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A069156	TECHNICAL REPORT ARBRL-TR-02148 THE H+02+OH+O REACTION, A COMPARISON OF POTENTIAL SURFACES
C FILE COPY	March 1979 DDC DC MAY 30 1979 B
	US ARMY ARMAMENT RESEARCH AND DEVELOPMENT COMMAND BALLISTIC RESEARCH LABORATORY ABERDEEN PROVING GROUND, MARYLAND
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REPORT DUCUMENTATION PAGE	READ INSTRUCTIONS
. REPORT NUMBER /2. GOVT AC	CESSION NO. 3. RECIPIENT'S CATALOG NUMBER
Technical Report ARBRI-TR-02148	
. TITLE (and Subtitio)	5. TYPE OF REPORT & PERIOD COVERED
The H+O <sub>2</sub> $\rightarrow$ OH+O Reaction, A Comparison of	
Potentiál Surfaces	BRL Report
	6. PERFORMING ORG. REPORT NUMBER
AUTHOR(a)	8. CONTRACT OR GRANT NUMBER(*)
Dr. Arthur Gauss, Jr.	
	10 PROGRAM ELEMENT PROJECT TASK
IC Army Dellistic Descende Laboratory	AREA & WORK UNIT NUMBERS
ATTN. DRDAR_BIR	
Aberdeen Proving Ground, MD 21005	RDT&E 1L161102AH43
1. CONTROLLING OFFICE NAME AND ADDRESS	12. REPORT DATE
US Armament Research & Development Command	MARCH 1979
US Army Ballistic Research Laboratory	13. NUMBER OF PAGES
ATTN: DRDAR-BL, APG, MD 21005 4. MONITORING AGENCY NAME & ADDRESS(II dillerent from Control	Iling Office) 15. SECURITY CLASS. (of this report)
	Unclassified
	15a. DECLASSIFICATION/DOWNGRADING SCHEDULE
6. DISTRIBUTION STATEMENT (of this Report)	
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Specific rate constants are calculated and the overall thermal rate constant is estimated at  $1600^{\circ}$  and  $2500^{\circ}$ K. In addition a difficulty with the single parameter Surface I has been corrected yielding Surface I'. Specific rates and an overall rate are estimated using those surfaces as well.

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## I. INTRODUCTION

The H+O<sub>2</sub>  $\rightarrow$  OH + O reaction is an endothermic (~ 17 kcal/mole) reaction<sup>1</sup> important in combustion. It continues to appear in reaction schemes<sup>2-4</sup> describing various combustion processes.

Trajectory calculations have been used to calculate cross sections, specific rate constants and estimates of the overall thermal rate constant for the reaction. A new potential surface (Surface II) has been used which fits the known properties of the HO<sub>2</sub> complex better than two

previous surfaces (Surfaces I and I'). The new surface is described by two adjustable Sato parameters instead of one as in the two previous surfaces.

Surface I and associated results were discussed in a previous paper<sup>5</sup> (called Paper I). Surface I' corrects a problem which was subsequently found in Surface I. Trajectory results for Surface I' are reported in this report along with those for Surface II. The classical trajectory program CLASTR<sup>6</sup>, suitably modified for these LEPS surfaces, was used for all the trajectory calculations. Surface I' shows little difference in the trajectory results relative to Surface I. It also has similar potential contours for all configurations compared.

The results of trajectory calculations on Surface II indicate enhancement of the reaction rate due to vibrational excitation of the  $O_2$ , but the magnitude of this effect is considerably less than on the previous single parameter surfaces. This vibrational enhancement of the rate is in agreement with other studies<sup>7,8</sup> on endothermic reactions.

