

Ab Initio Energies and Product Branching Ratios for the O + C₃H₆ Reaction

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Received: July 13, 2007; In Final Form: September 13, 2007

Intermediate and transition-state energies have been calculated for the O + C₃H₆ (propene) reaction using the compound ab initio CBS-QB3 and G3 methods in combination with density functional theory. The lowest-lying triplet and singlet potential energy surfaces of the O–C₃H₆ system were investigated. RRKM statistical theory was used to predict product branching fractions over the 300–3000 K temperature and 0.001–760 Torr pressure ranges. The oxygen atom adds to the C₃H₆ terminal olefinic carbon in the primary step to generate a nascent triplet biradical, CH₃CHCH₂O. On the triplet surface, unimolecular dissociation of CH₃CHCH₂O to yield H + CH₃CHCHO is favored over the entire temperature range, although the competing H₂CO + CH₃CH product channel becomes significant at high temperature. Rearrangement of triplet CH₃CHCH₂O to CH₃CH₂CHO (propanal) via a 1,2 H-atom shift has a barrier of 122.3 kJ mol^{−1}, largely blocking this reaction channel and any subsequent dissociation products. Intersystem crossing of triplet CH₃CHCH₂O to the singlet surface, however, leads to facile rearrangement to singlet CH₃CH₂CHO, which dissociates via numerous product channels. Pressure was found to have little influence over the branching ratios under most conditions, suggesting that the vibrational self-relaxation rates for $p \leq 1$ atm are negligible compared to the dissociation rates.

1. Introduction

The gas-phase reaction of O(³P) with short-chain alkenes C_nH_{2n} is fundamental in physical chemistry and in combustion. Ethene, C₂H₄, is a key intermediate in the oxidation of CH₄ and longer-chain hydrocarbons.^{1,2} The O–C_nH_{2n} system involves a large number of possible rearrangements, dissociations, and secondary reactions that can complicate determining the kinetic parameters and product species arising from the initial step. Mechanistically, the basics of the O + C_nH_{2n} reaction series are reasonably well understood, especially for the simplest case, O + C₂H₄. The O atom adds to the terminal olefinic carbon to form a triplet biradical RCH₂–CH₂O that undergoes rearrangement and/or dissociation involving a large number of reaction intermediates and transition states,^{3–5} complicated by the presence of the lower-energy singlet surface accessible by intersystem crossing from the nascent triplet biradical. Experiments have shown that for the prototypical O + C₂H₄ system, the initially formed CH₂–CH₂O triplet biradical follows two principal reaction pathways, (1) direct dissociation to H + CH₂CHO, the vinyloxy radical, and (2) intersystem crossing to form singlet CH₂–CH₂O, followed by 1,2 H-atom transfer and C–C bond cleavage to yield CH₃ + HCO.^{3,4,6–8}

High-energy collisions of O with C_nH_{2n} species, among other small molecules and radicals, are also of increasing interest in the context of the near-space environment. Above 80 km altitude, O atoms are prevalent in the atmosphere due to solar VUV photodissociation of O₂.⁹ Collisions of O with molecular species associated with space vehicle operation in low-earth orbit can produce highly excited, radiating products, owing to very

large relative collision velocities. For instance, collision of ambient O with desorbed H₂O from the Space Shuttle results in electronically excited OH(A) giving rise to A–X emission in the near UV and in physical excitation of several vibrational modes of H₂O, resulting in emission in the 1.7–6.3 μm near- and mid-infrared spectral region.¹⁰ For a satellite in a circular orbit at 300 km, the orbital velocity is 7.726 km s^{−1}, corresponding to a 3.59 eV center-of-mass (COM) collision energy for the O + C₃H₆ reaction assuming a stationary O atom. If thrusters are fired, the approximately 3.5 km s^{−1} exhaust velocity can add or subtract from the orbital velocity depending on the “attack angle” relative to the spacecraft orbit vector, giving rise to COM collision energies up to several electronvolts larger.¹¹ Partially or unburned fuel constituents in thruster exhaust can result in efficient production of spectrally bright emitters upon reaction with O. Laboratory data confirming this effect have been obtained for collisions of hyperthermal O with CH₄,¹² C₂H₂,¹³ hydrazines,¹⁴ N₂,¹⁵ and CO₂.¹⁶

Atomic oxygen has also been implicated in the degradation of low-earth orbit spacecraft surfaces.^{11,17} These effects can include inelastic scattering, chemical reactions with surfaces, surface-catalyzed reactions, and sputtering. For instance, osmium films, which are relatively inert for most ground-based applications, oxidize and degrade in low-earth orbit, releasing gaseous OsO₄. Hydrocarbon-based polymers are also slowly oxidized by impinging atomic oxygen to produce gas-phase metastables which could exhibit reactivity similar to the O + C_nH_{2n} system. At the same time, surface imperfections such as scratches, pin defects, and overall roughness can result in a drastically increased rate of degradation due to trapped oxygen atoms which can undercut an otherwise highly protective surface coating.

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The $O + C_nH_{2n}$ reaction for $n = 2-4$ has been studied in a number of laboratory experiments utilizing various techniques to form O atoms, notably discharge or Hg-sensitized flow¹⁸⁻²⁰ and laser photolysis of a NO_2 or SO_2 precursor.²¹⁻²⁴ Using the latter technique, Bersohn and co-workers reported detailed product state results for the reaction of O with several alkenes in a series of landmark publications.^{4,5,25-27} A number of crossed-beam experiments spanning almost four decades have also been reported.^{6-8,28-32} In terms of computation, the recent work of Nguyen et al.³³ offers an excellent summary of the theoretical studies done on the benchmark $O + C_2H_4$ system, including the relatively recent calculation of the triplet surface.³⁴ The Nguyen et al. reference sets out a comprehensive high-level study of intermediate and transition-state energies for the $O + C_2H_4$ system for both the triplet and singlet surfaces and goes on to predict product branching ratios using RRKM unimolecular reaction rate theory.

