MONTE CARLO CALCULATIONS OF DIATOMIC MOLECULE GAS FLOWS INCLUDING ROTATIONAL MODE EXCITATION

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A sample of 1000 model molecules was observed as the gas relaxed to a steady state starting from three specified initial states. The initial states considered are: (1) complete equilibrium, (2) nonequilibrium equipartition (i.e., all rotational energy states are assigned the mean energy level that obtains at equilibrium with a Boltzmann distribution at the translational temperature), and (3) nonequipartition (i.e., the mean rotational energy is different from the equilibrium mean value with respect to the translational energy states). Since only uniform flow is considered, the effect of elastic collisions is ignored in the Monte Carlo simulation.

In all cases investigated the present model satisfactorily simulated the principal features of the relaxation effects in nonequilibrium flow of diatomic molecules.
### NOMENCLATURE

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
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<tbody>
<tr>
<td>A</td>
<td>constants appearing in equations (23) and (25)</td>
</tr>
<tr>
<td>(A^{(6)})</td>
<td>integral appearing in cross section (see eq. (6))</td>
</tr>
<tr>
<td>B</td>
<td>constant associated with intermolecular potential (eq. (23))</td>
</tr>
<tr>
<td>b</td>
<td>impact parameter</td>
</tr>
<tr>
<td>C</td>
<td>constant associated with intermolecular potential (eq. (23))</td>
</tr>
<tr>
<td>(C_\alpha)</td>
<td>coefficient of wave function (eq. (13))</td>
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<tr>
<td>(C_6)</td>
<td>coefficient associated with (R^6) term (eq. (23))</td>
</tr>
<tr>
<td>c</td>
<td>molecular speed</td>
</tr>
<tr>
<td>(c_0)</td>
<td>rms molecule speed, (\sqrt{\frac{3kT}{M}})</td>
</tr>
<tr>
<td>D_\alpha</td>
<td>transition amplitude</td>
</tr>
<tr>
<td>d</td>
<td>strength associated with intermolecular potential</td>
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<tr>
<td>E</td>
<td>kinetic energy or energy</td>
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<tr>
<td>f</td>
<td>distribution function</td>
</tr>
<tr>
<td>g</td>
<td>relative speed</td>
</tr>
<tr>
<td>(\hbar)</td>
<td>Planck's constant divided by (2\pi)</td>
</tr>
<tr>
<td>I</td>
<td>quantity given in equation (25)</td>
</tr>
<tr>
<td>K</td>
<td>matrix whose element is defined by equation (20)</td>
</tr>
<tr>
<td>k</td>
<td>Boltzmann constant</td>
</tr>
<tr>
<td>m</td>
<td>magnetic quantum number; also molecular weight</td>
</tr>
<tr>
<td>N</td>
<td>number of simulated molecules within the cell</td>
</tr>
<tr>
<td>n</td>
<td>number density</td>
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<tr>
<td>P</td>
<td>transition probability</td>
</tr>
<tr>
<td>(\tilde{P})</td>
<td>modified transition probability</td>
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<td>Q</td>
<td>collision cross section</td>
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</table>
R intermolecular distance
T temperature
t time
V interaction potential
W eigenenergy of the rotational Hamiltonian
|α⟩ eigenfunction of the rotational Hamiltonian
Δτ increment of collision time
δ index of power associated with point centers of repulsion model
   (eq. (5)) or delta function (eq. (16))
λ quantity defined in equation (25)
μ reduced mass
ν collision frequency
σ effective collision diameter
τ characteristic collision time based on initial translational
   temperature, \(1/(n\pi\sigma^2C_0)\)
χ deflection angle defined in equation (1) or molecular orientation
   angle (eq. (22))
ψ wave function
ω difference eigenenergy

Subscripts:
C distance of closest approach of pair of molecules
i,j rotational states
max maximum value
0 initial
r rotational
t total
tr translational
α rotational state
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SUMMARY

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In all cases investigated the present model satisfactorily simulated the principal features of the relaxation effects in nonequilibrium flow of diatomic molecules.

INTRODUCTION

Understanding the energy balance, as well as the energy transfer mechanisms, within the internal excited states of a nonequilibrium rarefied flowing gas, is important, in indeed, to a number of problem areas within the broad categories of planetary reentry, combustion, and pollution. In this paper, results are presented on translation-rotation relaxation. Furthermore, these results are obtained by solving Boltzmann's equation by the Monte Carlo direct simulation technique, a method that has received considerable recent attention (refs. 1-11). A feature of this method is that it gives insight into the effects of relaxation on the microscopic level during molecular

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collisions; in particular, the instantaneous distributions of internal states can be continuously followed.

The method is described in detail elsewhere (refs. 1 and 6-9). Briefly, the flow is computed by following in detail several thousand model molecules that are allowed to interact with each other. The coordinates of each molecule in phase space (including rotational state) are at all times known. These coordinates change only during a collision and the modeling of these intermolecular encounters is the essence of an accurate simulation. To account for these encounters, a molecule and a near neighbor are each selected at random as are also their impact parameter and deflection angles - all in a manner representative of typical molecules undergoing encounters. They are accepted for an interaction or rejected depending on a selected rule that depends on cross section and, therefore, on intermolecular potential and the relative velocity of the collision pair. Since the initial coordinates (relative velocity, impact parameter, and pair of rotational states) are known, there remains only to find the final rotational state. This is found by computing the distribution (transition probability) of all final states accessible from the known initial states. The final state is then determined by a random selection from this distribution.

The procedure for handling the "translational" interactions parallels other investigations (refs. 1 and 6-9) treating monatomic gases. The procedure described in this paper differs, however, from other investigations of the treatment of internal state interactions. These other investigations fall into four categories: (1) semi-empirical, (2) classical, (3) semi-classical, and (4) quantum mechanical. The semi-empirical models, energy sink (ref. 11), and rough spheres and loaded spheres (ref. 12), while adequate for steady flows at or near equilibrium, lack sufficient physical detail to inspire confidence in their use for highly nonequilibrium flows.

The classical models (refs. 13 and 14), although consistent with the classical direct simulation Monte Carlo procedure used here, necessarily include approximations to make the models sufficiently tractable to a study of the type that is the subject of this report. Those approximations, although yielding appropriate macroscopic behavior for a nonequilibrium gas, do not adequately describe its microscopic behavior. For example, molecular encounters can occur that violate energy and momentum conservation. One, therefore, is at a loss as to how to treat the negative energies and momentum that arise during a simulation.

Semiclassical models (refs. 10, 15, and 16) appear to be based on physically realistic criteria; however, the Pearson and Hansen model was too simplified. Although this model would not seriously violate equilibrium concepts, it was subject to slow drifts from equipartition (ref. 10) (i.e., the rotational temperature would drift from equality with the translational temperature). Although, by heuristic arguments this model could be altered to qualitatively satisfy proper interaction behavior, we have based our investigation described here on the semiclassical model of Itikawa (ref. 16). This model is founded on more rigorous concepts and, in addition, allows for treatment of molecular collisions. The model also satisfies conservation of probability.
In this paper we treat translation-rotation interactions for a uniformly flowing gas far removed from solid boundaries. In fact, we assume that the rotational relaxation does not affect the flow; we are concerned only with understanding the rotational relaxation behavior. Results are given based on calculations starting from three different sets of initial conditions: (1) complete equilibrium, (2) nonequilibrium equipartition (distribution of energy is constant for all molecules - the energy value is based on the total energy within the rotational states at complete equilibrium that is to be uniformly distributed to each molecule; that is, total energy in rotation is physically correct but distributed incorrectly), and (3) nonequipartition (same as (2) except partition of energy within rotational states is not based on complete equilibrium). Throughout this paper the key mathematical relations essential to the Monte Carlo simulation are defined. Readers desiring a more comprehensive treatment of the method are referred elsewhere (refs. 1 and 6-9).

PROCEDURE

The essential aspects of the procedure are described in the introduction. The analytical relations peculiar to this investigation are described in the text that follows.

Selection Rule Defining the Occurrence of an Encounter

The key to an accurate simulation is the procedure for selecting the molecular pair to reach in a collision, determining whether a reaction occurs, advancing the time parameter in a systematic manner until the next collision occurs, and so on. Both the probability and time intervals are strongly dependent on collision frequency which, in turn, is dependent on the uncertain relation for intermolecular potential.

To aid this discussion, it is worthwhile to refer briefly to classical relations and how those relations depend on, for example, an intermolecular potential base and on a two-parameter repulsion between the point-center model.