- JANAF Tables, edited by D.R. Stull (Dow Chemical, Midland, MI, 1965-1967).
- M.B. Collset, D.W. Naegeli, and I. Glassman, Sixteenth Symposium on Combustion (Combustion Institute, Pittsburgh, 1977) p. 1023.
- T. Miyauchi, Y. Mori, and A. Imamura, Sixteenth Symposium on Combustion, (Combustion Institute, Pittsburgh, 1977) p.1073.
- 4. G. Dixon-Lewis and R.J. Simpson, Sixteenth Symposium on Combustion, (Combustion Institute, Pittsburgh, 1977) p. 1111.
- 5. A. Gauss, Jr., J. Chem. Phys. 68, 1689 (1978).
- 6. CLASTR is program No. 229 in the Quantum Chemistry Program Exchange Catalogue (QCPE, Indiana University, Chemistry Department).
- D.S. Perry, J.C. Polanyi and C. Woodrow Wilson, Jr., Chem. Phys. 3, 317 (1974).
- R.N. Porter, L.B. Sims, D.L. Thompson, and L.M. Raff, J. Chem. Phys. 58, 2855 (1973).

The results show clearly that the most significant changes in specific reaction rates between Surface II and Surfaces I and I' occur for the lower vibrational states (v=0,1,2) of the  $0_2$  whereas the higher states

(v=3,4,5,6) show less of an effect. Surface II is quite different from Surfaces I and I' in the high interaction region (all nuclei close together). That even small differences in the shape of the potential in this region can affect the reactivity in the lowest vibrational level has been demonstrated by other workers<sup>9,10</sup>.

### II. SURFACE I', DESCRIPTION AND TRAJECTORY RESULTS

The trajectory results for Surface I' are summarized in Tables I and II, similar to the presentation of results in Paper I. As in that study trajectories are run in sets for which the sum of the relative translational energy  $(E_{\rm p})$  plus the vibrational energy  $(E_{\rm v})$  is constant.

The majority of trajectory points were run with rotational quantum number J=1 (corresponding to about a hundreth of a kcal/mole of rotational energy). A number of points were run with J=21 (about 2 kcal/mole of rotational energy). Three types of trajectory events are defined: reactive, nonreactive, and complex. Complex trajectories are long trajectories that the CLASTR integrator cannot follow; they are not back integrable (energy and angular momentum are conserved). Both reactive and non-reactive events are back integrable. The complex trajectories have been further subdivided (Table II) into reactive complex and nonreactive complex trajectories according to the decision of the CLASTR integrator. Observe that in addition to some points for checking Surface I' results against those of Surface I (Paper I) additional points were run on Surface I' for vibrational levels v=3,2 and 1. They fill in nicely between the v=4 and v=0 results of Surface I. (See Paper I.) Specific rate constants have been evaluated for the v=3,2 and 1 levels assuming, as in Paper I, a constant average cross section over all translational energies down, of course, to the classical threshold. The maximum impact parameter used for the cross section calculations was 3 A. These specific rate constants are tabulated in Table III for two temperatures, 1600°K and 2500°K.

The difficulty with Surface I was that it had an infinity in the derivatives of the potential in the linear configuration  $(\theta=180^{\circ})$ . Surface I' eliminates this infinity. This infinity causes little difference in the trajectories results between Surfaces I and I' since the system very rarely approaches the linear or near linear configuration even on Surface I'. Because of the large potential hole the system

N. Sathyamurthy, J.W. Duff, C. Stroud, and L.M. Raff, J. Chem. Phys. 67, 3563 (1977).

N. Sathyamurthy, R. Rangarajan, and L.M. Raff, J. Chem. Phys. 64, 4606 (1976).

definitely prefers bent configurations. Also Surfaces I and I' are nearly identical in well depth and equilibrium configuration. The potential contours are similar for all values of the H-O-O angle.

