1)/(W(J)*SUM1)+*1.5)*(1.+RT(1)/2./PP1(J)/REYN
0C767   414*          1*(U(1)-4.*V(2)/0+V(3)/0))
0C771   415*          USLIP=RT(1)/P(1)/REYN/AMACH*((6.2832*STAGFS*T(1)/CAMMA/TF5)**C.51)*
0C771   416*          1*(2.-AC1)/AC/2.*((4.*U(2)-3.*U(1)-U(3))/2./Q
0C772   417*          TSLIP=(2.-AC1)/AC*(3.1416**C.5)
0C773   418*          TSLIP=(TSLIP*(AMACH*RT(1)*GAMMA/PR/REYN/2./((GAMMA-1.)*((GAMMA*TF5/
0C773   419*          12./T(1)/STAGFS)*C.5)*(T(2)-T(1))/Q-2.5*P(1)*RT(1)*
0C773   420*          *AM1CH**3.
0C773   421*          2*((CAMMA*TF5/STAGFS)*C.5)/RO(1)/REYN/((2.*T(1))*C.5)*EL1/PR*
0C773   422*          3SUM2)*SUM3
0C774   423*          TSLIP=(TSLIP+SUM4)/SUM5
0C775   424*          USCONT=(USLIP-U(1))/U(1)
0C776   425*          USCONT=USCONT
0C777   426*          USCONT=ABS(USCONT)
0C100   427*          IF(USCONT.GT.EPSI) U(1)=U(1)+USCONS/USCONT*EPSI*U(1)
0C1002  428*          IF(USCONT.LE.EPSI) U(1)=USLIP
0C1004  429*          TSCONT=(TSLIP-T(1))/T(1)
0C1005  430*          TSCONT=TSCONT
0C1006  431*          TSCONT=ABS(TSCONT)
0C1007  432*          IF( TSCONT.GT.EPSI) T(1)=T(1)+TSCONS/TSCONT*EPSI*T(1)
0C111   433*          IF( TSCONT.LE.EPSI) T(1)=TSLIP
0C1013  434*          RT(1)=(1.458C-05*(T(1)*STAGFS)*C.5)/((T(1)*STAGFS+110.4)*RTFS)
0C1013  435*          C CORRECTION TO V PROFILE
0C1014  436*          DO 70 N=2,NDIV
0C1017  437*          V2=V(N+1)-2.*V(N)+V(N-1)
0C1020  438*          P1=P(N+1)-P(N-1)
0C1021  439*          V1=V(N+1)-V(N-1)
0C1022  440*          RT1=RT(N+1)-RT(N-1)
0C1023  441*          U1=U(N+1)-U(N-1)
0C1023  442*          C SPHERC/CYLINDER OPTION FOLLOWS
0C1024  443*          IF(XBOD<0.9 .GT. 0.2) GO TO 67
0C1024  444*          C SPHERE
0C1026  445*          CAPW=-RT(N)*V2/Q**Z+D.375*REYN/Q*(P1+RO(N))*V(N)*V17-V17/Q
0C1026  446*          1*(RT(N)/X(N)+RT1/4./Q)-RT(N)*U1/4./Q/X(N)
0C1027  447*          CAPV=CAPW*(U(N)+V(N))/X(N)*T3.5*RT(N)/X(N)+RT1/2./Q)

```

DATE 1208

MAIN

DATE 120877

```

01030 448*      DERV=2.*RT(N)/Q**2+0.375*REYN/Q*RO(N)*V1+3.5*RT(N)/X(N)**2          003052
01C30 449*      1+RT1/2./Q/X(N)                                               003052
01031 450*      GO TO 68                                              003072
01C31 451*      CYLINDER                                              003072
01C32 452*      C 67 CAPV=RT(N)*V2/Q**2+0.375*REYN/Q*(P1+RO(N)*V(N)*V1)-V1/Q/2.0          003074
01C32 452*      1*(RT(N)/X(N)+RT1/2.0/Q)-RT(N)*U1/8.C/Q/X(N)                                003074
01033 454*      CAPV=CAPV+(U(N)+V(N))/X(N)*(7.0*RT(N)/X(N)+RT1/Q)/4.0          003126
01C34 455*      DERV=2.C*RT(N)/Q**2+C.375*REYN/Q*RO(N)*V1+(7.0*RT(N)/X(N)**2          003145
01034 456*      1+RT1/Q/X(N))/4.0                                              003145