In this case, where the intermolecular potential $V(R)$ is spherically symmetric, the deflection angle of an encounter, $\chi(b,g)$, which depend on impact parameter $b$ and relative velocity of approach, $g$, and the $k$th moment "transport" cross section, $Q^{(k)}$, are given by

$$\chi(b,g) = \pi - 2 \int_{R_C}^{\infty} (bdR/R^2) / \sqrt{1 - (b/R)^2 - V(R)/(1/2)\mu g^2} \tag{1}$$

$$Q^{(k)}(g) = 2\pi \int_0^{\infty} (1 - \cos^2 \chi)bdb \tag{2}$$
where $R_C$ and $\mu$ are the distance of closest approach and reduced mass, respectively. (For example, see chap. 8, ref. 12.) The collision frequency $\nu$ is then given by

$$\nu = nQ(\ell)g$$

(3)

From this relation, we can compute the collision time $\Delta t$ of an encounter and the elapsing time $t$. There results

$$\Delta t = \frac{2}{N} \frac{1}{\nu}$$

$$t = \sum \Delta t$$

(4)

where $N$ is the number of particles in a simulated cell. The collision frequency is not, however, accurately known in general.

In the case when the potential can be described by

$$V(R) = \frac{d}{R^\delta}$$

(5)

the frequency is given by

$$\nu = 2\pi n \left( \frac{\delta d}{\frac{1}{2} \mu} \right)^{2/\delta} A(\ell) (\delta) g^{(\delta-4)/\delta}$$

(6)

If $\nu_{\text{max}}$ and $g_{\text{max}}$ are the maximum values possible in a cell then the dimensionless ratios in the equation

$$\left( \frac{\nu}{\nu_{\text{max}}} \right) = \left( \frac{g}{g_{\text{max}}} \right)^{(\delta-4)/\delta}$$

(7a)

define a curve for specified values of $\delta$. If one accepts the above relation as representative of molecular encounters, then any point in the region below the curve represents a valid encounter, and points above the curve are invalid. We can then use the relation as a "selection rule," defining whether an encounter occurs or not.

Rather than accept the above relations as completely valid, we also investigated results using the linear relation

$$\frac{\nu}{\nu_{\text{max}}} = B + (1 - B) \frac{g}{g_{\text{max}}}$$

(7b)
where \( B \) is an adjustable parameter that gives the best results in the case \( B \approx 0.3 \). The reasons underlying this choice are described later.

Discussion of transition probabilities is given in the next section. Given the fact that a rotational transition has occurred, however, trajectories are required by the method so that the particle coordinates in phase-space can be recomputed.

Collision Dynamics

The relative velocity after the collision is obtained by knowing the rotational energy and momentum before and after a collision. These relations are classical relations given by

\[
(g')^2 = g^2 - \frac{2}{\mu} (E_{r1}' - E_{r1} + E_{r2}' - E_{r2})
\]

and

\[
b' = \frac{[gb - (M_{r1}' - M_{r1} + M_{r2}' - M_{r2})/\mu]}{g'}
\]

where \( E_{r2} \) and \( M_{r2} \) are rotational energy and momentum before a collision and a prime denotes value after a collision, the transition probabilities as well as calculation of a trajectory are based on \(|g' - g|/g \ll 1\) and \(|b' - b|/b \ll 1\); that is, the relative velocities and impact parameters are only slightly perturbed as a result of the rotational transitions.

The deflection angle (eq. (1)), although also dependent on intermolecular potential, can be adequately approximated to obtain trajectory results by the infinite-rise, rigid-spherical molecule of diameter \( \sigma \). Such a relation yields

\[
\chi(b) = 2 \cos^{-1} \left( \frac{b}{\sigma} \right)
\]

To properly account for the effect of inelastic collisions, we use the following mean value:

\[
\bar{\chi}(b) = \frac{[\chi(b) + \chi(b')]}{2}
\]

The derivation of the relation for rotational transition probabilities is briefly reviewed in the next section.

ROTATIONAL TRANSITION PROBABILITY

The details of the method and its applicability are discussed in reference 16. Only major derivations and results will be presented here for the purpose of direct application to the Monte Carlo simulation.
The interaction considered in the calculation of rotational transition probability is described by the reaction:

\[ N_2(j_1) + N_2(j_2) \rightarrow N_2(j'_1) + N_2(j'_2) \]  

(12)

The calculation is based on a semi-classical theory. To make numerical calculations tractable, several approximations are made. Whether these approximations are valid is difficult to assess, except that they lead to the correct qualitative behavior in the several applications considered. The total wave function of the system is expanded in terms of a set of wave functions based on a "rotational" Hamiltonian and given by:

\[ \psi = \sum_\alpha C_\alpha(t) |\alpha\rangle \exp \left( -\frac{i}{\hbar} W_\alpha t \right) \]  

(13)

Here \( \alpha \) specifies the rotational state of the molecules and \( W_\alpha \) and \( |\alpha\rangle \) are, respectively, the eigenenergy and eigenfunction of that state. The time-dependent coefficient \( C_\alpha(t) \) is then determined by:

\[ i\hbar \frac{dC_\alpha}{dt} = \sum_\alpha \langle \alpha' | V | \alpha \rangle \exp(i\omega_{\alpha'\alpha} t) C_\alpha \]  

(14)

where \( V \) is the interaction potential and \( \omega_{\alpha'\alpha} = \frac{(W_{\alpha'} - W_\alpha)}{\hbar} \). We separate the Hamiltonian into isotropic (spherically symmetric) and nonisotropic parts. Since our interest is in inelastic collisions, the elastic process due to the isotropic part of the potential, \( V(0) \), is eliminated by introducing the distorted-wave type coefficient defined by:

\[ D_\alpha(t) = C_\alpha(t) \exp \left\{ \frac{i}{\hbar} \int_{-\infty}^t V(0)[R(t')] dt' \right\} \]  

(15)

where \( R(t) \) is the distance separating gravity centers of the molecules at time \( t \). The coefficient \( D_\alpha \) satisfies the equation

\[ i\hbar \frac{dD_\alpha}{dt} = \sum_\alpha \left[ \langle \alpha' | V | \alpha \rangle - V(0)\delta_{\alpha'\alpha} \right] \exp(i\omega_{\alpha'\alpha} t) D_\alpha \]

\[ = \sum_\alpha \langle \alpha' | \tilde{V} | \alpha \rangle \exp(i\omega_{\alpha'\alpha} t) D_\alpha \]  

(16)

and the initial condition

\[ D_\alpha(t = -\infty) = \begin{cases} 1 & \text{for } \alpha = \alpha_0 \\ 0 & \text{for } \alpha \neq \alpha_0 \end{cases} \]  

(17)

The second line in equation (16) defines the reduced matrix element \( \langle \alpha' | \tilde{V} | \alpha \rangle \).
The transition probability for the process \( \alpha_0 \rightarrow \alpha \) is given by

\[
P(\alpha_0 \rightarrow \alpha) = |D_\alpha(t = \infty)|^2
\]  

(18)

In a rigorous treatment, the state \( \alpha \) depends on the rotational angular momenta, \( j \), and the projections, \( m \), of both molecules. For the present problem, we are only interested in the probability averaged over the \( m \) states. We use Rabitz method (ref. 17) to eliminate the \( m \)-dependence of the interaction matrix (effective potential method). We solve equation (16) with:

\[
\alpha \equiv (j_1, j_2), \quad D_\alpha \equiv D_{j_1j_2} \quad \text{and} \quad \langle \alpha' | \tilde{V} | \alpha \rangle \equiv \langle j_1', j_2' | \tilde{V}^{\text{eff}} | j_1j_2 \rangle
\]

This treatment is discussed in more detail in appendix A.

We can further approximate the solution of equation (16) by introducing the exponential approximation (refs. 18-20).

\[
D_{\alpha'}(\infty) = \langle \alpha' | \exp \tilde{K} | \alpha_0 \rangle
\]

\[
= \sum_{n=0}^{\infty} \frac{1}{n!} \langle \alpha' | \tilde{K}^n | \alpha_0 \rangle
\]  

(19)

where \( \tilde{K} \) is a matrix whose element is defined by

\[
\langle \alpha_i | \tilde{K} | \alpha_j \rangle = -\frac{i}{\hbar} \int_{-\infty}^{\infty} dt \langle \alpha_i | \tilde{V}^{\text{eff}} | \alpha_j \rangle \exp(i\omega_{ij}t)
\]  

(20)

The element of \( \tilde{K}^n \) is evaluated by

\[
\langle \alpha' | \tilde{K}^n | \alpha_0 \rangle = \sum_{\alpha_1\alpha_2, \ldots, \alpha_{n-1}} \langle \alpha' | \tilde{K} | \alpha_{n-1} \rangle \langle \alpha_{n-1} | \tilde{K} | \alpha_{n-2} \rangle \cdots \langle \alpha_1 | \tilde{K} | \alpha_0 \rangle
\]  

(21)

For \( N_2 + N_2 \), we select the following interaction potential as the relevant interaction (refs. 21-22),

\[
V = V^{(0)}(R) + V^{(1)}(R)[P_2(\cos x_1) + P_2(\cos x_2)]
\]

\[
+ V^{(2)}(R)P_2(\cos x_1)P_2(\cos x_2)
\]  

(22)

Here \( P_2 \) is the Legendre function of order two, \( R \) is the separation distance between the two molecules as before, and \( x_i \) is the angle between the
directions of the inter-molecular vector and the axis of $i$th molecule; the two molecules are not necessarily in a single plane. $V^{(2)}$ induces a simultaneous rotational transition in both molecules in first order. Each term of the potential is assumed to have the form

$$
\begin{align*}
V^{(0)}(R) &= C \exp(-\alpha R) - \frac{C_6}{R^6} \\
V^{(1)}(R) &= AC \exp(-\alpha R) \\
V^{(2)}(R) &= BC \exp(-\alpha R)
\end{align*}
$$

(23)

(See appendix A for further discussion.)