The only difference in the mathematical form of the two surfaces I and I' is in the Sato parameter. The Sato parameter for Surface I' is

$$\Delta' = \left\{ 1 - e^{-\left( \frac{x_{a'b}}{b} \right)^2} \right\} \left\{ 0.53(\sin 2\pi h)^2 - 0.25(\cos \pi h)^2 + 0.2(\frac{\pi}{2})^2 \right\}$$

TABLE I. Summary of the reactive and complex trajectories for Surface'. The total number of trajectories for each initial 0, vibra-

E <sub>R</sub> +E <sub>v</sub>	02			
kcal	Vibrational			Total
mole	Level	Reactions	Complexes	Trajectories
		J=1 RESULTS		
33	6	16	16	300
29	3 2 1	1 1 0	6 0 0	500 500 500
24	{ 4 3	7 4	24 15	300 500
22	{ 2 1	1 0	10 1	500 500
18	3 2 1	1 0 1	72 18 3	500 500 500
14	{ 2 1	0 0	44 18	500 500
		J=21 RESULTS		
33	6	15	12	300

tional level in each energy set  $(E_R + E_v)$  are given also.

E <sub>R</sub> +E <sub>v</sub>	02			
kcal mole	Vibrational Level	Total Complexes	Reactive Complexes	Non-Reactive Complexes
		J=1 RESU	LTS	
33	6	16	4	12
29	$\begin{cases} 3\\ 2 \end{cases}$	6 0	1 0	5 0
	(1	0	0	0
24	{ 4 3	24 15	5 2	19 13
22	{ <sup>2</sup> 1	10 1	1 0	9 1
18	$ \left\{\begin{array}{c} 3\\2\\1 \end{array}\right. $	72 18 3	5 1 0	67 17 3
14	{ 2 1	44 18	0 0	44 18
		J=21 RESU	LTS	
33	ó	12	4	8

TABLE II.	The complex th	rajectories ta	bulated in	Table	I are divided
	into reactive	and non react	ive groups	using	the decision
	of the CLASTR	integrator.			

TABLE III. Specific rate constants (at 1600°K and 2500°K) for Surface I' are given for  $0_2$  initially in the vibrational levels v=3,2,1. The populations of the vibrational levels relative to the v=0 level are shown. All calculations were done with an initial  $0_2$  rotational quantum number (J) of one. The errors quoted are the Monte Carlo statistical errors.

Temperature (°K)	Vibrational Level	Specific Rate Constant cm <sup>3</sup> molecule sec	Relative Level Population
160091	. 3	$(6.7 \pm 4.4) \times 10^{-12}$	.016
1000 K	1	$(1.3 \pm 1.3) \times 10$ $(0.29 \pm 0.29) \times 10^{-12}$	.063
2500°K	3 2 1	$\begin{array}{rrrr} (8.3 & \pm & 5.5) \times & 10^{-12} \\ (1.9 & \pm & 1.9) \times & 10^{-12} \\ (0.61 & \pm & 0.61) \times & 10^{-12} \end{array}$	.073 .171 .410

where

$$b = 0.6 \times 10^{-8}$$
 cm.

$$Th = Arcsin(r_1/x_2 sin \theta)$$

and

3

$$r_a = (r_1^2 + r_2^2/4 - r_1r_2 \cos \theta)^{\frac{1}{2}}$$

See Fig. 1 for a pictorial representation of the parameters  $x_{a}^{},$  Th,  $\theta,$   $r_{1}^{},$   $r_{2}^{},$  and  $r_{3}^{}.$ 

The total rate constant may be evaluated using the specific rate constant data from Surface I' (Table III) and Surface I (Paper I). The total rate constant is given by (only odd J appear in the sum for  $0_2$ )

where

$$K(T) = \sum_{v,J} F_{BC}(v,J) k_{vJ}$$
$$F_{BC} = \frac{f_J(2J+1)e}{Q_{Jv}}$$

 $F_{BC}$  is the rotational-vibrational distribution function (Boltzmann distribution) and  $Q_{Jv}$  is the rotational-vibrational partition function. The total rate constant evaluated from the trajectory results for Surfaces I and I' at 1600°K is given by

$$K_{\rm T}(1600^{\circ}K) = (3.0 \pm 2.0) \times 10^{-13} \frac{{\rm cm}^3}{{\rm molecule sec}}$$