Given the ongoing importance and rich chemistry inherent in the series of $O + C_nH_{2n}$ reactions, together with the large amount of experimental data, it makes sense to use currently available computational tools to better understand the stationary states and likely reaction pathways following formation of the nascent triplet biradical. The Nguyen et al. theoretical study mentioned above demonstrates the tools and methods that can be applied to these systems. While that work focused on the prototypical $O + C_2H_4$ reaction, the existing information regarding the higher alkenes suggests that ab initio computations and RRKM theory be applied to longer-chain alkenes to gauge the similarities and differences between the electronic structure and product state predictions. This work applies similar ab initio quantum mechanical calculations and RRKM unimolecular rate theory to the next larger system, $O + C_3H_6$.

II. Methodology

II.A. Quantum Chemistry Models. The reaction system was modeled using the Gaussian 03 software package³⁵ running on a Pentium-based personal computer, together with GaussView for visualization. Three levels of calculation were pursued, density functional theory (DFT) and the compound complete basis set with quadratic configuration interaction (CBS-QB3)³⁶ and Gaussian-3 (G3)³⁷ ab initio methods. The DFT method was used to survey the potential energy surface (PES) through the calculation of energies for intermediates and transition states and to follow the internal reaction coordinate (IRC). The CBS-QB3 and G3 methods were used to obtain more accurate energies than those provided by DFT-level results. When possible, the average of the calculated CBS-QB3 and G3 energy values was input into the RRKM analysis, described below. The lowest-lying triplet and singlet surfaces of the $O-C_3H_6$ system were investigated.

The Becke three-parameter/Lee-Yang-Parr density functional (B3LYP)³⁸ was used within the DFT approach, with unrestricted electron pairing on both the triplet and singlet surfaces. As discussed below, the B3LYP results provided important inputs for the higher-level methods. The method includes correlation interactions while running at a speed comparable to lower-level Hartree-Fock calculations. A 6-31+G(d,p) basis set was used in the B3LYP method for the initial survey. Energies for individual molecules were found by using a geometry optimization. The sum of the energy of any two species, molecule, radical, or atom, was taken as the energy of the two species at infinite separation. All species were treated as neutrals.

The energy along the various reaction pathways was calculated by first locating a transition state. The transition state was

found using the quadratic synchronous transit (QST2) method within Gaussian that requires input of reactant and product geometries and energies. The transition state was confirmed by checking that the resulting Hessian contained only one imaginary frequency, consistent with a first-order saddle point on the PES. To ensure that the transition state connected the assumed reactants and products, an IRC calculation was performed to follow the energy path from the transition-state structure to the reactants and products located on opposite sides of the PES saddle point.

Intermediates and transition states that were amenable to the DFT method using the B3LYP/6-31+G(d,p) level of calculation in the initial PES survey were also examined at a higher level of theory. The B3LYP method with a larger basis set, 6-311+G(3df,2p), was used to calculate species energies. The initial B3LYP/6-31+G(d,p) survey had found transition-state geometries with enough accuracy that the "Berny" optimization method in Gaussian³⁹ could be employed successfully to obtain transition-state structures at the B3LYP/6-311+G(3df,2p) level using the initial geometries from the B3LYP/6-31+G(d,p) calculation.

The compound CBS-QB3 method takes advantage of the initial B3LYP geometry survey to calculate more accurate energy values.^{36,40} For finding minimum-energy conformations, the CBS-QB3 method can be run as a single job. For transition states, the CBS-QB3 method was started at its second step, a second-order Moller-Plesset (MP2) single-point energy calculation, after reading in the results of a B3LYP/6-311G(2d,d,p) geometry optimization. The transition states calculated using the B3LYP/6-311+G(3df,2p) basis were reoptimized using the 6-311G(2d,d,p) basis to be consistent with energy minima calculated using CBS-QB3/6-311G(2d,d,p). The results were then incorporated into the second step of the CBS-QB3 method.

The seven-step G3 prescription employs quadratic configuration interaction with single and double excitations and triple excitations added perturbatively (QCISD(T)).³⁷ Given that the G3 method involves a geometry optimization using a full MP2 step, it is well suited for the calculation of transition-state energies and frequencies. To calculate transition-state energies, the G3 steps were run separately, starting with Hartree-Fock and MP2 transition-state geometry optimizations followed by single-point calculations. The results of these calculations were then compiled to obtain the G3 transition-state energies as outlined by Curtiss et al.³⁷

The B3LYP/6-31+G(d,p) survey provided a fairly inexpensive method to construct potential energy surfaces for the $O-C_3H_6$ system. Nevertheless, steps for which the reaction coordinate is more complicated could not be easily characterized using the QST2 method at the B3LYP/6-31+G(d,p) level. Notably, these include reactions for which there is no or a very small potential barrier, such as the initial $O + C_3H_6$ addition step and singlet-surface homolytic bond dissociation of C_3H_6O intermediates.

On the triplet surface, the initial O-atom addition reaction required the additional efforts associated with a relaxed PES scan at the DFT level followed by a G3 IRCMax calculation. The IRCMax calculation uses the DFT geometry for subsequent single-point energy calculations rather than a geometry obtained with the MP2 method, which is the normal optimized geometry in the G3 compound calculation. This method assumes that the DFT calculations produce geometries close to those of the actual reaction coordinate and that subsequent higher-level single-point energy calculations within the compound G3 method based on

this geometry provide more accurate energies than does the DFT method by itself.⁴¹

On the singlet surface, the homolytic bond dissociation reactions were modeled using a relaxed PES with a full unrestricted MP2/6-31G(d) calculation to obtain a minimum-energy path. Transition states were found by optimizing a structure near the energy maximum using the Berny algorithm and a full unrestricted MP2/6-31G(d) calculation. The resulting transition-state structure could then be used in the remaining G3 calculations to obtain G3 energies.

II.B. RRKM Methodology. The MultiWell suite of programs was used to analyze the reaction rates and branching ratios.^{42,43} The MultiWell suite contains programs to calculate moments of inertia, internal state sums and densities, reaction rates, and branching ratios. The program is designed to solve the RRKM master equation, described in detail in a number of classic kinetics texts.^{44,45} RRKM theory uses internal state sums (for transition states) and densities (for intermediates) to calculate unimolecular reaction rates. Rotational states are included in the MultiWell program, which approximates all molecules as symmetric tops with two equal moments of inertia. This approximation is made by taking the geometric mean of the two most similar moments, input into MultiWell as a two-dimensional external moment.