In the present calculation the trajectory $R(t)$ is determined by solving the classical equation of motion approximated by

$$
\frac{1}{2} \mu \left( \frac{dR}{dt} \right)^2 = E - C e^{-\alpha R} + \frac{C_6}{R_C^6} - \frac{E b^2}{R_C^2}
$$

(24)

where $\mu$ is the reduced mass of the system, $b$ is the impact parameter, $R_C$ is the distance of closest approach, and $E$ is the kinetic energy of the relative motion. This provides a semiclassical version of the modified wave number approximation (ref. 23) with the assumption that the gradients of the attractive part of the potential and the centrifugal force are much smaller than those of the repulsive potential, in the region where most of the rotational transition takes place. The solution of equation (24) can be obtained analytically and the time integration in equation (20) can be performed readily. We have, then

$$
\langle \alpha_i | \tilde{K} | \alpha_j \rangle = -i \langle \alpha_i | \alpha_j \rangle \tilde{I}
$$

(25)

with

$$
\tilde{I} = \frac{\pi}{\alpha \lambda \Lambda} \coth \left( \frac{\pi}{2\Lambda} \right)
$$

$$
\lambda = K (2\mu \tilde{E})^{-1/2}
$$

$$
\Lambda = \frac{\alpha}{2 |\omega_{\alpha_i \alpha_j}|} \left( \frac{2\tilde{E}}{\mu} \right)^{1/2}
$$

and

$$
\tilde{E} = E \left( 1 - \frac{b^2}{R_C^2} \right) + \frac{C_6}{R_C^6}
$$
Two important properties of the transition probability should be mentioned: the conservation of probabilities and detailed balancing. The exponential approximation self-ensures the conservation of probabilities:

\[
\sum_{j_1',j_2'} P(j_1j_2 + j_1'j_2') = 1
\]  

(26)

To satisfy the detailed balancing relation, we choose, as the kinetic energy \( E \) in the calculation of \( \langle \alpha_i | V_{\text{eff}} | \alpha_i' \rangle \), the mean value of the initial and final channel energies (i.e., \( E = (1/2)(E_i + E_j) \)). The channel energy \( E_i \) for the \( i \)th channel is defined by

\[
E_i = E_t - W_{\alpha_i} = (E_0 + W_{\alpha_0}) - W_{\alpha_i}
\]  

(27)

where \( E_0 \) is the initial relative kinetic energy and \( E_t \) is the total energy (which is conserved during the collision). This procedure results in a symmetry relation

\[
P(j_1j_2 + j_1'j_2';E_0) = P(j_1'j_2' + j_1j_2;E_0')
\]  

(28)

where

\[
E_0 + W_{j_1j_2} = E_0' + W_{j_1'j_2'},
\]

In order to have a properly detailed balance, we modify our result by

\[
\tilde{P}(j_1j_2 \rightarrow j_1'j_2') = \frac{(2j_1< + 1)(2j_2< + 1)}{(2j_1 + 1)(2j_2 + 1)} P(j_1j_2 \rightarrow j_1'j_2')
\]  

for \((j_1j_2) \neq (j_1'j_2')\), and

\[
\tilde{P}(j_1j_2 \rightarrow j_1j_2) = 1 - \sum_{j_1'j_2'(#j_1j_2)} \tilde{P}(j_1j_2 \rightarrow j_1'j_2')
\]  

(29)

Here \( j_1< \) designates the smaller of \( j_1 \) and \( j_1' \). These modified transition probabilities, \( \tilde{P} \), satisfy both the detailed balancing relation and the conservation of probabilities.

The program listings for calculating rotational transition probability are presented in appendix B.
RESULTS AND DISCUSSION

Calculations were performed using three different sets of initial conditions. These initial conditions, described in the introduction, are: (1) complete equilibrium, (2) nonequilibrium equipartition, and (3) non-equipartition. The first case, equilibrium, was run to test whether the method remains in equilibrium for long computational times; that is, to verify that the model would not drift to improper internal distribution (ref. 10). The second case tests whether, indeed, the model has an internal mechanism to drive itself to equilibrium within a reasonable physical time scale. The third case provides insight into relative internal time scales to reach (1) a quasi-Boltzmann distribution characterized by a rotation temperature $T_{\text{rot}} \neq T$ and then (2) the time scale for this quasi-distribution to decay to equilibrium $T_{\text{rot}} = T$. The simulations, therefore, permit us to observe energy partitioning and the relaxation mechanisms, as well as relaxation rates. Because we are here interested only in rotational transitions that lead to a final equilibrium state, all elastic collisions have been ignored in order to expedite the calculations.

Equilibrium Case and Collision Frequency

Many of the physical properties for one-dimensional calculations have been based on the hyperbolic function trajectory $\text{sech}(\alpha t)$ (ref. 15); some others are based on the effective potential, and the classical equations of motion are solved for this interaction potential including the step by step energy conservation.

Collision frequency, however, cannot be evaluated analytically for these potentials (see eqs. 1-3). It is also not feasible to use the Monte Carlo method to compute and thereby describe intermolecular potentials numerically. Nevertheless, we can semi-empirically determine a macroscopically (statistically) correct collision frequency by using the results of equation (7) for the equilibrium case; that is, we determine the most probable index of power $\delta$ in equation (7) by varying $\delta = 4$ (Maxwellian molecules) to $\delta = \infty$ (hard-sphere molecules). The simulation should maintain equilibrium when the proper value of $\delta$ is chosen. The effect of the parameter $\delta$ on rotational energy is sensitive to the higher velocity collisions. We find that rotational temperature increases as the parameter $\delta$ increases. Figure 1 shows the computed rotational temperature history from an initial equilibrium state as the value of $\delta$ is varied. The proper value for which the model (point centers of repulsion) fits close to the present model, is found to be approximately $C = 0.45$, or $\delta = 7.5$, where $C = (\delta - 4)/\delta$. Also presented in the same figure are the results of the temperature variation obtained by using the linear selection formula of equation (7b) where the parameter $B$ was varied from 0 to 0.5. The best value of $B$ that satisfies this selection rule seems to be $B = 0.3$. Temperature histories computed using this value ($C = 0.45$) are shown next.

In figures 2 through 7, translational and rotational temperatures for different initial conditions are plotted as functions of nondimensional
collision time. These temperatures should asymptotically approach the equilibrium values shown by the dotted line at the end of the time scale. Also shown in figure 2 for several time intervals, \( t/\tau = 0, 5, 10, \) and \( 20 \), are the translational and rotational distribution functions. The translational distribution function \( f_{\text{tr}} \) is plotted as a function of molecular speed ratio \( c/c_0 \), where \( c_0 \) is the rms speed defined by \( \sqrt{5kT/M} \). The rotational distribution function \( f_r \) is presented in terms of the rotational energy level \( j \) for the same collision times corresponding to the translational distribution functions. Mean collision time \( \tau \) is calculated based on the initial translational temperature, kept always at room temperature \( T_{\text{tr}} = 320^\circ \text{K} \). Note that initial equilibrium distribution functions for rotational and translational energies are selected at random from the Rayleigh and Maxwellian distributions, respectively, (their distributions, therefore, do not represent analytical functions), and all subsequent distributions evolve from the present simulation using the Itikawa model.

Throughout the testing, all distributions and temperatures represented equally valid equilibrium states. Figures 3a and 3b show the comparison of the Monte Carlo solutions with theoretical (equilibrium) functions for the non-dimensional time at \( t/\tau = 15 \) and \( 20 \). Since the Monte Carlo solution - the so-called Klimontovich function (ref. 24) - does not, in general, present a smooth function, a mean distribution function (Boltzmann solution), time averaged over the last five collision times (from \( t/\tau = 15 \) to \( 20 \)), is shown in figure 3c. This mean value can be compared with the theoretical value when the gas approaches equilibrium. The agreement with theory seems to be very good.