This value has been evaluated neglecting the possible contribution of reactive complexes. The error quoted is the Monte Carlo statistical error. The contribution from the v=0 level has been taken as zero. The experimental value of Schott<sup>11</sup> is

$$K_{\rm E}(1600^{\circ}{\rm K}) = 1.35 \times 10^{-12} \frac{{\rm cm}^3}{{\rm molecule sec.}}$$

At 2500°K the rate constant from the trajectory results of Surfaces I and  $I^\prime$  is

$$K_{\rm T}(2500^{\circ}{\rm K}) = (1.7 \pm 0.9) \times 10^{-12} \frac{{\rm cm}^3}{{\rm molecule sec}}.$$

11. G.L. Schott, Combustion and Flame, 21, 357 (1973).





The sea burners

The experimental result of Schott<sup>11</sup> is

$$K_{\rm E}(2500^{\circ}{\rm K}) = 5.9 \times 10^{-12} \frac{{\rm cm}^3}{{\rm molecule sec.}}$$

As with the  $1600^{\circ}$ K result the trajectory result is a minimum rate constant (neglecting reactive complexes). If the decision of the CLASTR integrator in regard to the reactivity of the complexes is assumed correct, then both trajectory results would be doubled, bringing the trajectory results within a factor of two of experiment.

#### III. SURFACE II

Surface II contains two adjustable Sato parameters. The general LEPS potential for a three atom system has the following form  $^{12}$ 

$$V = Q_{1}' + Q_{2}' + Q_{3}' - (\alpha_{1}'^{2} + \alpha_{2}'^{2} + \alpha_{3}'^{2} - \alpha_{1}' \alpha_{2}' - \alpha_{1}' \alpha_{2}' - \alpha_{1}' \alpha_{2}' + D_{2}^{e}$$

where

$$Q_{i}' = \frac{Q_{i}}{1+\Delta_{i}} = \frac{D_{i}^{e}}{4(1+\Delta_{i})} \left\{ \left(3+\Delta_{i}\right) e^{-2\beta_{i}(r_{i}-r_{io})} - \left(2+6\Delta_{i}\right) e^{-\beta_{i}(r_{i}-r_{io})} \right\}$$
  
and  
$$\alpha_{i}' = \frac{\alpha_{i}}{1+\Delta_{i}} = \frac{D_{i}^{e}}{4(1+\Delta_{i})} \left\{ \left(1+3\Delta_{i}\right) e^{-2\beta_{i}(r_{i}-r_{io})} - \left(6+2\Delta_{i}\right) e^{-\beta_{i}(r_{i}-r_{io})} \right\}$$

i = 1,2,3.

The Morse parameters for the two diatomics OH and  $\mathbf{0}_2$  are given in Paper I.

In the present case the two Sato parameters are: 1)  $\Delta = \Delta_1 = \Delta_3$ (OH parameter) and 2)  $\Delta_2$  (O<sub>2</sub> parameter). The Sato parameter  $\Delta$  is closely similar to the  $\Delta'$  defined for Surface I' in the previous section. The Sato parameter  $\Delta$  is given by

 $\Delta = 0.7 \left\{ \frac{-\left( \frac{x_{a}}{b} \right)^2}{1 - e^{-\left( \frac{x_{a}}{b} \right)^2}} \right\} \left\{ 0.53(\sin 2\pi h)^2 - 0.25(\cos \pi h)^2 + 0.2(\pi/2)^2 \right\}$ (see Fig. 1).

12. J.T. Muckerman, J. Chem. Phys. 56, 2997 (1972).

The other Sato parameter  $\Delta_2$  is given by

$$\Delta_{2} = \left\{ \sin(c \cdot r_{2}) + .15 \cdot \left[ \sin(c \cdot r_{2}) \right]^{8} \right\}$$
  
\* 
$$\left\{ 0.53(\sin 2Th)^{2} - 0.25(\cos Th)^{2} + 0.2 \left(\frac{\pi}{2}\right)^{2} \right\}$$

where

 $c = \pi/5.2 \times 10^{-8} cm.$ 

The other parameters, Th and  $r_2$  have been defined previously (Fig. 1).