Vibrational wave functions and frequencies were obtained from the Gaussian 03 results. The MP2 level of calculation performed as part of the G3 compound method provided the $3N - 6 = 24$ vibrational frequencies, less the one imaginary frequency associated with the transition states. The O–C₃H₆ system possesses internally hindered rotations, identified through visual inspection of the program animation of the internal mode displacements. For instance, triplet CH₃CH₂CHO has two hindered rotations, the first along the CH₃CH₂–CHO bond and the second along the CH₃–CH₂CHO bond, with symmetries of two and three, respectively. Gaussian 03 treats all internal degrees of freedom as harmonic oscillators, and hindered rotations appear as very low frequency vibrations when thus approximated.

For the hindered rotors, the harmonic oscillator assumption results in a grossly inaccurate calculation of the sums of states. These degrees of freedom were treated with the method described by Knyazev for one-dimensional hindered rotors.⁴⁶ The Knyazev method calculates sums of rotational states based on the potential barrier to the rotation, V . The sums are calculated from the rotational constant, B (cm^{−1}), the reduced moment of inertia, I_r (amu Å²), the effective harmonic frequency, ω (cm^{−1}), and the symmetry number n using eqs 1 and 2

$$B = \frac{16.85763}{I_r} \quad (1)$$

$$V = \frac{1}{B} \left[\frac{\omega}{n} \right]^2 \quad (2)$$

The moments of inertia for the two-dimensional external moment and for the two hindered rotors were calculated using the coordinates obtained from the G3 calculations. The coupling between the internal rotors was ignored, and they were treated as two independent one-dimensional modes.

Once all of the internal degrees of motion were properly assigned, the sums and densities of states were calculated using MultiWell routines based on the Stein–Rabinovitch extension to the Beyer–Swinehart algorithm.⁴⁷ MultiWell uses a two array system, one array for lower energy and a second array for higher energy. A grain size of 10 cm^{−1} with 250 bins was used for the

lower-energy array for energies from 0 to 2500 cm^{−1}, while a grain size of 341.365 cm^{−1} with 250 bins was used for the higher-energy array up to 85000 cm^{−1}. The program interpolates values to produce an output with 10 cm^{−1} resolution from 0 to 85000 cm^{−1}. The RRKM master equation was solved using Gillespie's exact stochastic method.^{48,49}

For the pressure dependence, the collider was taken to be CH₃CH(O)CH₂ (propylene oxide). The Lennard-Jones collision parameters σ and ϵ/k_B for CH₃CH(O)CH₂ were estimated by adding the difference between the values for CH₂(O)CH₂ and CH₂CH₂ to those for C₃H₆, resulting in $\sigma = 4.6$ Å and $\epsilon/k_B = 475$ K. The collisional energy transfer efficiencies were calculated using a monoexponential model⁵⁰ with the exponential constant $\alpha = 100 + 0.015^*E$ cm^{−1}, where E is the internal energy. Reaction rates and branching ratios in the 300–3000 K temperature and 0.001–760 Torr pressure ranges were investigated, with the assumption that the system is chemically activated through O + C₃H₆ recombination. Note that the internal energy in the O + C₃H₆ system at a given temperature can be approximated as

$$E = E_v + E_{R,T} = \sum_j \frac{v_j e^{-v_j/kt}}{1 - e^{-v_j/kt}} + 3kT \quad (3)$$

where the sum is over the vibrational modes j and $3kT$ accounts for the external rotational and translational degrees of freedom. The 300–3000 K temperature range corresponds to a range of 10.6–427 kJ mol^{−1} internal energy in the O + C₃H₆ system.

III. Results and Discussion

III.A. Calculated Energy Values. The calculated intermediate and transition-state energy values for the O–C₃H₆ system are summarized in Tables 1 and 2 for the triplet and singlet surfaces, respectively. The B3LYP results are reported for two different basis sets and with and without zero-point energy correction. The corresponding energy levels are plotted in Figure 1 (triplet system) and Figure 2 (singlet system), with solid lines indicating reaction pathways that may or may not be favorable. In the tables and figures, reaction intermediates corresponding to energy minima are denoted by ³INT(i) for the triplet states and ¹INT(i) for the singlet states. Likewise, transition states corresponding to first-order energy saddle points are denoted ³TS(i) and ¹TS(i) for the triplets and singlets, respectively. The energy zero is defined as the sum of the O(³P) and C₃H₆ separated reactant energies for both surfaces.

III.A.1. Triplet Surface. From Figure 1, it can be seen that the O–C₃H₆ triplet surface is characterized by three bound intermediates, denoted ³INT(1), ³INT(2), and ³INT(3). The initial step involves addition of the O atom to the terminal carbon of the C=C bond in propene, generating the relatively stable secondary carbon biradical ³INT(2). The reaction coordinate is complicated by an initial energy minimum at ³INT(1), which is a weakly bound complex with a <10 kJ mol^{−1} barrier to further rearrangement. The reaction of O + C₃H₆ to form the CH₃CHCH₂O triplet biradical ³INT(2) is therefore effectively without barrier. However, ³INT(1) is important with respect to constructing the RRKM model discussed below.