Nonequilibrium Case

The result of the equipartition, nonequilibrium test is shown in figure 4. The gas initially starts in the equipartition state with the rotational energy of all molecules assigned the mean equilibrium energy corresponding to the level of \( j = 10 \), and where the translational energy is specified as in the previous case. Several energy distribution functions are shown at nondimensional times of \( 0, 1, 2, 3, 5, 20, \) and \( 35 \). The simulation again seems very good. The distribution functions for \( t/\tau = 30 \) and \( 35 \), and the mean distribution functions averaged over these time intervals, are compared with theoretical calculations in figure 5; once again the agreement is good. Notice that only even-numbered rotational energy levels are occupied. This follows since we have considered a homonuclear model to be initially in an even-numbered level, and we have not allowed changes in nuclear spin.

The last test investigated is for the nonequipartition and nonequilibrium case. All gases initially start with the rotational energy level of \( j = 12 \) (corresponding to a rotational temperature of \( T_r = 455^\circ \text{K} \)) and where the translational energy is selected at random from the Maxwellian distribution corresponding to a translational temperature of \( 320^\circ \text{K} \). In figure 6, both the rotational and translational temperatures approach an equilibrium state, corresponding to a temperature of \( 374^\circ \text{K} \). This occurs in an exponential manner. Transition distribution functions are shown at \( t/\tau = 0, 1, 2, 3, 5, \) and \( 20 \) in the same figure. The present model also appears to perform very
well in this case. This is the case (initial rotational temperature higher than translational temperature) for which the modified Pearson-Hansen model failed to show satisfactory performance when extended to higher temperatures (ref. 10).

The distribution functions for \( t/\tau = 15 \) and \( 20 \), and the mean distribution functions of rotational and translational energy, averaged over \( t/\tau = 15 \) to \( 20 \), are shown in figure 7. Comparison with theoretical distributions is very satisfactory.

Notice that in both figures 4 and 6 the gas relaxes asymptotically to equilibrium. The approach is rapid initially and quite slow finally.

Of all the models tested to date, the semiclassical model of Itikawa appears most satisfactory. However, this model with 1000 molecules consumes about 100 sec of the CDC 7600 machine computing time to travel one characteristic collision time. The model, therefore, may still need further simplification to permit its practical use in the more complex molecular simulations, such as, for example, shock wave structure and gas-surface interactions.

**CONCLUSION**

The Itikawa model when used with the appropriate representations for collision frequency provides an adequate physical description of a homonuclear diatomic gas in rotational relaxation. This model appears to hold the most immediate promise for further application to more complex problems. However, because of the computational time required to do rigorous calculations at each Monte Carlo collision event, subsequent investigations will require simplification of the algorithm. Nevertheless, the present model will permit one to examine the principal features of rotational effects in nonequilibrium flow of diatomic molecules, such as shock wave structure.

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APPENDIX A

INTERMOLECULAR POTENTIAL AND INTERACTION MATRIX ELEMENT

Interaction Potential

Each term of the interaction potential (eq. (23)) is assumed to be defined by the representations given in the text. The first term, $V^{(0)}$, can be determined fairly reliably from either the result of molecular beam experiments (ref. 21) or the analysis of transport coefficients (ref. 22). These parameters are $C = 3.44 \times 10^3$ eV, $C_6 = 73.4$ eV Å$^{-6}$ and $\alpha = 3.16$ Å$^{-1}$. Unfortunately, we have little information about the anisotropy of the interaction potential for $N_2 + N_2$. Therefore, we adopt the form given by $V^{(1)}$ and $V^{(2)}$ in equation (23), similarly as in reference 15, and regard $A$ and $B$ as adjustable parameters. The values employed in this report are $A = B = 0.2$.

Interaction Matrix

Applying the effective potential method (ref. 17) to the interaction potential given by equations (22) and (23), we can calculate the matrix element as follows:

$$
\langle j_1'j_2'|\tilde{V}^{\text{eff}}|j_1j_2\rangle = \langle j_1'j_2'|\tilde{V}^{\text{eff}}|j_1j_2\rangle - V^{(0)}(R)\delta_{j_1',j_1}\delta_{j_2',j_2} = (j_1'j_2'|j_1j_2)C \exp[-\alpha R(t)] (A1)
$$

$$
(j_1'j_2'|j_1j_2) = \frac{A}{\sqrt{5}} \left[ (-1)^{j_2'}(2j_2+1)^{-1/2} \begin{pmatrix} j_1 & j_1 & 2 \\ 0 & 0 & 0 \end{pmatrix} \delta_{j_2',j_2} + (-1)^{j_1}(2j_1+1)^{-1/2} \begin{pmatrix} j_2 & j_2 & 2 \\ 0 & 0 & 0 \end{pmatrix} \delta_{j_1',j_1} \right]
\left[ \begin{array}{c} 1 \\ \frac{1}{\sqrt{5}} \\ \frac{1}{\sqrt{5}} \\ \frac{1}{\sqrt{5}} \end{array} \right]
\sum_{m=0,2,4} (2m+1)^{1/2} \begin{pmatrix} 2 & 2 & m \\ 0 & 0 & 0 \end{pmatrix} (A2)
$$

where $\begin{pmatrix} j_1' & j_1 & j \\ 0 & 0 & 0 \end{pmatrix}$ is the 3-j symbol and $L = \max(j_1 + j_2, j_1' + j_2')$. 
APPENDIX B