Surface II yields an equilibrium position for HO<sub>2</sub> in better agreement with recent experimental results than either Surface I or I'. For Surface II the minimum energy is at  $r_1=0.99$ ,  $r_2=1.30$ ,  $\theta=105^\circ$  versus experimental values of  $r_1 = 0.985$ ,  $r_2 = 1.36$ ,  $\theta = 106^{\circ}$ .<sup>13</sup> Surface II is also a better fit to the shape of the potential minimum than either I or I'. The force constants reported from the analysis of experimental results are  $f_{OH} = 6.5 \text{ mdyne/Å}$ ,  $f_{OO} = 5.8 \text{ mdyne/Å}$  and  $f_{\theta} = 1.088 \text{ mdyne/Å}$ .<sup>14</sup> The values from the new two parameter LEPS Surface II are  $f_{OH}$ = 6.1 mdyne/Å,  $f_{OO}$ =5.1 mdyne/Å, and  $f_{\theta}$ =0.65 mdyne/Å. The new two parameter surface, like the previous surfaces, fits the angular variation of the potential as given by Gole and Hayes (Fig. 2)<sup>15</sup>. For  $\theta \leq 95^\circ$  there is some divergence between Surface II and the CI results; the LEPS fitting function is not sufficiently flexible to closely follow the ab initio data in this region. However, the CI calculation itself is very limited, using only minimal basis set SCF generated orbitals. The differences between Surface II and Surface I can be seen by comparing the potential plots for Surface II at  $\theta$ =80,108,140, and 180° (Figs. 3, 4, 5, 6) with the corresponding plots in Paper I.

#### IV. TRAJECTORY RESULTS FOR SURFACE II.

The trajectory results for Surface II are summarized in Tables IV and V. The mode of presentation is similar to that in Paper I and to that for Surface I' in Section II of this article. Trajectories were run for

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Figure 2. Comparison (at  $r_1 = 0.96$ Å,  $r_2 = 1.23$ Å) of the angular dependence of the LEPS potential energy Surface II with the ab initio surfaces calculated by Gole and Hayes.







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Figure 4. Contour diagram (in eV) of the LEPS potential Surface II for  $\theta=108^{\circ}$ ; the potential well is near its greatest depth (~2 eV.) at this angle.



Figure 5. Contour diagram (in eV) of the LEPS potential Surface II for  $\theta{=}140^\circ.$ 





E <sub>R</sub> +E <sub>v</sub>	°2			
kca1	Vibrational			Total
mole	Level	Reactions	Complexes	Trajectories
		J=1 Resu	lts	
38	6	22	0	500
33	6	26	5	500
	6	22	12	500
29	23	17	2	750
	0	10	5	1350
24	13	14	10	750
24	10	16	5	1350
10	13	3	26	750
10	10	4	30	1350
		J = 21 R	esults	
33	6	34	5	500
1.0	13	8	30	750
10 ,	10	7	24	1350

TABLE IV. Displayed here are the reactive and complex trajectories for Surface II. The total number of trajectories is also given.

TABLE V. The complex trajectories for Surface II shown in Table IV are here divided into reactive and non-reactive groups according to the decision of the CLASTR integrator.

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$\frac{E_R + E_v}{\frac{kcal}{mole}}$	0 <sub>2</sub> Vibrational Level	Total Complexes	Reactive Complexes	Non-Reactive Complexes
		J=1 RESU	ILTS	
38	6	0	0	0
33	6	5	0	5
29	$ \left\{\begin{array}{c} 6\\ 3\\ 0 \end{array}\right. $	12 2 5	2 0 1	10 2 4
24	$\begin{cases} 3 & \cdot \\ 0 & \cdot \end{cases}$	10 5	1 0	9 5
18	$\begin{cases} 3\\ 0 \end{cases}$	26 30	1 3	25 27
		J-21 RES	ULTS	
33	6	5	0	5
18	$\begin{cases} 3\\ 0 \end{cases}$	30 24	2 2	28 22

three different values of  $E_R + E_v$  ( $E_R$ , relative translational, +  $E_v$ , vibrational energy) for each vibrational level v=0, 3 and 6. Trajectories were run for one value of  $E_R + E_v$  for the v=4 level. The minimum cross section data of Fig. 7 are fitted with the following function

$$S_{r_{vJ}} = S_{r_{vJ}}^{o} + \frac{1}{\kappa} \tan^{-1} \left\{ \lambda (E - E_{o}) \right\}$$