01034 457*      C TO FIND CCCV                                              003145
01035 458*      68 SIGV=A35*EPSI*DCRV*V(N)/CAPV1                                     003174
01C36 459*      IF(SIGV.LT.1.)CCCV=SIGV                                         003174
01040 460*      IF(SIGV.GE.1.)CCCV=1.                                           003202
01C42 461*      70 V(N)=V(N)-CCCV*CAPV/DERV                                         003210
01044 462*      V(2)=(V(3)-4.*DELTA*U(1))*EFFR)/4.                                     003216
01C44 463*      C TO FIND CORRECTION TO U PROFILE                               003216
01045 464*      DO 80 N=2,NDIV                                              003224
01C50 465*      U2=U(N+1)-2.*U(N)+U(N-1)                                         003245
01051 466*      U1=U(N+1)-U(N-1)                                              003252
01052 467*      RT1=RT(N+1)-RT(N-1)                                         003255
01C53 468*      V1=V(N+1)-V(N-1)                                              003260
01C54 469*      CAFU=-RT(N)*U2/Q**2+REYN*RC(N)*V(N)*U1/2./Q          003263
01C54 470*      SPHCRE/CYLINDER OPTION FOLLOWS                               003263
01C55 471*      IF(XBCD-0.9 .GT. C.2) GO TO 71                                         003274
01055 472*      SPHCRE                                              003274
01C57 473*      CAFU=CAPU+(U(N)+V(N))/X(N)*(8.*RT(N)/3./X(N)+RT1/2./Q+REYN*RO(N) 003276
01C57 474*      1*U(N))                                              003276
01C60 475*      DERV=2.*RT(N)/Q**2+8.*RT(N)/3./X(N)**2+RT1/2./Q/X(N)+REYN*RO(N) 003314
01056 475*      1*(2.*U(N)+V(N))/X(N)                                         003314
01061 477*      GO TO 72                                              003335
01061 478*      C CYLINDER                                              003335
01C62 479*      71 CAFU=CAPU+(U(N)+V(N))/X(N)*(7.*RT(N)/3./X(N)+RT1/2./Q+PEYN*RO(N) 003337
01C62 480*      1*U(N))                                              003337
01C63 481*      DERV=2.*RT(N)/Q**2+7.C*RT(N)/3./X(N)**2+RT1/2./Q/X(N)+REYN*RO(N) 003362
01063 482*      1*(2.0*U(N)+V(N))/X(N)                                         003362
01C64 482*      73 CAFU=CAPU+RT(N)*V1/6.C/Q/X(N)-U1/6.[RT(N)/X(N)+RT1/74.C/70] 003404
01064 484*      1+2.0*P2(N)*REYN/X(N)                                         003404
01064 485*      C TO FIND CCCU                                              003404
01065 486*      SIGU=ABS((EPSI*DERU*U(N)/CAPU)                                         003431
01C66 487*      IF(SIGU.LT.1.)CCCU=SIGU                                         003437
01070 488*      IF(SIGU.GE.1.)CCCU=1.                                         003445
01C72 489*      80 U(N)=U(N)-CCCU*CAPU/DERU                                         003453