PROGRAM LISTING FOR ROTATIONAL TRANSITION PROBABILITY CALCULATION

*DECK MAIN
PROGRAM MAIN( INPUT, OUTPUT, TAPF5=INPUT, TAPF6=OUTPUT )
CROSS SECTION CALCULATION
COMMON / LIMIT/ LL1MIN, LL1MAX, LL2MIN, LL2MAX
DIMENSION PWAVE(20,20), SIGMA(20,20), LABCSA(16)
COMMON SIG(20)
COMMON / CM1/ PWAVE, EKIN, L10, L20, BIMP, NMAX, NPRINT, L1PAR, L2PAR
COMMON / CM2/ JEL1, JEL2, LABCSA, LLMAX
COMMON / CM3/ VA, VB, IPRT1, IPRT2, IPRT3, IPRT4
COMMON / CMVR1/ VC, VALPHA, VC6, BBIMP, EEEE
COMMON / CV1/ ABCC(40,40,9)
JMAX = 20
PAI2 = 6.283185
C** INITIAL CLEAR OF ABCC, SHOULD BE MADE FOR EACH VA, VB
DO 5009 I=1,40
DO 5009 J=1,40
DO 5009 K=1,9
ABCC(I,J,K) = 0.0
5009 CONTINUE
5010 CONTINUE
DO 5016 I=1,JMAX
DO 5016 J=1,JMAX
SIGMA(I,J) = 0.0
PWAVE(I,J) = 0.0
5016 CONTINUE
C** INPUT ** IMPACT PARAMETER (IN ANGSTROM)
C** CALCULATION FROM 'BIMPI' TO 'BIMPF' WITH STEP 'DBIMP'
READ(5,4) BIMPI, DBIMP, BIMPF
4 FORMAT (3F10.0)
IF (BIMPI.LT.0.0) GO TO 5999
C** INPUT ** CROSS SECTION CALCULATION (ICROS=1) OR NOT (0)
READ(5,3) ICROS
C** INPUT ** RELATIVE KINETIC ENERGY (IN EV)
READ(5,1) EKIN
1 FORMAT (F10.0)
C** INPUT ** INITIAL ROTATIONAL STATES
READ(5,2) L10, L20
2 FORMAT (2I5)
C** INPUT ** MAX NO. OF TERMS IN EXP
   READ(5,3) NMAX
3 FORMAT(15)
C** INPUT ** INDEX FOR PRINT OUT
   READ(5,6) IPRT1,IPRT2,IPRT3,IPRT4
6 FORMAT(415)
C** INPUT ** POTENTIAL PARAMETERS FOR SPHERICAL PART
C** V(R) = VC*EXP(-VALPHA*R)-VC6/R**6
C** V IN EV, R IN ANGSTROM
   READ(5,7) VC,VALPHA,VC6
7 FORMAT(3F10.0)
C** INPUT ** POTENTIAL PARAMETERS FOR NON-SPHERICAL PART
   READ(5,5) VA,VB
5 FORMAT(2F5.0)
C** INPUT ** WHEN IPRTSG=0, PRINT PARTIAL SUM OVER BIMP
   READ(5,3) IPRTSG
C** INPUT ** LIMITATION OF RANGE OF LL1,LL2
C** LL1 = LL1MIN-LLMAX, LL2=LL2MIN-LL2MAX
   READ(5,8) LL1MIN,LL1MAX,LL2MIN,LL2MAX
8 FORMAT(415)
   LL1MIN= LL1MIN+1
   LL1MAX= LL1MAX+1
   LL2MIN= LL2MIN+1
   LL2MAX= LL2MAX+1
   J1MIN= (LL1MIN+1)/2
   J1MAX= (LL1MAX+1)/2
   J2MIN= (LL2MIN+1)/2
   J2MAX= (LL2MAX+1)/2
WRITE(6,5900)
1 BIMP1,DRIMP,BIMP2,
2 ICROS ,
3 FKN,
4 L10,L20 ,
5 NMAX,
6 IPRT1,IPRT2,IPRT3,IPRT4,
7 VC,VALPHA,VC6,
8 VA,VB,
9 IPRTSG
5900 FORMAT(1H1/5X,1OH INPUT DATA/}
1 5X,18HBIMP1,DRIMP,BIMP2/,3F10.2/}
2 5X, 6HICROS/,15/
3 5X, 5HEKIN/,F10.6/
4 5X, 8HL10,L20/,215/
5 5X, 5HNMAX/,15/
6 5X,24HIPRT1,IPRT2,IPRT3,IPRT4/,415/
7 5X,14HVC,VALPHA,VC6/,3F10.4/
8 5X,6HVA,VR/,2F5.2/
9 5X,7HPRTS/G/,15/)
WRITE(6,5902) L1MIN, L1MAX, L2MIN, L2MAX
5902 FORMAT(1HO/5X,4H L1=I4,3H - ,I4,3X,4H L2=I4,3H - ,I4/) C********** NPRINT
NPRINT= NMAX-3
BIMP= BIMPI
IF(ICROS.EQ.0) IPRT4=0
IF(ICRUS.EQ.0) GO TO 5500
H3= DBIMP/3.0*PAI2
NSUM= 0
MPRINT=0
H3B= H3*BIMP
CALL PROB
5100 CONTINUE
DO 5111 I=J1MIN,J1MAX
DO 5111 J=J2MIN,J2MAX
SIGMA(I,J)= SIGMA(I,J)+H3B*PWAVE(I,J)
5111 CONTINUE
BIMP=BIMP+DBIMP
CALL PROB
H4B=4.0*H3*BIMP
DO 5112 I=J1MIN,J1MAX
DO 5112 J=J2MIN,J2MAX
SIGMA(I,J)=SIGMA(I,J)+H4B*PWAVE(I,J)
5112 CONTINUE
BIMP= BIMP+DBIMP
CALL PROB
H3B=H3*BIMP
DO 5113 I=J1MIN,J1MAX
DO 5113 J=J2MIN,J2MAX
SIGMA(I,J)= SIGMA(I,J)+H3B*PWAVE(I,J)
5113 CONTINUE
IF(IPRTSG.EQ.1) GO TO 5120
GO TO 5300
5120 CONTINUE
IF(BIMP.GT.BIMPF-0.5*DBIMP) GO TO 5200
NSUM= NSUM+2
GO TO 5100
5200 CONTINUE
MPRINT =1
IF(IPRTSG .EQ.0) GO TO 5400
5300 CONTINUE
C******************SEARCH FOR MAX OF SIGMA
SMAX=0.0
DO 5305 I=J1MIN, J1MAX
DO 5305 J=J2MIN, J2MAX
IF(SIGMA(I,J) .GT. SMAX) SMAX= SIGMA(I,J)
5305 CONTINUE
C********** PRINT
     L1PARO=MOD(L10,2)+1
     L2PARO=MOD(L20,2)+1
     WRITE(6,5901) BIMPI,BIMP,NSUM,L10,L20,EKIN
5901 FORMAT(1H1/5X,23HCROSS SECTION FOR BIMP=F6.2,3H = ,F6.2,7H (NSUM=
       112,1H),3X,5H(L10=I2,1X,4HL20=I2,1H),3X,5H(EKIN=F13.5//3X,2HL2)
     DO 5311 JINV=J2MIN, J2MAX
       J= J2MIN+J2MAX-JINV
     L2= 2*J-3*L2PARO
C********** NORMALIZATION TO SMAX
     DO 5310 I=J1MIN, J1MAX
       SIG(I)= SIGMA(I,J)/SMAX
     5310 CONTINUE
     JF= J1MIN+15
     IF(JF.GT. J1MAX) JF= J1MAX
     WRITE(6,5911) L2,(SIG(I),I=J1MIN,JF)
5911 FORMAT(1HO,3X,I2,16(IPF6.3,1X))
     5311 CONTINUE
     WRITE(6,5912) (LABCSA(I),I=1,16)
5912 FORMAT(1HO,4X,16(3X,I2,2X),3H L1)
     SMAX= 0.1*SMAX
     WRITE(6,5919) SMAX
5919 FORMAT(1HO/3X,15HNORMALIZATION FACTOR,2X,F13.5,2X,11HANGSTROM**2)
     SEL= -SIGMA(JEL1,JEL2)
C******  ELASTIC
     WRITE(6,5918) SEL
5918 FORMAT(1HO,4X,29HTOTAL INELASTIC CROSS SECTION,2X,F13.5)
     IF(MPRINT.EQ.1) GO TO 5400
     GO TO 5120
5400 CONTINUE
C***** ROTATIONAL CONSTANT IN EV
     BROT= 0.2512E-3
     S= 0.0
     SWW= 0.0
     DO 5415 I=J1MIN, J1MAX
       LI= 2*I-3+L1PARO
       DO 5414 J=J2MIN, J2MAX
         IF(I.EQ.JEL1.AND.J.EQ.JEL2) GO TO 5414
         L2= 2*J-3*L2PARO
5414 CONTINUE
5415 CONTINUE
*DECLK PROB
SUBROUTINE PROB