The constants in this expression for the various vibrational levels are shown in Table VI. Interpolation was used to determine the constants for the levels for which there is no data (v=1,2,5) or only one datum (v=4). Note that the v=4 trajectory point fits the interpolated curve well. The thresholds for each cross section function of the three lowest levels (v=0,1,2) are a few tenths of a kcal/mole below the classical thresholds. From these analytical cross section curves the specific rate constants may be derived using the usual expression

$$k_{vJ} = \left\{ \frac{8kT}{\pi\mu_A, BC} \right\}^{\frac{1}{2}} \int_{O} S_{r_{vJ}} f(T, E_R) dE_R$$

where v is the vibrational quantum number of  $0_2$ , J is the rotational quantum number of  $0_2$  (always odd either 1 or 21), T is the absolute temperature,  $\mu_{A,BC}$  is the reduced mass of H, $0_2$ ,  $E_R$  is the collision energy (relative translational energy), and  $S_{r_{VJ}}$  is the cross section given by the function previously described. Also

$$f(T,E_R) = \frac{E_R}{(kT)^2} e^{-E_R/kT}$$

The specific rate constants are shown in Table VII at two temperatures,  $1600^{\circ}$ K and  $2500^{\circ}$ K. If rotational effects can be neglected and assuming a Boltzmann distribution of vibrational energy, the overall thermal rate constant can be estimated (assuming also vibrational levels above v=6 make negligible contributions).

In the case of the v=6 and v=0 states the J=21 data do not show statistically greater reactivity than the J=1 data. For the v=6, J=1 state  $S_{r_{6,1}} = (1.47 \pm 0.28) \text{Å}^2$ , for v=6, J=21,  $S_{r_{6,21}} = (1.92 \pm 0.32) \text{Å}^2$ , for the v=0, J=1 state  $S_{r_{0,1}} = (.0838 \pm 0.0418) \text{Å}^2$ , and for the v=0, J=21 state we have  $S_{r_{0,21}} = (.1466 \pm 0.0552) \text{Å}^2$ . For the v=3 state the J=21

### TABLE VI. The parameters for the cross section fitting function

 $S_{r_{vJ}} = S_{vJ}^{0} + \frac{1}{\kappa} \tan^{-1} \lambda(E-E_{0})$  are given here for each vibrational level. Parameters were adjusted for cross sections fits to the v=0,3 and 6 levels data only, the parameters for other levels were interpolated. Rotational quantum number J is equal to one (J=1).

0 <sub>2</sub>	S <sup>°</sup> r <sub>v</sub> J		Eo
Vibrational Level	(J=1) (Å <sup>2</sup> )	к (А <sup>-2</sup> )	kcal mole
6	.60	122.5	-8.0
5	.49	150	-3.7
4	. 388	189	0.7
3	. 30	245	5.0
2	.228	322	9.4
1	.176	418	1.37
ō	.15	490	18.0

 $\lambda = 1/2$  FOR ALL LEVELS

data show statistically somewhat greater reactivity than the J=1 data. The results are for the J=1 state  $S_{r_{3,1}} = (1.131 \pm 0.652) \text{Å}^2$  and for J=21,  $S_{r_{3,21}} = (3.016 \pm 1.061) \text{Å}^2$ . In all cases above the J=21 data are larger  $r_{3,21}$  than the J=1 results. The total thermal rate constant at 1600°K and

than the J=1 results. The total thermal rate constant at 1000 K and 2500°K will be calculated assuming no rotational dependence but the above data indicate that these total thermal rates may be 50 to 100% low due to rotation. This rotational dependence should be examined with more care in the future.