Fortuitously, ³TS(1) was found using the QST2 method during the initial B3LYP/6-31+G(d,p) survey. The B3LYP/6-311++G-(3df,2p) method failed to find ³TS(1), as did the B3LYP/6-311G(2d,d,p) step within the CBS-QB3 method. Since the CBS-QB3 method optimizes initial geometries with a DFT step, it is not surprising that both the CBS-QB3 and DFT methods can

TABLE 1: Triplet Surface Ab Initio Energies (kJ mol⁻¹, 0 K) Relative to O(³P) + CH₃CHCH₂ (¹A), Computed Using the B3LYP (Two Different Basis Sets), CBS-QB3, and G3 Methods; For the Blank Entries, the Structure and Energy Could Not Be Determined

species		B3LYP				CBS-QB3	G3
		6-31+(d,p)		6-311++(3df,2p)			
		w/o ZPE	w/ ZPE	w/o ZPE	w/ ZPE		
O(³ P) + CH ₃ CHCH ₂ (¹ A)		0.0	0.0	0.0	0.0	0.0	0.0
O(³ P) + CH ₃ CHCH ₂ (¹ A)	³ INT(1)						-12.3
CH ₃ CHCH ₂ O (³ A)	³ INT(2)	-123.5	-124.2	-123.8	-125.0	-104.3	-99.8
CH ₃ CH ₂ CHO (³ A)	³ INT(3)	-148.7	-144.2	-151.3	-147.4	-127.3	-124.3
CH ₃ CH (³ A) + H ₂ CO (¹ A')		-23.2	-38.3	-30.7	-46.4	-30.1	-29.2
CH ₃ CHCHO (² A) + H(² S)		-62.5	-85.1	-71.9	-94.7	-84.4	-78.3
CH ₃ (² A) + CH ₂ CHO (² A'')		-113.6	-132.5	-120.7	-139.9	-114.9	-115.1
CH ₃ CH ₂ (² A') + HCO (² A')		-105.0	-124.0	-116.6	-135.9	-111.5	-111.3
CH ₃ CH (³ A) + H ₂ (¹ Σ _g) + CO(¹ Σ _g)		17.9	-27.3	2.6	-41.4	-29.9	-25.4
saddle points	³ TS(1)	-24.7	-23.6				-4.8
	³ TS(2)	7.7	-1.8	3.2	-7.2	18.4	22.1
	³ TS(3)	-22.9	-32.0	-29.1	-38.8	-23.2	-20.4
	³ TS(4)	-45.4	-62.6	-51.1	-68.5	-50.8	-43.5
	³ TS(5)	-7.8	-17.7	-14.3	-24.7	10.8	16.9
	³ TS(6)	-84.7	-90.2	-104.1	-09.6	-81.6	-87.7

TABLE 2: Singlet Surface Ab Initio Energies (kJ mol⁻¹, 0 K) Relative to O(³P) + CH₃CHCH₂ (¹A), Computed Using the B3LYP (Two Different Basis Sets), CBS-QB3, and G3 Methods; For the Blank Entries, the Structure and Energy Could Not Be Determined

species		B3LYP				CBS-QB3	G3
		6-31+(d,p)		6-311 ++ (3df,2p)			
		w/o ZPE	w/ ZPE	w/o ZPE	w/ ZPE		
CH ₂ CH ₂ (¹ A _g) + H ₂ CO (¹ A')		-317.1	-322.0	-330.9	-336.1	-333.3	-328.3
CH ₃ CH(O)CH ₂ (¹ A)	¹ INT(1)	-357.1	-341.8	-362.6	-347.3	-365.0	-358.1
CH ₃ CH ₂ CHO (¹ A')	¹ INT(2)	-450.6	-438.1	-456.4	-444.3	-454.1	-449.6
CH ₃ CHCHOH (¹ A)	¹ INT(3)	-410.1	-397.2	-421.9	-408.6	-419.2	-415.0
CH ₃ CCH + H ₂ O (¹ A ₁)		-291.0	-298.2	-314.2	-320.8	-329.1	-326.1
CH ₂ CH ₂ (¹ A _g) + H ₂ (¹ Σ _g) + CO(¹ Σ _g)		-276.0	-311.1	-297.6	-331.0	-343.6	-337.9
saddle points	¹ TS(1)	-31.3	-32.9	-37.7	-39.7	-39.8	-36.6
	¹ TS(2)	-113.7	-114.1	-117.9	-118.6	-139.1	
	¹ TS(3)	-120.2		-129.4	-129.6	-124.4	-112.2
	¹ TS(4)	-163.6	-167.1	-169.3	-173.1	-173.8	-169.2
	¹ TS(5)	-69.3	-76.4	-79.7	-87.1	-85.9	-79.6
	¹ TS(6)	47.6	21.7	21.6	-4.7	11.5	10.4
	¹ TS(7)						-80.6
	¹ TS(8)						-90.3

fail to identify difficult saddle points. The G3 method uses a full MP2 step to optimize the initial geometry, with MP2 methods known for their accuracy with respect to transition-state geometries. The additional investment of the MP2 step in the G3 method enabled the saddle point to be found.

Following the energy gradient from ³TS(1) leads to the formation of the triplet biradical intermediate ³INT(2). The B3LYP/6-31+G(d,p) and IRCMax calculations of the relaxed PES for CH₃CHCH₂O redissociation to the reactants are shown in Figure 3, along with illustrations of the associated structures along the reaction coordinate. The IRCMax calculation yields an upper limit for the ³TS(1) energy using the DFT structure, while the ³INT(1) energy is calculated using the more accurate MP2 geometry. Thus, an upper limit for the local energy barrier governing the rearrangement of the weakly bound ³INT(1) intermediate is equal to the difference between the ³TS(1) and ³INT(1) energies, or 7.5 kJ mol⁻¹.

Aside from redissociation via the entrance channel, the biradical ³INT(2) has two dissociative pathways leading to products. One pathway leads to H₂CO + CH₃CH via C–C bond cleavage, while the second leads to the formation H + CH₃CHCHO through C–H bond breaking. As can be seen from Figure 1, the H + CH₃CHCHO pathway has a lower barrier; it is also doubly degenerate as there are two equivalent H atoms which can be lost. In addition, any product H₂CO could

conceivably dissociate in a secondary process to H₂ + CO. The H₂ and CO products are accessible within the energy limits of the system, but the activation energy associated with the dissociation of H₂CO is calculated to be 338.7 kJ mol⁻¹ using the G3 method, effectively eliminating this channel on the triplet surface.

In addition to the two dissociative paths from ³INT(2), there is also the possibility of a 1,2 H-atom shift to form triplet propanal, CH₃CH₂CHO, ³INT(3). However, a very large energy barrier of 122.3 kJ mol⁻¹ is calculated for this rearrangement, and it is not accessible for thermal reactant energies. For any ³INT(3) that is formed, two product channels are available, both involving C–C bond cleavage. Cleavage of the CH₃CH₂–CHO bond is the much lower energy pathway and results in the formation of HCO + CH₃CH₂. The competing dissociation, H₃C–CH₂CHO → CH₃ + CH₂CHO, exhibits an approximately three times larger activation energy.