01072 490*      C SPECIES PRODUCTION RATE                               003453
01C74 491*      DO 90 N=1,NDIV1                                              003461
01077 492*      W5=T(N)*STAGFS                                         003474
01100 493*      W6=UNIR*W5                                              003476
01101 494*      W7=118000./W6                                              003500
01102 495*      W8=W7**1.5.                                         003503
01103 495*      W9=224900./W6                                              003510
01104 497*      W10=150000./W6                                              003513
01105 499*      W11=W1C**2                                              003516
01106 499*      W12=39100./W6                                              003520
01107 500*      W13=7080./W6                                              003523
01110 501*      W14=75500./W6                                              003526
01111 502*      W15=128500./W6                                              003531
01112 503*      W16=85520./W6                                              003534

```

MAIN

			DATE 12C877
C1113	504*	W17=63290./W6	003537
C1114	505*	W18=W5**0.5	003542
C1115	506*	W19=W5**2.5	003547
C1116	507*	W20=500./W6	003554
C1117	508*	W210=-10./W6	003557
C1120	509*	IF(W7.CT.BC.)G0 TO 81	003562
C1122	510*	FK1=2.5*10.**11*W18*W9/EXP(W7)	003566
C1123	511*	G0 TO 82	003576
C1124	512*	31 FK1=C.	003600
C1125	513*	52 RK1=2.08*10.**8*W5*W8	003601
C1126	514*	IF(W9.GT.BC.)G0 TO 83	003604
C1130	515*	FK2=1.7*10.**12*W18*W9/EXP(W8)	003610
C1131	516*	G0 TO 34	003620
C1132	517*	23 FK2=C.	003622
C1133	518*	34 RK2=9.444*10.**10*W18*W9	003623
C1134	519*	IF(W10.CT.BC.)G0 TO 85	003626
C1136	520*	FK3=7.*10.**10*W18*W11/EXP(W10)	003632
C1137	521*	G0 TO 86	003642
C1140	522*	35 FK3=C.	003644
C1141	523*	85 RK3=1.75*10.**10*W18*W11	003645
C1142	524*	IF(W12.CT.BC.)G0 TO 87	003650
C1144	525*	FK4=3.2*10.**9*W5/EXP(W12)	003654
C1145	526*	G0 TO 89	003663
C1146	527*	37 FK4=C.	003665
C1147	528*	IF(W13.GT.BC) G0 TO 881	003665
C1151	529*	RK4=13.33*10.**9*W5/EXP(W13)	003672
C1152	530*	G0 TO 932	003700
C1153	531*	881 RK4=C.	003702
C1154	532*	382 IF(W14.CT.BC.)G0 TO 91	003703
C1156	533*	FK4=7.*10.**13/EXP(W14)	003706
C1157	534*	G0 TO 92	003714
C1160	535*	91 FK4=C.	003716
C1161	536*	IF(W20.CT.BC.) G0 TO 921	003716
C1163	537*	RK4=1.55*10.**13/EXP(W20)	003723
C1164	538*	G0 TO 922	003730
C1165	539*	921 RK4=C.	003732
C1166	540*	392 IF(W15.GT.BC.)G0 TO 93	003733
C1170	541*	FK4=9.1*10.**24/W19/EXP(W15)	003736
C1171	542*	G0 TO 34	003745
C1172	543*	93 FK4=C.	003747
C1173	544*	IF(W16.GT.BC.)G0 TO 95	003750
C1175	545*	RK4=9.79*10.**23/W19/EXP(W16)	003753
C1176	546*	G0 TO 96	003762
C1177	547*	94 RK4=C.	003764
C1200	548*	36 IF(W17.CT.BC.)G0 TO 97	003765