The total rate constant at 1600°K from the trajectory work on Surface II is

 $K_{\rm T}(1600^{\circ} \text{K}) = (1.45 \pm 0.49) \times 10^{-2} \frac{\text{cm}^3}{\text{molecule sec.}}$ 

This result is quite close to the experimental value given earlier (see Section II). It must be kept in mind that reactive complexes have been neglected in this calculation and these will augment the result, as most likely will the rotational contribution. TABLE VII - Specific rate constants (at 1600°K and 2500°K) for Surface II are given. These are derived from the cross section curves shown in Fig. 7. Errors are Monte Carlo errors. All calculations were done with initial 0<sub>2</sub> rotational quantum

Temperature(°K)	Vibrational Level	Specific Rate Constant cm <sup>3</sup> molecule sec.	Relative Level Population
	6	$(7.4\pm1.5) \times 10^{-11}$	$3.12 \times 10^{-4}$
	5	$(5.9\pm1.5) \times 10^{-11}$	$1.13 \times 10^{-3}$
	4	$(4.1\pm1.2) \times 10^{-11}$	$4.22 \times 10^{-3}$
1600°K	3	$(2.0\pm0.7) \times 10^{-11}$	$1.6 \times 10^{-2}$
	2	$(6.4\pm2.2) \times 10^{-12}$	6.3 $\times 10^{-2}$
	1	$(1.8\pm0.7) \times 10^{-12}$	$2.46 \times 10^{-1}$
	0	$(5.1\pm1.8) \times 10^{-13}$	1
	6	$(9.4\pm1.9) \times 10^{-11}$	$5.71 \times 10^{-3}$
	5	$(7.5\pm1.7) \times 10^{-11}$	$1.3 \times 10^{-2}$
	4	$(5.5\pm1.4) \times 10^{-11}$	$3.02 \times 10^{-2}$
2500°K	3	$(3.2\pm0.9) \times 10^{-11}$	$7.27 \times 10^{-2}$
	2	$(1.5\pm0.5) \times 10^{-11}$	$1.71 \times 10^{-1}$
	1	$(6.6\pm2.1) \times 10^{-12}$	$4.1 \times 10^{-1}$
	0	$(3.0\pm1.0) \times 10^{-12}$	1

number J = 1. Vibrational level populations relative to the v=0 level are also shown.

At 2500°K the total rate constant is given by -12 cm<sup>3</sup>

 $K_{\rm T}(2500^{\circ}K) = (7.9 \pm 2.4) \times 10^{-12} \frac{{\rm cm}^3}{{\rm molecule sec.}}$ 

This result is some 30% higher than the experimental result of Schott quoted earlier (Section II). The trajectory result is low because of the reasons stated above for  $K_T(1600^{\circ}K)$  and because of the neglect of vibrational levels above v=6 which will contribute more significantly



Figure 7. The minimum cross sections are shown as a function of translational energy. Monte Carlo statistical error bars are included at each trajectory point. The data are fitted with the functions  $S_{r_{vJ}} = S_{vJ}^{o} + \frac{1}{\kappa} \tan^{-1} \left\{ \lambda(E-E_{o}) \right\}$  where the

constants are displayed in Table VI.

than at 1600°K. The error result quoted with each rate constant is the Monte Carlo statistical error inherent in the trajectory method.

#### V. DISCUSSIONS AND CONCLUSIONS

By using two Sato parameters which are functions of the coordinates an improved LEPS surface (Surface II) for  $HO_2$  has been constructed. The

equilibrium geometry of this surface agrees more closely with experiment than the geometry of either Surface I or I'. However, the greatest improvement is in the force constants for the  $HO_2$  complex which are much better than those of either Surface I or I' and match the experimental

values quite well.