It is known that the formyl radical HCO is a major product of the O + CH₃CHCH₂ reaction. Nevertheless, on the triplet surface, only ³INT(3) dissociates to yield HCO as a product. Given the large energy barrier to formation of ³INT(3), any HCO product is likely formed from singlet surface dissociation (see below).

III.A.2. Singlet Surface. The O–C₃H₆ singlet surface is also characterized by three bound intermediates, denoted ¹INT(1),

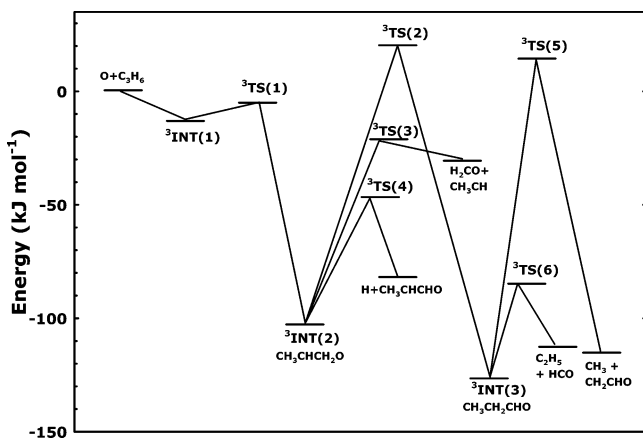


Figure 1. Schematic C₃H₆O triplet surface intermediate and transition-state energies and reaction pathways, referenced to $E = 0$ kJ mol⁻¹ for the O(³P) + C₃H₆ separated reactants. The energy values were taken as the average of the CBS-QB3 and G3 energies tabulated in Table 1.

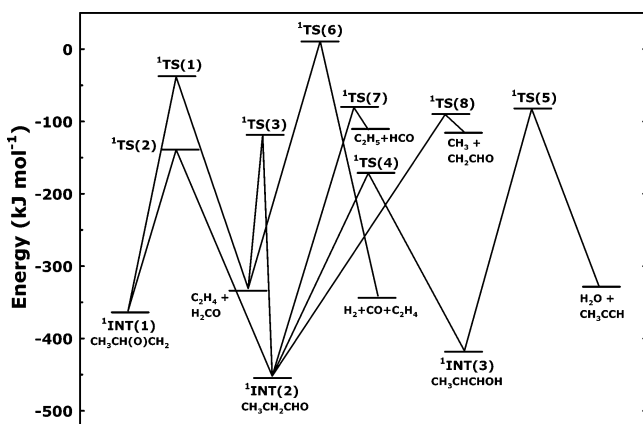


Figure 2. Schematic C₃H₆O singlet surface intermediate and transition-state energies and reaction pathways, referenced to $E = 0$ kJ mol⁻¹ for the O(³P) + C₃H₆ separated reactants. The energy values were taken as the average of the CBS-QB3 and G3 energies tabulated in Table 2.

¹INT(2), and ¹INT(3). The first intermediate, propylene oxide, ¹INT(1), has a dissociation channel to form C₂H₄ + H₂CO. Though the activation energy for further dissociation of H₂CO on the triplet surface is prohibitive, on the singlet surface, the barrier (¹TS(6)) associated with H₂CO dissociation lies only about 10 kJ mol⁻¹ above the O(³P) + C₃H₆ reactant energy. A second pathway involves rearrangement via C(O)C ring opening of ¹INT(1) to form singlet propanal CH₃CH₂CHO, ¹INT(2).

From ¹INT(2), singlet propanal, there are four dissociation/rearrangement pathways in addition to the entrance path. One of the four pathways involves tautomerization to the alcohol CH₃CHCHOH, ¹INT(3), which can subsequently dissociate to form H₂O + CH₃CCH. A second dissociative path from ¹INT(2) produces C₂H₄ + H₂CO through a 3,1 H-atom shift. This is the second route to C₂H₄ + H₂CO on the singlet surface, the first resulting from dissociation of the oxide ¹INT(1), as discussed above.

The two remaining reaction pathways from ¹INT(2) involve homolytic bond cleavage of the C–C bonds to form the radical pairs C₂H₅ + CHO and CH₃ + CH₂CHO. Accurately calculating homolytic bond cleavage on a singlet surface is often problematic. Our attempts to locate transition states for these reactions using DFT-level PES surveys were unsuccessful. However, a PES using a full unrestricted MP2/6-31G(d) calculation did pass through a maximum for both dissociation paths. Transition states were found by performing a Berny optimization, using the same

full unrestricted MP2/6-31G(d) method, for a structure near the maximum of the PES for each reaction. In both cases, a structure was obtained with a single imaginary frequency. The IRCs were followed at the MP2 level in both directions to ensure that the transition state connected the reactants and products. The transition-state structures were then used in the remaining sequence of steps to obtain G3 energies. These transition states are reported as ¹TS(7) and ¹TS(8) in Table 2 and Figure 2. The two dissociation pathways have very similar activation energies, in contrast to those for triplet propanal, ³INT(3), for which the pathways display significant differences in activation energy.

It should be noted that repeated stationary-state optimization runs using the DFT and MP2 methods starting from the molecular geometry of the triplet biradical failed to locate an analogous open-chain singlet biradical intermediate, ¹CH₃-CHCH₂O. Instead, the singlet system calculation converged on either the ¹INT(2) (aldehyde) or ¹INT(3) (enol) geometry. We could not locate a stable, open-chain singlet biradical, consistent with the computational results of Nguyen et al.³³ and other workers⁵¹ for the singlet PES of C₂H₄O.