C1202	549*	FK4=6.4*10.**9*W18/EXP(W17)	003770
C1203	550*	G0 TO 98	003777
C1204	551*	97 FK4=C.	004001
C1205	552*	98 RK4=1.79*10.**19/W5/EXP(W210)	004002
C1206	553*	W2=R0(N)*R0FS	004010
C1207	554*	W3=W2**2	004013
C1210	555*	R1=W3*(CN(N)/WTN+CNO(N)/WTN0+2.*CN2(N)/WTN2+9.*C02(N)/WT02+	004015
C1210	556*	125.*C01(N)/WT0)	004015
C1211	557*	R1=R1*(FK1*C02(N)/WT02-W2*RK1*C01(N)**2/WT0**2)	004041
C1212	558*	R2=W3*(C0(N)/WT0+C02(N)/WT02+CNO(N)/WTN0+2.47*CN2(N)/WTN2+2.15*	004055
C1212	559*	110.**5*CN(N)/W5/WTN)	004055

MAIN

DATE 120877

O1213	567*	R2=R2*(FK2*CN2(N)/WTN2-W2*RK2*CN(N)**2/WTN**2)	004077
O1214	561*	R3=W3*(C02(N)/WT02+CN2(N)/WTN2+2D.*(CNO(N)/WTNO+C0(N))/WT0+1CN(N)/WTN))	004113
O1215	562*	R3=R3*(FK3*CN0(N)/WTN0-W2*RK3*CN(N)*C0(N)/WTN/WT0)	004113
O1216	564*	R4=W3*(FK4*CN0(N)*C0(N)/WTN0/WT0-RK4*C02(N)*CN(N)/WT02/WTN)	004127
O1217	565*	R5=W3*(FK5*CN2(N)*C0(N)/WTN2/WT0-RK5*CN0(N)*CN(N)/WTNO/WTN)	004143
O1220	566*	R6=W3*(FK6*CN2(N)*C02(N)/WT02/WTN2-RK6*(CNO(N)/WTN0)**2)	004160
O1221	567*	R7=W3*(FK7*CN(N)*C0(N)/WTN/WT0-RK7*(CNO(N)/WTNOI)**2)	004175
O1222	561*	W4=ROFS*UFS/RADS	004211
O1223	568*	W02(N)=WT02*(-R1+R4-R6)/W4	004225
O1224	570*	WN2(N)=-WTN2*(R2+R5+R6)/W4	004227
O1225	571*	WN0(N)=WTN0*(-R3-R4+R5+R6)/W4	004234
O1226	572*	WN(N)=WTN*(2.*R2+R3+R4+R5-R7)/W4	004241
O1227	573*	WN1(N)=WT0*(2.*R1+R2-R4-R5-R7)/W4	004251
O1230	574*	WN01(N)=WTNOI*R7/W4	004262
O1231	575*	WEL(N)=WTEL*R7/W4	004273
O1232	575*	CONTINUE	004277
O1233	577*	CONTINUATION TO CONCENTRATION PROFILES	004310
O1234	573*	DO 100 N=2,NDIV	004310
O1237	579*	RT1=RT(N+1)-RT(N-1)	004315
O1240	580*	W21=-R0(N)+V(N)/2./Q	004320
O1241	581*	W22=(2.*RT(N)/X(N)+RT1/2./Q)/2./Q/REYN/SC1	004325
O1242	582*	W27=RT(N)/Q**2/REYN/SC1	004337
O1243	587*	DEI:C=-2.*RT(N)/REYN/SC1/Q**2	004344
O1243	584*	OXYGEN MOLECULE	004344
O1244	585*	C021=C02(N+1)-C02(N-1)	004351
O1245	586*	C023=C02(N+1)-2.*C02(N)+C02(N-1)	004354
O1246	587*	CAP02=W02(N)+(W21+W22)*C021+W23*C023	004361
O1247	583*	SIG02=ABS(EPSI*DERC*C02(N)/CAP02)	004367
O1250	589*	IF(SIG02.LT.1.)CCC02=SIG02	004377
O1252	593*	IF(SIG02.GE.1.)CCC02=1.	004405
O1254	591*	C02(N)=C02(N)-CCC02*CAP02/DERC	004413
O1254	592*	NO ION	004413
O1255	593*	W221=(2.*RT(N)/X(N)+RT1/2./Q)/2./Q/REYN/SC2	004420
O1256	594*	W231=RT(N)/Q**2/REYN/SC2	004430
O1257	595*	DERC1=-2.*RT(N)/REYN/SC2/Q**2	004435
O1260	596*	CNOII=CNOI(N+1)-CNOI(N-1)	004442
O1261	597*	CNGI3=CNOI(N+1)-2.*CNOI(N)+CNOI(N-1)	004445
O1262	598*	CAPNOI=WN0I(N)+(W21+W221)*CNOI1+W231*CNOI3	004452
O1263	599*	SIGNOI=ABS(EPSI1*DERC1*CNOI(N)/CAPNOI)	004460
O1264	600*	IF(SIGNOI.LT.1.)CCCN0I=SIGNOI	004465
O1266	601*	IF(SIGNOI.GE.1.)CCCN0I=SIGNOI	004473
O1270	602*	CNOI(N)=CNOI(N)-CCCN0I*CAPNOI/DERC1	004501
O1270	603*	NITROGEN ATOM	004501
O1271	604*	CN1=CN(N+1)-CN(N-1)	004506