C************ MAIN PROGRAM FOR THE CALC. OF TRANS. PROB.
COMMON /LIMLS/ LMIN,LLMAX,LL2MIN,LL2MAX
DIMENSION PLODD(20,20),PEV(20,20),PWA(20,20)
DIMENSION AKSUM(20,20),AKSUM1(20,20)
DIMENSION LABCSA(16)
COMMON /MV1/ AMATRX(20,20,9)
COMMON /MV2/ VBB,VAA,BROT,ETOT,BRC,VALP
COMMON /CMV1/ VC,VALPHA,VC6,BIMP,EAAA
COMMON /MV3/ NOUNT
COMMON /CM1/ PWA,EKIN,L10,L20,BB,B,NMAX,NPRINT,L1PAR,L2PAR
COMMON /CM2/ JEL,JEL2,LABCSA,LLMAX
COMMON /CM3/ VA,VB,IPRT1,IPRT2,IPRT3,IPRT4
   ICLOCK=0
   BIMP=BBB
   JMAX=20
   LMAX= 2*JMAX-2
C************ REDUCED MASS IN AMU
   RMASS= 14.02
C************ ROTATIONAL CONSTANT IN EV
   BROT= 0.2512E-3
VVALP= 0.045723*VALPHA/ SQRT(RMASS)
VAA= 0.4472136*VA
VBB= 0.2*VB=0.6298283
ETOT= EKIN+ FLOAT(L10*(L10+1)+L20*(L20+1))*BROT
EB= ETOT/BROT
EB1= SQRT(EB)
LLMAX= INT(EB1)+1
IF(IPRT1.EQ.1) GO TO 11
WRITE(6,901) BIMP,L10,L20,EKIN,ETUT,LLMAX,EB,VC,VALPHA,VC6,VA,VB
901 FORMAT(1H1,5X,5HBIMP=F6.2,5HANGST,5X,6H( L10=I2,5H L20=I2,2H )
1 5X,5HEKIN=F13.5,3HEV ,3X,5HETOT=E13.5,3X,6HLLMAX=I2,
2 3X,3HEB=F8.2/6X,3HVC=F10.2,2X,7HVALPHA=F8.3,2X,4HVC6=F7.2,2X,
3 3HVA=F7.2,2X,3HVB=F7.2/)
11 CONTINUE
IF(LLMAX.GT.LMAX) LLMAX=LMAX
IF(LL1MAX.GT.LLMAX) LL1MAX= LLMAX
IF(LL2MAX.GT.LLMAX) LL2MAX= LLMAX
DC 18 I=1,JMAX
DO 18 J=1,JMAX
PEVN(I,J)= 0.0
PADD(I,J)= 0.0
PWAVE(I,J)= 0.0
AKSUMO(I,J)=0.0
AKSUM1(I,J)= 0.0
DO 16 M=1,9
AMATRIX(I,J,M)= 0.0
16 CONTINUE
18 CONTINUE
L10= L10+1
L20= L20+1
JFL1=(L10+1)/2
JEL2=(L20+1)/2
L1PAR= MOD(L10,2)+1
L2PAR= MOD(L20,2)+1
IF(L1PAR.LT.LL1MIN) L1PAR= LL1MIN
IF(L2PAR.LT.LL2MIN) L2PAR= LL2MIN
CL0= FLOAT((2*L10+1)*(2*L20+1))
PE0= 0.0
C*********** N= 1
N=1
LL1I= LL10
LL2I= LL20
NCOUNT= 0
DC 59 K=1,9
CALL VMATRIX(LL1I,LL2I,K)
59 CONTINUE
   LL1FMN= MAX0(LL10-2,LL1PAR)
   LL1FMX= MIN0(LL10+2,LL1MAX)
   LL2FMN= MAX0(LL20-2,LL2PAR)
   LL2FMX= MIN0(LL20+2,LL2MAX)
   K= 1
   JJ11= (LL1I+1)/2
   JJ12= (LL2I+1)/2
   DO 64 I=1,3
   LL2F= LL20-4+2*I
   DO 63 J=1,3
   LL1F= LL10-4+2*J
   IF(LL2F.LT.1.OR.LL1F.LT.1) GO TO 62
   JJF1= (LL1F+1)/2
   JJF2= (LL2F+1)/2
   AKSUMO(JJF1, JJF2) = AMATRX(JJI1, JJ12, K)
62 K=K+1
63 CONTINUE
64 CONTINUE
   TCTPW= 0.0
   TCTPC= 0.0
   DO 74 LL1F=LL1FMN,LL1FMX,2
   DO 73 LL2F=LL2FMN,LL2FMX,2
   JJF1= (LL1F+1)/2
   JJF2= (LL2F+1)/2
   PODD(JJF1, JJF2)= AKSUMO(JJF1, JJF2)
   PEVN(JJF1, JJF2)= 0.0
   IF(LL1F.EQ.LL1I.AND.LL2F.EQ.LL2I) PEVN(JJF1, JJF2)=1.0
   PWAVE(JJF1, JJF2)= PODD(JJF1, JJF2)**2*4.0+PEVN(JJF1, JJF2)**2
   P=PWAVE(JJF1, JJF2)
   L01=LL1F-1
   L02=LL2F-1
   IF(IPRT2.EQ.1 ) GO TO 72
   PCORCT= P
   WRITE(6,902) N, L01, L02, P, PCORCT, PODD(JJF1, JJF2), AKSUMO(JJF1, JJF2)
902 FORMAT(1H0,2X,2HN=12,3X,5H L1F=I2,2X,5H L2F=I2,5X,6HPWAVE=E13.5, 1 3X,7HPCORCT=E13.5,3X,5HPODD=E13.5,5X,6HAKSUM=E13.5)
72 CONTINUE
   TOTPW= TOTPW+P
73 CONTINUE
74 CONTINUE
   IF(IPRT1.EQ.1 ) GO TO 75
   WRITE(6,904) N, TOTPW, ICLOCK, NCOUNT, TOTPC
904 FORMAT(1H0,1X,1H0,1H0,1H0,1H0,1H0,12X,1H0,1H0,1H0,1H0,1H0,1H0,1H0,1H0,1H0,1H0,1H0)
75 CONTINUE
   IF(NMAX.EQ.1 ) GO TO 300
N = 2
C**************************100
100 CONTINUE
   NCCUNT = 0
   N2 = 2*N-2
   LL2IMX= MAX0(LL20-N2,N2PAR)
   LL2IMX= LL20+N2
   IF(LL2IMX .GE. LLMAX) LL2IMX= LLMAX
   DO 149 LL2I=LL2IMN,LL2IMX,2
       JJ2= (LL2I+1)/2
       E2= SQRT(EFLOAT(LL2I-1)**2)
       LLMAX1= INT(E2)+1
       LL1I=LL10+N2
       JJ1I=(LL1I+1)/2
       IF(LLMAX1 .GE. LLMAX1) LLMAX1= LLMAX1
       IF(LL1I .GT. LLMAX1) GO TO 130
   DO 114 K=5,8,3
       CALL VMA TRX(LL1I,LL2I,K)
   114 CONTINUE
   DO 119 K=3,9,3
       CALL VMA TRX(LL1I,LL2I,K)
   119 CONTINUE
   IF(LL1I-2 .LT. 1) GO TO 125
   AMATRX(JJ1I,JJ2I,4) = AMATRX(JJ1I-1,JJ2I,6)
   K=4
   IF(LL2I .EQ. LL20-N2 .OR. LL2I .EQ. LL20+N2) CALL VMA TRX(LL1I,LL2I,K)
   AMATRX(JJ1I,JJ2I,7) = AMATRX(JJ1I-1,JJ2I+1,1)
   K=7
   IF(LL2I .EQ. LL20-N2 .OR. LL2I .EQ. LL20+N2) CALL VMA TRX(LL1I,LL2I,K)
   IF(LL2I-2 .LT. 1) GO TO 130
   AMATRX(JJ1I,JJ2I,1) = AMATRX(JJ1I-1,JJ2I-1,9)
   K=1
   IF(LL2I .EQ. LL20-N2 .OR. LL2I .EQ. LL20+N2) CALL VMA TRX(LL1I,LL2I,K)
   125 CONTINUE
   IF(LL2I-2 .LT. 1) GO TO 130
   AMATRX(JJ1I,JJ2I,2) = AMATRX(JJ1I,JJ2I-1,8)
   K=2
   IF(LL2I .EQ. LL20-N2) CALL VMA TRX(LL1I,LL2I,K)
   130 CONTINUE
   LL1I=LL10-N2
   JJ1I= (LL1I+1)/2
   IF(LL1I .LT. L1PAR) GO TO 149
   DO 134 K=5,8,3
       CALL VMA TRX(LL1I,LL2I,K)
   134 CONTINUE
DO 139 K=1,7,3
CALL VMATRX(LL1I,LL2I,K)
139 CONTINUE
AMATRX(JJ1I,JJI2,6)= AMATRX(JJ1I+1,JJI2,4)
K = 6
IF(LL2I.EQ.LL20-N2.OR.LL2I.EQ.LL20+N2) CALL VMATRX(LL1I,LL2I,K)
AMATRX(JJ1I,JJI2,9)= AMATRX(JJ1I+1,JJI2+1,1)
K=9
IF(LL2I.EQ.LL20+N2-2.OR.LL2I.EQ.LL20-N2) CALL VMATRX(LL1I,LL2I,K)
IF(LL2I-2.LT.1) GO TO 149
AMATRX(JJ1I,JJI2,2)= AMATRX(JJ1I,JJI2-1,8)
K = 2
IF(LL2I.EQ.LL20-N2) CALL VMATRX(LL1I,LL2I,K)
AMATRX(JJ1I,JJI2,3)= AMATRX(JJ1I+1,JJI2-1,7)
K = 3
IF(LL2I.EQ.LL20-N2+2.OR.LL2I.EQ.LL20-N2) CALL VMATRX(LL1I,LL2I,K)
149 CONTINUE
C********* 150
LL1IMN = MAXO(LL10-N2+2, L1PAR)
LL1IMX = LL10+N2-2
IF(LL1IMX.GE.LL1MAX) LL1IMX = LL1MAX
DC 199 LL1I=LL1IMN, LL1IMX, 2
JJ1I = (LL1I+1)/2
EB2 = SQRT(EB- FLOAT(LL1I-1)**2)
LLMAX1 = INT(EB2)+1
LL2I = LL20+N2
JJ1J = (LL2I+1)/2
IF(LLMAX1.GE.LL2MAX) LLMAX1 = LL2MAX
IF(LL2I.GT.LLMAX1) GO TO 180
DO 164 K=5,9
IF(LL1I.EQ.LL10+N2-2.OR.K.EQ.6) GO TO 163
CALL VMATRX(LL1I,LL2I,K)
GO TO 164
163 CONTINUE
AMATRX(JJ1I,JJI2,6)= AMATRX(JJ1I+1,JJI2,4)
164 CONTINUE
IF(LL2I-2.LT.1) GO TO 175
AMATRX(JJ1I,JJI2,2)= AMATRX(JJ1I,JJI2-1,8)
AMATRX(JJ1I,JJI2,3)= AMATRX(JJ1I+1,JJI2-1,7)
IF(LL1I-2.LT.1) GO TO 180
AMATRX(JJ1I,JJI2,1)= AMATRX(JJ1I-1,JJI2-1,9)
175 CONTINUE
IF(LL1I-2.LT.1) GO TO 180
AMATRX(JJ1I,JJI2,4)= AMATRX(JJ1I-1,JJI2,6)
180 CONTINUE
LL2I=LL20-N2
JJ12= (LL2I+1)/2
IF(LL2I.LT.L2PAR) GO TO 199
DO 184 K=5,6
IF(LL1I.EQ.LL10+N2-2.AND.K.EQ.6) GO TO 183
CALL VMATRX(LL1I,LL2I,K)
GO TO 184
183 CONTINUE
AMATRX(JJ1I,JJ12,6)= AMATRX(JJ1I+1,JJ12,4)
184 CONTINUE
DO 189 K=1,3
CALL VMATRX(LL1I,LL2I,K)
189 CONTINUE
AMATRX(JJ1I,JJ12,8)= AMATRX(JJ1I,JJ12+1,2)
AMATRX(JJ1I,JJ12,9)= AMATRX(JJ1I+1,JJ12+1,1)
IF(LL1I-2.LT.1) GO TO 199
AMATRX(JJ1I,JJ12,4)= AMATRX(JJ1I-1,JJ12,6)
AMATRX(JJ1I,JJ12,7)= AMATRX(JJ1I-1,JJ12+1,3)
C*********** 199
199 CONTINUE
LL1FMX=LL10+2*N
LL2FMX= LL20+2*N
LL1FMN= LL10-2*N
IF(LL1FMN.LT.L1PAR) GO TO 201
LL1I=LL1FMN+2
200 CONTINUE
LL2FMN=LL20-2*N
IF(LL2FMN.LT.L2PAR) GO TO 202
LL22= LL2FMN+2
GO TO 205
201 CONTINUE
LL1FMN= L1PAR
LL1I= L1PAR
GO TO 200
202 CONTINUE
LL2FMN=L2PAR
LL22= L2PAR
205 CONTINUE
LL22F= LL2FMX-2
IF(LL2FMX.LE.LL2MAX) GO TO 206
LL2FMX= LL2MAX
LL22F= LL2MAX
206 CONTINUE
DO 249 L2=LL22,LL22F,2
LL11F= LL1FMX-2
EB2 = SQRT(EB - FLOAT(L2-1)**2)
LLMAX1 = INT(EB2) + 1
IF(LLMAX1 .GE. LLMAX) LLMAX1 = LLMAX
IF(LL1F .GE. LLMAX1) LL1F = LLMAX1
DO 248 L1 = LL11, LL1F, 2
  K1 = 1
  K2 = 0
L22 = L2 - 2
L11 = L1 - 2
IF(L2 .LT. LL1F) L22 = LL1F
IF(L1 .LE. LL1F) L11 = LL1F
IF(L2 .LE. LL1F) K1 = 4
IF(L1 .LE. LL1F) K2 = 1
K = K1
J1 = (L1+1)/2
J2 = (L2+1)/2
AK = AKSUM0(J1, J2)
L2P2 = L2 + 2
DO 219 LL2 = L22, L2P2, 2
  K = K+K2
  L1P2 = L1 + 2
  DO 218 LL1 = LL11, L1P2, 2
    JJ1 = (LL1+1)/2
    JJ2 = (LL2+1)/2
    AKSUM1(JJ1, JJ2) = AK*AMATRX(J1, J2, K)+AKSUM1(JJ1, JJ2)
    K = K+1
218 CONTINUE
219 CONTINUE
C************ PRINT OF AMATRX
IF(N .LT. IPRT3) GO TO 248
L01 = L1 - 1
L02 = L2 - 1
IF(L2 .GE. LL22F-1 .AND. LL20+2*N .LE. LLMAX+2) GO TO 220
IF(L1 .GE. LL11F-1 .AND. LL10+2*N .LE. LLMAX+2) GO TO 220
IF(L2 .EQ. LL22 .AND. LL20-2*N .GE. LL1F-2) GO TO 220
IF(L1 .EQ. LL11 .AND. LL10-2*N .GE. LL1F-2) GO TO 220
GO TO 248
220 CONTINUE
WRITE(6, 906) L01, L02
906 FORMAT(1H0, 5X, 15HCHECK OF AMATRX, 5X, 3HL1=I2, 2X, 3HL2=I2)
WRITE(6, 907) (AMATRX(J1, J2, J), J=1, 9)
907 FORMAT(1H5, 2X, 9E13.5)
248 CONTINUE
249 CONTINUE
C************ 250
LL1FMX = LL10+2*N
TOTPW= 0.0
DO 299 LL2F=LL2FMN,LL2FMX,2
LL1F= LL1FMX
EB2= SQRT(EB- FLOAT(LL2F-1)**2)
LLMAX1= INT(EB2)+1
IF(LLMAX1.GE. LL1MAX) LLMAX1= LL1MAX
IF(LL1FMX.GE. LL1MAX1) LL1F= LL1MAX1
DO 298 LL1F=LL1FMN,LL1F,2
JJF1= (LL1F+1)/2
JJF2= (LL2F+1)/2
AKSUM1(JJF1,JJF2)= AKSUM1(JJF1,JJF2)/ FLOAT(N)
IF(MOD(N,2).NE.0) GO TO 294
AKSUM1(JJF1,JJF2)= -4.0*AKSUM1(JJF1,JJF2)
PEVN(JJF1,JJF2)= PEVN(JJF1,JJF2)+ AKSUM1(JJF1,JJF2)
PWAVE(JJF1,JJF2)= PADD(JJF1,JJF2)**2*4.0*PEVN(JJF1,JJF2)**2
GO TO 295
294 CONTINUE
PADD(JJF1,JJF2)= PADD(JJF1,JJF2)+ AKSUM1(JJF1,JJF2)
PWAVE(JJF1,JJF2)= PADD(JJF1,JJF2)**2*4.0*PEVN(JJF1,JJF2)**2
295 CONTINUE
P= PWAVE(JJF1,JJF2)
L01= LL1F-1
L02= LL2F-1
C******* PRINT OF PWAVE
IF(N.LE.NPRINT) GO TO 297
IF(PE.GE.0.1E-4) GO TO 296
IF(LL1F.EQ.LL1FMN) GO TO 296
IF(LL1F.GT.LL1FMX-2) GO TO 296
GC TO 297
296 CONTINUE
PCORCT= P
WRITE(6,902) N,L01,L02,P,PCORCT,PADD(JJF1,JJF2),AKSUM1(JJF1,JJF2)
297 CONTINUE
TOTPW= TOTPw+P
AKSUM0(JJF1,JJF2)= AKSUM1(JJF1,JJF2)
AKSUM1(JJF1,JJF2)= 0.0
298 CONTINUE
299 CONTINUE
PE1=PWAVE(JEL1,JEL2)
PE10= ABS((PE1-PEO)/PE1)
IF(IPRT1.EQ.1) GO TO 3904
WRITE(6,904) N,TOTPw,ICLOCK,NCOUNT,TOTPC
904 FORMAT(1H0//2X,8H**** N=I2,5X,13HSUM OF PWAVE= El3.5,
15X,7HICLOCK=I10,3HSEC,5X,7HNCOUNT=I10,5X,6HTOTPC=El3.5)
CONTINUE
IF(AOS(T0TPW-1.0) LT 0.1E-3. AND PE1. LT 0.1E-3) GO TO 300
IF(N .EQ. NMAX) GO TO 300
N=N+1
PE0= PE1
GC TO 100