III.B. RRKM Model. For the purposes of the RRKM calculation, a model of the triplet surface was constructed including two intermediates and five product channels (see Figure 1). The biradical CH₃CHCH₂O (³INT(2)) was assigned to the first well and triplet CH₃CH₂CHO (³INT(3)) to the second well. Four channels arise from the biradical ³INT(2), (1) redissociation to the reactants O + C₃H₆, (2) a 1,2 H-atom shift to form ³INT(3), and dissociation to yield either (3) H + CH₃-CHCHO or (4) H₂CO + CH₃CH. From propanal ³INT(3), there are three reaction paths, (1) rearrangement back to ³INT(2), and dissociation to yield (2) CH₃ + CH₂CHO or (3) C₂H₅ + HCO.

Similarly, a model of the singlet surface was constructed with three intermediates and four product channels (Figure 2). ¹INT(1) was assigned to the first well, ¹INT(2) to the second, and ¹INT(3) to the third. From the first well, two reactions are possible, dissociation to yield C₂H₄ + H₂CO and rearrangement to form ¹INT(2). ¹INT(2) can (1) rearrange via a 2,1 H-atom shift and ring closure to form ¹INT(1), (2) tautomerize to the enol isomer CH₃CHCHOH, ¹INT(3), (3) undergo a 3,1 H-atom shift and C–C bond scission to form C₂H₄ + H₂CO, or (4) dissociate along one of its C–C bonds to produce the radical pairs C₂H₅ + HCO or CH₃ + CH₂CHO. Of the reactions described above, the dissociation pathways producing the HCO and CH₂CHO radicals are thought to be dominant.⁴ Secondary reactions involving the dissociation of HCO to H + CO⁵² and the dissociation of the vinyloxy radical CH₂CHO to several different products have also been studied.^{5,53} From the third well ¹INT(3), in addition to the entrance channel, there is one product channel to form CH₃CCH + H₂O. The model was run under the same conditions (pressure, temperature, and collisional relaxation parameters) as those described above for the triplet surface, with two exceptions, (1) the reactants were taken to have a “thermal” energy distribution rather than a “chemically activated” distribution because we had no activating channel to reference with the same O(³P) + C₃H₆ energy zero, and (2) the G3 energy results were used directly.

On both surfaces, rearrangements between intermediates were allowed to be reversible, while dissociative reactions were assumed to be irreversible.

III.B.1. Branching Ratios on the Triplet Surface. The results of the RRKM calculation for the triplet surface model appear in Table 3 for the 300–3000 K temperature range and high (760 Torr) and low (0.001 Torr) pressures. Pressure was found to have little influence over the branching ratios,

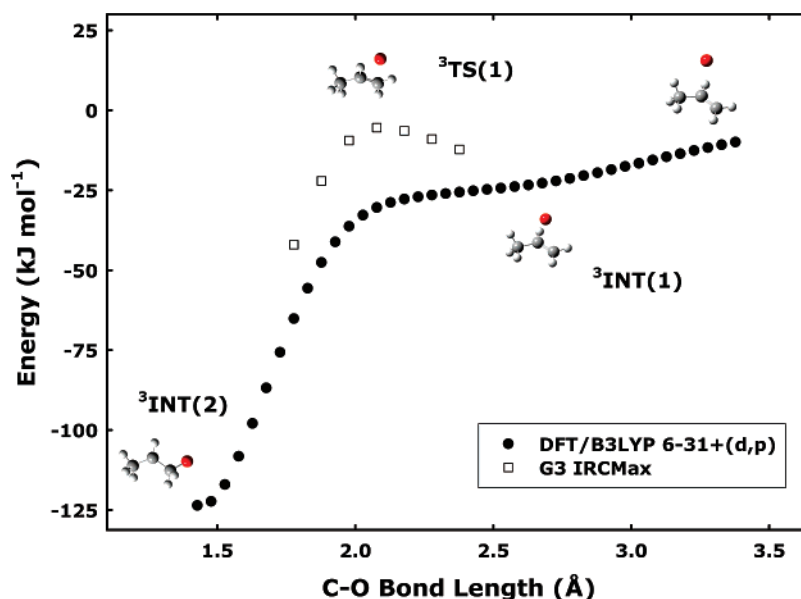


Figure 3. Potential energy profile for redissociation of triplet $\text{CH}_3\text{CHCH}_2\text{O}$ to $\text{O}(^3\text{P}) + \text{C}_3\text{H}_6$, with the other degrees of freedom relaxed as the terminal C–O bond length is increased. Calculations were performed at the B3LYP level with a 6-31+(d,p) basis set and by the G3 IRCMax method for geometries near the transition state $^3\text{TS}(1)$. The energies are relative to the energy of the separated $\text{O}(^3\text{P}) + \text{C}_3\text{H}_6$ reactants.

TABLE 3: Percent Product Distribution for Triplet $\text{CH}_3\text{CHCH}_2\text{O}$ Dissociation Channels with Yield $\geq 0.5\%$, Computed at Two Different Pressures Using the RRKM Model Described in the Text

<i>T</i> (K)	760 Torr				0.001 Torr			
	H + CH_3CHCHO	$\text{H}_2\text{CO} +$ CH_3CH	O + CH_3CHCH_2	$\text{C}_2\text{H}_5 +$ HCO	H + CH_3CHCHO	$\text{H}_2\text{CO} +$ CH_3CH	O + CH_3CHCH_2	$\text{C}_2\text{H}_5 +$ HCO
300	95.3	4.7	0.0	0.0	94.7	5.3	0.0	0.0
600	89.8	10.2	0.0	0.0	89.2	10.7	0.2	0.0
900	84.3	15.4	0.3	0.0	84.0	15.6	0.3	0.1
1200	79.3	19.9	0.7	0.2	79.1	20.1	0.7	0.1
1500	75.8	22.9	1.0	0.3	75.1	23.6	1.0	0.3
1800	72.8	25.3	1.4	0.5	73.3	24.9	1.4	0.5
2100	70.9	26.6	1.6	0.8	71.3	26.0	1.8	0.8
2400	69.0	27.6	2.1	1.2	69.6	26.9	2.4	1.1
2700	68.3	27.4	2.4	1.7	68.3	27.6	2.5	1.5
3000	66.9	27.7	3.0	2.1	66.9	28.1	3.1	1.9