O1272	605*	CN3=CN(N+1)-2.*CN(N)+CN(N-1)	004511
O1273	606*	CAPN=WN(N)+(W21+W22)*CN1+W23*CN3	004516
O1274	607*	SIGN1=ABS(EPSI*DERC*CN(N)/CAPN)	004523
O1275	608*	IF(SIGN1.LT.1.)CCCN=SIGN1	004530
O1277	609*	IF(SIGN1.GE.1.)CCCN=1.	004536
O1301	610*	CN(N)=CN(N)-CCCN*CAPN/DERC	004544
O1301	611*	OXYGEN ATOM	004544
O1302	612*	C01=C0(N+1)-C0(N-1)	004551
O1303	613*	C02=C0(N+1)-2.*C0(N)+C0(N-1)	004554
O1304	614*	CAP0=W0(N)+(W21+W22)*C01+W23*C03	004561
O1305	615*	SIG0=ABS(EPSI*DERC*C0(N))/CAP0	004566

1. Report No. NASA TP-1227	2. Government Accession No.	3. Recipient's Catalog No.	
4. Title and Subtitle A Numerical Solution of the Navier-Stokes Equations for Chemically Nonequilibrium, Merged Stagnation Shock Layers on Spheres and Two-Dimensional Cylinders in Air		5. Report Date May 1978	
7. Author(s) Kenneth D. Johnston and William L. Hendricks*		6. Performing Organization Code	
9. Performing Organization Name and Address George C. Marshall Space Flight Center Marshall Space Flight Center, Alabama 35812		8. Performing Organization Report No. M-254	
12. Sponsoring Agency Name and Address National Aeronautics and Space Administration Washington, D.C. 20546		10. Work Unit No.	
11. Contract or Grant No.			
13. Type of Report and Period Covered Technical Paper			
14. Sponsoring Agency Code			
15. Supplementary Notes Prepared by Systems Dynamics Laboratory, Science and Engineering. *Lockheed Huntsville Research and Engineering Center			
16. Abstract Results of solving the Navier-Stokes equations for chemically nonequilibrium, merged stagnation shock layers on spheres and two-dimensional cylinders are presented. The effects of wall catalysis and slip are also examined. The thin shock layer assumption is not made, and the thick viscous shock is allowed to develop within the computational domain. The results show good comparison with existing data. Due to the more pronounced merging of shock layer and boundary layer for the sphere, the heating rates for spheres become higher than those for cylinders as the altitude is increased.			
17. Key Words (Suggested by Author(s))		18. Distribution Statement	
		STAR Category 34	
19. Security Classif. (of this report) Unclassified	20. Security Classif. (of this page) Unclassified	21. No. of Pages 102	22. Price \$6.25

* For sale by the National Technical Information Service, Springfield, Virginia 22161

National Aeronautics and
Space Administration

Washington, D.C.
20546

Official Business

Penalty for Private Use, \$300

THIRD-CLASS BULK RATE

Postage and Fees Paid
National Aeronautics and
Space Administration
NASA-451



3 1 1U.D. 042778 S00903DS

DEPT OF THE AIR FORCE
AF WEAPONS LABORATORY
ATTN: TECHNICAL LIBRARY (SUL)
KIRTLAND AFB NM 87117

S

POSTMASTER: If Undeliverable (Section 158
Postal Manual) Do Not Return

NASA