300 CONTINUE
C********** ELASTIC
DELST1= PEVN(JEL1, JEL2)
DELST2= -2.0*PODD(JFL1, JEL2)
C********** FINAL PRINT
MPRINT=0
L1PARO= MOD(110, 2)+1

301 CONTINUE
WRITE(6,901) BIMP, LLC, L20, EKIN, ETOT, LLMAX, EB, VC, VALPHA, VC6, VA, VB
WRITE(6,910) N, TOTPW, ICLOCK, TOTPC

910 FORMAT(1H5, 5X, 5HNMAX=I2, 2X*, 6HTOTPW=E13.5, 2X, 6HCULOCK=I5, 3HSEC, 1)
2X, 6HCULOCK=E13., 5//3X, 2HLZ)
IMN= LL2FMN
IMX= LL2FMX
IF(MOD(LL2FMX-L2PAR, 2).NE. 0) IMX= LL2FMX-1
DO 319 I= IMN, IMX, 2
LL2F= IMX+IMN-1
J2= (LL2F+1)/2
LL1F= LL1FMX
EB2= SQRT(EB- FLOAT(LL2F-1)**2)
LLMAX1= INT(EB2)+1
IF(LLMAX1.GE.LL1MAX) LLMAX1= LL1MAX
IF(LL1FMX.GE.LLMAX1) LL1F= LLMAX1
J1MX= (LL1F+1)/2
J1Mx1= J1MX
J1MN= (L1PAR+1)/2
IF(J1MX-J1MN.GT.15) J1MX= J1MN+15
LO2= LL2F-1
WRITE(6,911) LO2, (PWRITE(J1, J2), J1=J1MN, J1MX)

911 FORMAT(1HO/3X, I2, 16(1PF6.3, 1X))
IF(MPRINT.EQ.1) GO TO 319
L1MIN= MIN0(LL20, LO2)
CL10= FLOAT(2*L1MIN2+1)
DO 318 J1= J1MN, J1MX1
L01= 2*J1-3+L1PARO
L1MIN= MIN0(L10, L01)
CL1= CL10* FLOAT(2*L1MIN+1)
P= PWAVE(1, J2)
PCCRC= CL1/CL0*P
TOTPC = TOTPC + PCORCT
PWAVE(J1, J2) = PCORCT