TABLE 4: Percent Product Distribution for Singlet $\text{CH}_3\text{CH}_2\text{CHO}$ Dissociation Channels, Computed at Three Different Pressures Using the RRKM Model Described in the Text; The Three Principal Product Channels Are Shown

<i>T</i> (K)	760 Torr ^a			0.5 Torr			0.001 Torr		
	$\text{H}_2\text{CO} +$ C_2H_4	$\text{C}_2\text{H}_5 +$ HCO	$\text{CH}_3 +$ CH_2CHO	$\text{H}_2\text{CO} +$ C_2H_4	$\text{C}_2\text{H}_5 +$ HCO	$\text{CH}_3 +$ CH_2CHO	$\text{H}_2\text{CO} +$ C_2H_4	$\text{C}_2\text{H}_5 +$ HCO	$\text{CH}_3 +$ CH_2CHO
300	36.9	6.5	8.3	56.5	20.7	22.8	56.4	19.8	23.8
600	35.1	27.8	28.4	32.7	33.4	33.9	31.5	34.4	34.2
900	15.3	43.6	40.3	13.8	45.4	40.8	13.8	44.2	42.0
1200	4.7	52.5	42.7	4.8	52.3	42.9	5.0	51.9	43.0
1500	1.9	54.1	44.0	2.1	55.1	42.8	2.0	54.4	43.6
1800	0.8	55.3	43.8	0.7	56.0	43.2	0.8	56.1	43.1
2100	0.7	56.6	42.7	0.5	56.5	43.0	0.4	56.3	43.2
2400	0.6	56.7	42.7	0.6	57.7	41.7	0.4	56.7	42.8
2700	0.6	57.0	42.3	0.6	57.7	41.7	0.7	56.9	42.5
3000	1.0	56.6	42.4	1.0	57.2	41.7	1.1	56.2	42.7

^a At atmospheric pressure, 48.3, 8.8, and 0.8% of the energized $\text{CH}_3\text{CH}_2\text{CHO}$ intermediate is stabilized prior to dissociation at temperatures of 300, 600, and 900 K, respectively.

suggesting that the vibrational self-relaxation rates for $p \leq 1$ atm are negligible compared to the triplet intermediate inverse lifetimes.

For reactant internal energies greater than 20.2 kJ mol^{-1} , there is sufficient energy to overcome the barrier to convert $^3\text{INT}(2)$ into $^3\text{INT}(3)$ by the 1,2 H-atom shift. Yet, even at the highest temperature of 3000 K, with equipartition suggesting that each vibrational mode has up to 24.9 kJ mol^{-1} of energy, products resulting from the dissociation of $^3\text{INT}(3)$ are very minor, about

2.5%. It appears that nearly all of the dissociation products from the triplet surface result from the dissociation of the biradical $^3\text{INT}(2)$. Nguyen et al. obtained similar results for the $\text{O} + \text{C}_2\text{H}_4$ system.³³

For the reactions of the triplet biradical, $\text{CH}_3\text{CHCH}_2\text{O}$ ($^3\text{INT}(2)$), the activation barrier is lowest for H-atom loss to form $\text{H} + \text{CH}_3\text{CHCHO}$ and highest for the 1,2 H-atom shift to form triplet $\text{CH}_3\text{CH}_2\text{CHO}$ ($^3\text{INT}(3)$). The curvature of the PES in the vicinity of the transition state also strongly influences

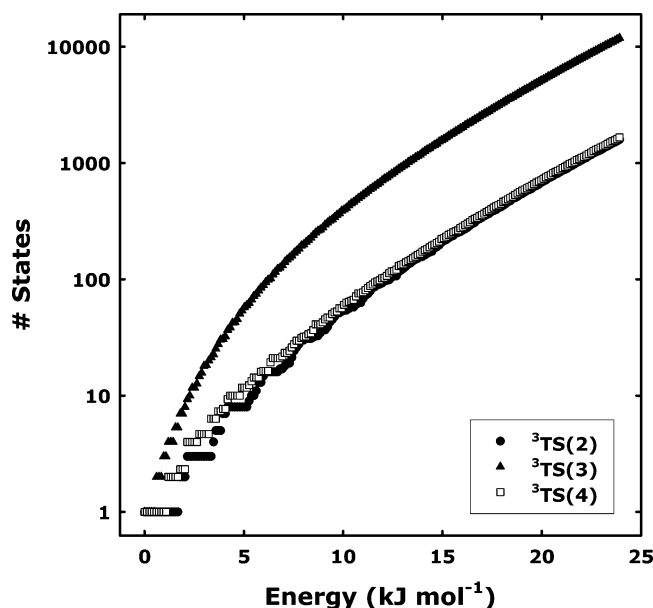


Figure 4. Semilog plot for three triplet transition-state sums as a function of energy above threshold, governing the dissociation of the triplet biradical CH₃CHCH₂O, ³INT(2). The larger ³TS(3) state sums lead to an enhancement of the H₂CO + CH₃CH channel as the energy increases. See Figure 1 for transition-state definitions.

the reaction rate. Given that the reaction rate is proportional to the transition-state sum of states, a faster rate is predicted for a PES with less curvature and a larger internal state sum. As an indication of this effect, Figure 4 shows the computed sums of states for the ³TS(2), ³TS(3), and ³TS(4) transition states from threshold to 23.9 kJ mol⁻¹ (2000 cm⁻¹) above the respective transition-state energy. It can be seen that the ³TS(3) state possesses about six times as many internal states as ³TS(2) or ³TS(4) at any given excess energy. The H + CH₃CHCHO product channel dominates at low temperature via the low threshold energy transition state ³TS(4). However, the higher-energy path producing CH₃CH + H₂CO becomes dominant as the temperature (or energy) increases, and the ³TS(3) internal state sum rapidly becomes greater than that for ³TS(4). (Consider as an analogy water held back by a dam; the water level connotes energy, while the dam connotes an energy barrier. For low levels, the water leaks through a small hole in the dam—analogue to the formation of H + CH₃CHCHO—while for high levels, the water flows much faster over the top of the dam—analogue to the formation of H₂CO + CH₃CH.)