As for Surfaces I and I' the trajectory results for Surface II show significant enhancement in the H+O2 reaction rate due to vibrational excitation of the O<sub>2</sub> molecule. An enhancement factor of ~ 150 at 1600°K and ~ 30 at 2500°K (between the v=0 and v=6 levels) is achieved for Surface II as indicated by the specific rate constants. These factors are not nearly so large as those for Surfaces I and I'. The reason they are not so large is clear from the specific reaction rate tables. At the v=6 level the specific rate constants (at 1600° and 2500°) for Surface II are very nearly equal to those for Surface I. However, as the vibrational level is decreased the Surface II specific rates do not drop nearly so rapidly. Other workers<sup>9,10</sup> have demonstrated that small changes in the inner reflective wall of potential surfaces can dramatically affect reactivity in the lowest vibrational level. The same sort of vibrational effect is noted here but the effect extends to levels above the lowest level. However, there are rather significant differences in the potential contours between Surface II and Surfaces I and I' in the high interaction region in our case.

An added benefit of the new potential surface (Surface II) is the dramatic decrease in complexes formed for  $O_2$  initially in the upper

vibrational levels (v=3 and above). Thus the minimum cross sections and specific rate constants quoted for these levels are less uncertain due to complexes than those rates in Paper I. Even if all the complexes were assumed to be reactive (a worst possible case) then the specific rate constant for the v=3 level would be some two (at  $2500^{\circ}$ K) or three (at  $1600^{\circ}$ K) times larger than the minimum rates. If a more reasonable assumption is made, such as taking the decision of the CLASTR integrator as correct for the complex trajectories, then the minimum rate constant is only some 10% low (at both  $1600^{\circ}$ K and  $2500^{\circ}$ K). The v=6 level is considerably less uncertain than the v=3 level. Even for the worst case (all complexes reactive) for v=6, the minimum rate would be low by some 25% at  $2500^{\circ}$ K and 10% at  $1600^{\circ}$ K. For the v=0 level there is a significant

increase in complexes for Surface II over Surface I but there is also a considerable increase in reactivity. In the v=0 level if all the complexes were reactive the specific rate constant would be some three (at  $2500^{\circ}$ K) or four (at  $1600^{\circ}$ K) times larger than the minimum rate. Assuming here that the CLASTR integrator decision is correct for the complex trajectories, then the v=0 specific rate constant is only some 20 percent low at  $2500^{\circ}$ K and  $30^{\circ}$  low at  $1600^{\circ}$ K. In conclusion, it can be said in view of the above results for the specific rate constants that the total minimum constant rate would be in the worst case a factor of three too low due to the complexes. It seems much more likely to be low by some 20 to  $30^{\circ}$ . It would be nice to eliminate the complex problem altogether with an improved integrator for the trajectories. Since the last paper (Paper I) several variable step integrators 15, 17 have been tried with no improvement over the CLASTR integrator.

As noted above the reactivity of at least some of the complexes causes the total rate constants calculated in Section III to be low. Also these rates will probably be low because of the greater reactivity of higher rotational states. As noted in Section III the higher rotational levels (data for J=21 level) are more reactive than the lower levels (data for J=1 level) perhaps by a factor of two. More trajectories should be run at various rotational levels to see if this tendency to greater reactivity is real or just due to statistical error.

Also the contribution of higher vibrational levels above v=6 should be investigated particularly at 2500°K. However, including these high vibrational levels (v > 6) means that some method for including the H +  $O_2(^1\Delta_p)$  reaction would also have to be devised.

While it appears that the total rate constants calculated from trajectories for Surface II will be higher than experiment, the rate constant from the data for Surfaces I and I' will be low. Total rate constants estimated for Surfaces I and I' are some 25% of the experimental rates (at 1600°K and 2500°K) if the complexes are neglected. If it is assumed that the CLASTR integrator gives at least a statistically correct decision then these rates will be 50% of experiment.

As more accurate potential surface data become available for this reaction it should be possible to further refine the modified LEPS surface. Better dynamics calculations should then be possible.

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## ACKNOWLEDGEMENTS

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The author would like to thank Dr. Thomas O'Brien for reviewing some of the calculations of the potential surface derivatives. The author appreciates also the suggestions of Dr. George F. Adams.

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