318 CONTINUE
319 CONTINUE
LABCSA(1) = L1PAR-1
DO 320 I=2,16
LABCSA(1) = LABCSA(1) + (I-1)*2
320 CONTINUE
WRITE(6,912) (LABCSA(J),J=1,16)
912 FORMAT(1H0,4X,16(3X,I2,2X),3H L1)
IF(MPRINT.EQ.1) GO TO 9906
C******* PRINT OF PCORCT WITH PELASTIC MODIFIED
PWAVE(JEL1,JEL2) = 1.0 - TOTPC + PWAVE(JEL1,JEL2)
IF(IPRT4.EQ.1) GO TO 1000
MPRINT=1
GO TO 301
9906 CONTINUE
WRITE(6,9909) TOTPC
9909 FORMAT(1H0/2X,35H PCORCT WITH PELASTIC MODIFIED, 5X, 1 7H(TOTPC=EL3.5,2H )/
1000 CONTINUE
PWAVE(JEL1,JEL2) = PWAVE(JEL1,JEL2) - 1.0
WRITE(6,9911) DELST1,DELST2
9911 FORMAT(1H0/1X,28H FOR ELASTIC SCATTERING, 3X, 3HD1=EL3.5, 1 2X, 3HD2=EL3.5)
RETURN
END

*DECK VMAT
SUBROUTINE VMATRIX(LL1,LL2,K)
COMMON /MV1/ AMATRIX(20,20,9)
COMMON /MV2/ VBB,VAA,BROT,ETOT,BRC,VVALP
COMMON /CMVR1/ VC,VALPHA,VC6,BIMP,EIJ
COMMON /MV3/ NCOUNT
COMMON /CV1/ ABCC(40,40,9)
JJ1 = (LL1+1)/2
JJ2 = (LL2+1)/2
L1J = LL1-1
L2J = LL2-1
IF(K.GT.3) GO TO 1011
L2I = L2J-2
IF(L2I.LT.0) GO TO 1510
GO TO 1100
1011 IF(K.GT.6) GO TO 1012
L2I = L2J
GC TO 1100
1012 L2I = L2J+2
1100 CONTINUE
   IF(MOD(K,3).NE.1) GO TO 1111
   L1I = L1J-2
   IF(L1I.LT.0) GO TO 1510
   GO TO 1200
1111 IF(MOD(K,3).NE.2) GO TO 1112
   L1I = L1J
   GO TO 1200
1112 L1I = L1J+2
1200 CONTINUE
   ABC = ABC(L1I,L2I,K)
   IF(ABC.NE.0.0) GO TO 1299
   CALCULATION OF VEFF(L1I,L2I / L1J,L2J )
       CC1 = CG20(L1I,L1J)
       CC2 = CG20(L2I,L2J)
1211 CONTINUE
   IF(K.LE.5) CSIGN = (-1.0)**(L1J+L2J)
   IF(K.GE.6) CSIGN = (-1.0)**(L1I+L2I)
   C = FLOAT((2*L1I+1)*(2*L2I+1)*(2*L1J+1)*(2*L2J+1))**0.25 * CSIGN
1212 CONTINUE
   B = VBB*CC1*CC2
1213 CONTINUE
   A = 0.0
   IF(L2I.NE.L2J) GO TO 1215
   A = VAA*CC1/SQRT(FLOAT(2*L2I+1))
   IF(MOD(L2I,2).NE.0) A = -A
1215 IF(L1I.NE.L1J) GO TO 1219
   AA = VAA*CC2/SQRT(FLOAT(2*L1J+1))
   IF(MOD(L1J,2).NE.0) AA = -AA
   A = A+AA
1219 CONTINUE
   ABC = C*(B+A)
   ABC(L1I,L2I,K) = ABC
1299 CONTINUE
   WI = BROAT*FLOAT(L1I*(L1I+1)+L2I*(L2I+1))
   WJ = BROAT*FLOAT(L1J*(L1J+1)+L2J*(L2J+1))
   W1J = ABS(WI-WJ)
   E1 = ETOT-WI
   EJ = ETOT-WJ
   IF(E1.LE.0.0.OR.EJ.LE.0.0) GO TO 1510
   E1J = 0.5*(E1+EJ)
   CALL ROOT(RC)
BRC = 1.0 - (BIMP/RC)**2 + VC6/EIJ/RC**6
EIJ1 = EIJ*BRC
IF(K.EQ.5) GO TO 1500
IF(LII.EQ.L2J.AND.K.EQ.3) GO TO 1500
IF(LII.EQ.L2J.AND.K.EQ.7) GO TO 1500
ATJ = VVALP* SQRT(EIJ1)/WIJ
EW = EIJ1/WIJ
DBALPH = EW/AIJ
APAI = 1.570796327/AIJ
F = EXP(-APAI)
FAIJ = 2.0*APAI*F/(1.0-F*F)
AAA = ABC*DBALPH*FAIJ
C*** 999 CHECK PRINT
999 CONTINUE
1498 CONTINUE
NCOUNT = NCOUNT+1
1499 CONTINUE
AMATRX(JJ1, JJ2, K) = AAA
RETURN
1500 CONTINUE
AAA = ABC/VVALP* SQRT(EIJ1)
GO TO 1498
1510 CONTINUE
AAA = 0.0
GO TO 1499
END

SUBROUTINE ROOT(RC)
C**** REVISED 6/26/74
COMMON /CMVR1/ VC, VALPHA, VC6 , BIMP , EKIN
RMIN = 1.12
RR0 = ALOG(VC/EKIN)/VALPHA
IF(RR0.GE.4.1) RR0 = 4.1
N = 1
3099 CONTINUE
RR=RR0
3100 CONTINUE
RL = EKIN*BIMP**2/RR**2
V = VC* EXP(-VALPHA*RR)
VL = -VALPHA*V
IF(VC6.EQ.0.0) GO TO 3101
VR = VC6/RR**6
V = V-VR
VL = VL+6.0*VR/RR
3101 CONTINUE
  F = (V+RL-EKIN)/(2.0*RL/RR-V1)
  IF(ABS(F/RR).LT.0.1E-5) GO TO 3199
  IF(N.GE.100) GO TO 3299
  RR = RR+F
  IF(RR.LT.RMIN) GO TO 3900
  N=N+1
  GO TO 3100
3199 CONTINUE
  RC = RR
  RETURN
3299 CONTINUE
  WRITE(6,998) RR,F,RR0
998 FORMAT(1HO/5X,14HERROR N GT 100,3X,3HRR=F13.5,3X,2E13.5/)
  RR=RR0
  GO TO 3199
3900 CONTINUE
  RRG = 0.5*(RR0+RMIN)
  GO TO 3099
END

*DECK CG20
FUNCTION CG20(J1,J2)
  IF(J2.EQ.J1+2.OR.J2.EQ.J1-2) GO TO 8001
  IF(J2.EQ.J1) GO TO 8002
  C = 0.0
  GO TO 8100
8001 CONTINUE
  IF(J2.EQ.J1+2) J=J1
  IF(J2.EQ.J1-2) J = J2
  X1= FLOAT(J+2)/FLOAT(2*J+5)
  X2= FLOAT(J+1)/FLOAT(2*J+3)
  X3= 1.0/FLOAT(2*J+1)
  C = SQRT(1.5*X1*X2*X3)
  GO TO 8099
8002 CONTINUE
  J= J1
  X1= FLOAT(J+1)/FLOAT(2*J+3)
  X2= FLOAT(J)/FLOAT(2*J+1)
  X3= 1.0/FLOAT(2*J-1)
  C = - SQRT(X1*X2*X3)
8099 IF(MOD(J,2).NE.0) C = -C
8100 CG20= C
RETURN
END
REFERENCES


Figure 1.- Variation of rotational temperature depending on specified values for the potential parameters.
Figure 2.- Temperature and distribution functions: Complete equilibrium initial conditions.
Figure 3.- Translational and rotational distribution functions: Complete equilibrium initial conditions.
Figure 3.- Continued.

(b) $t/\tau = 20$

Figure 3.- Continued.
(c) Mean value.

Figure 3.- Concluded.
Figure 4.- Temperature and distribution functions: Equipartition initial conditions.
Figure 5.- Translational and distribution functions: Equipartition initial conditions.

(a) $t/\tau = 30$
Figure 5.- Continued.

(b) $t/\tau = 35$

Figure 5.- Continued.
Figure 5.- Concluded.

(c) Mean value.
Figure 6.- Temperature and distribution functions: Nonequipartition initial conditions.
Figure 7.- Translational and rotational distribution functions: Nonequipartition initial conditions.
Figure 7.- Continued.

(b) $t/\tau = 20$

Figure 7.- Continued.
Figure 7.- Concluded.

(c) Mean value end.
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—National Aeronautics and Space Act of 1958

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