The RRKM analysis shows that rearrangement of ³INT(2) to ³INT(3) on the triplet surface is not favored even in the presence of significant excess energy. However, intersystem crossing of ³INT(2) from the triplet surface to the singlet surface may occur. Our ab initio work has shown that for singlet CH₃CHCH₂O, the 1,2 H-atom shift exhibits a small or no barrier, similar to others³³ findings for singlet CH₂CH₂O. (Recall the discussion above that the calculations could not identify a bound CH₃CHCH₂O singlet intermediate.) Intersystem crossing of ³INT(2) to the singlet surface would then result in the formation of singlet CH₃CH₂CHO, ¹INT(2), which can undergo several different rearrangements or directly dissociate as described below. On the triplet surface, HCO can only be formed through dissociation of ³INT(3), which according to our RRKM analysis is a minor channel at best. Instead, any HCO product is likely formed from dissociation of ¹INT(2) following intersystem crossing.

III.B.2. Branching Ratios on the Singlet Surface. Table 4 lists the singlet surface product branching ratios for the same

TABLE 5: Percent Product Distribution for C₃H₆O Triplet and Singlet Dissociation Channels for Two Temperatures, Computed at 0.001 Torr Using the RRKM Model Described in the Text; The Principal Product Channels Are Shown

T (K)	triplet surface			singlet surface		
	H + CH ₃ CHCHO	H ₂ CO + CH ₃ CH	C ₂ H ₅ + HCO	H ₂ CO + C ₂ H ₄	C ₂ H ₅ + HCO	CH ₃ + CH ₂ CHO
300	95	5	0	57	20	23
≥2000	70	28	2	1	56	43

300–3000 K temperature range and for 760, 0.5, and 0.001 Torr pressures. The low-temperature dissociation is dominated by the formation of C₂H₄ + H₂CO, with the formation of the C₂H₅/HCO and CH₃/CH₂CHO radical pairs quickly becoming dominant as the temperature increases. The H₂O + CH₃CCH product channel arising from the enol CH₃CHCHOH (¹INT(3)) intermediate does not occur at any temperature. From Figure 2, it would seem reasonable to assume that the C₂H₄ + H₂CO and H₂O + CH₃CCH product channels would be at least as probable as the C₂H₅ + HCO and CH₃ + CH₂CHO channels at high temperature based on energy barrier considerations. However, the corresponding transition-state sums exhibit a two to three order-of-magnitude advantage for the C–C bond dissociation pathways compared to those of the other pathways. Thus, the alkyl–aldehyde radical pairs dominate the high-temperature product yield for the same reasons as those described in section III.B.1 for the dissociation of ³INT(2). Note that for the lower temperatures (≤600 K) and atmospheric pressure (760 Torr), much of the energized CH₃CH₂CHO intermediate population is stabilized by collisions prior to rearrangement or dissociation. With the 760 Torr collision rate calculated as 1.46 × 10¹⁰ s⁻¹ and half of the CH₃CH₂CHO stabilized, the 300 K chemically activated lifetime is estimated to be on the order of 100 ps or larger.

III.C. Summary of RRKM Branching Ratios. Table 5 summarizes the major product distribution from both surfaces at low (300 K) and high (≥2000 K) temperatures. For the triplet surface, the H + CH₃CHCHO product channel dominates, with a significant contribution from the H₂CO + CH₃CH channel at high temperature. For the singlet surface, H₂CO + C₂H₄ is the principal product channel at low temperature, with the C₂H₅/HCO and CH₃/CH₂CHO radical pairs the nearly exclusive products at high temperature. Formaldehyde (H₂CO) may arise from either the singlet or triplet surfaces, though it is likely to be produced from the singlet surface at low temperature and from the triplet surface at high temperature. The presence of CH₃CH or C₂H₄ coproduct could conceivably be used as an indication of the source of H₂CO, CH₃CH indicating triplet surface dissociation and C₂H₄ indicating singlet surface dissociation.

IV. Summary

The lowest-lying triplet and singlet potential energy surfaces for the O + C₃H₆ reaction have been characterized using the DFT and ab initio CBS-QB3 and G3 theoretical methods. With the intermediate and transition-state energies, geometries, and vibrational frequencies as inputs, RRKM statistical theory was then used to predict product branching fractions as a function of reactant temperature and pressure. The potential energy surface for the O + C₃H₆ reaction is similar to that reported by Nguyen et al. for the O + C₂H₄ system³³ but for the additional features associated with H-atom transfers and C–C bond cleavage involving the extra C₃H₆ carbon atom. In both systems, products arise from reactions of the initially formed CH₃CHCH₂O or CH₂CH₂O biradical. Rearrangement of the nascent

triplet biradical $\text{CH}_3\text{CHCH}_2\text{O}$ to $\text{CH}_3\text{CH}_2\text{CHO}$ (propanal) is hindered both by a large activation energy barrier and by a low sum of states for the transition state. The triplet system yields mostly $\text{H} + \text{CH}_3\text{CHCHO}$ products at all temperatures. The nascent singlet biradical $\text{CH}_3\text{CHCH}_2\text{O}$, accessed through inter-system crossing, quickly rearranges to singlet propanal. Subsequent singlet surface dissociation yields mostly $\text{C}_2\text{H}_4 + \text{H}_2\text{CO}$ at low temperature and a roughly 1:1 proportion of $\text{C}_2\text{H}_5 + \text{HCO}$ and $\text{CH}_3 + \text{CH}_2\text{CHO}$ at high temperature. Aside from evidence that singlet $\text{CH}_3\text{CH}_2\text{CHO}$ is stabilized at 1 atm at low temperature, pressure was found to have little influence over the branching ratios, suggesting that the vibrational self-relaxation rates for $p \leq 1$ atm are negligible compared to the dissociation rates.

Acknowledgment. This research was supported by the Air Force Office of Scientific Research under Task 2303ES/92VS04COR. G.D.B. gratefully acknowledges an AFOSR 2006 Summer Faculty Fellowship and the Welch Foundation for its support of the LeTourneau University chemistry department under Award BL-0028.

Supporting Information Available: Calculated intermediate and transition-state geometries. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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