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# Photodissociation spectroscopy and dynamics of CH<sub>3</sub>O and CD<sub>3</sub>O

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The photodissociation spectroscopy and dynamics of the methoxy radical  $\tilde{A}(^2A_1) \leftarrow \tilde{X}(^2E)$  transition have been investigated using fast radical beam photofragment translational spectroscopy. The  $\tilde{A}$  state of both  $CH_3O$  and  $CD_3O$  is observed to predissociate via curve crossings with one or more repulsive electronic states. The photofragment yield spectrum consists of the C-O stretch progression and combination bands based on this mode. The major product channel is  $CH_3$  ( $CD_3$ ) + O, with a threshold 3775 cm<sup>-1</sup> above the zero-point level of the  $\tilde{A}$  state. The product translational energy distributions reveal  $\nu_2$  umbrella excitation in  $CH_3$  ( $CD_3$ ), yielding considerable insight into the dissociation dynamics for this channel.  $CD_3O$  shows a mode-specific dynamical effect in which energy deposited in parent umbrella motion preferentially populates umbrella motion in the fragment. The product channels  $CH_2 + OH$  ( $CD_2 + OD$ ) and  $D + CD_2O$  are also observed, with evidence that the methylene fragment is predominantly in the excited  $\tilde{a}(^1A_1)$  state.

#### I. Introduction

The methoxy radical (CH<sub>3</sub>O) is a remarkable molecule and has become one of the most widely studied polyatomic free radicals since it was first observed in emission of ethyl nitrate photolysis in 1953.<sup>1</sup> Methoxy has drawn such intense scrutiny because it serves as a paradigm for a surprisingly large number of fundamental physical and chemical interactions. It is important in the atmosphere due to its participation in smog chemistry<sup>2</sup> and is an intermediate in the combustion of cool flames of methane,<sup>3</sup> oxygenated fuels containing CH<sub>3</sub>OH, and fuel-lean ignition processes.<sup>4</sup> Although not yet detected in interstellar gas, it is quite likely that CH<sub>3</sub>O exists in this medium because its close relatives CH<sub>3</sub>OH and CH<sub>2</sub>O are found in abundance.<sup>5</sup>

In the field of photodissociation dynamics, the rich photochemistry of methoxy provides archetypal cases of two distinct dissociation mechanisms. On the ground state surface, dissociation to H + CH<sub>2</sub>O is impeded by an activation barrier. Stimulated emission pumping (SEP) experiments show narrow resonances both below and above the barrier, addressing the long-standing question of mode-specific vs statistical unimolecular decomposition.<sup>6</sup> In a preliminary report,<sup>7</sup> we presented the first unambiguous evidence for predissociation of CH<sub>3</sub>O following excitation of the ultraviolet  $\tilde{A}(^2A_1) \leftarrow \tilde{X}(^2E)$  transition. In contrast with barrier-impeded ground state dissociation, UV dissociation of methoxy, yielding CH<sub>3</sub> + O, provides a textbook example of predissociation via coupling to purely repulsive excited state potential energy surfaces. In this paper we give a full exposition covering our experiments on the ultraviolet photodissociation dynamics of CH<sub>3</sub>O and CD<sub>3</sub>O.

Although the literature on CH<sub>3</sub>O is quite extensive, the most germane works from the viewpoint of photodissociation dynamics are the spectroscopic<sup>6–31</sup> and theoretical investigations<sup>32–40</sup> of the rotational, vibrational, and electronic structure of methoxy. Within the  $C_{3\nu}$  point group, the CH<sub>3</sub>O ground state is of E symmetry due to a 2-fold electronic degeneracy. Consequently, CH<sub>3</sub>O is subject to vibronic coupling via the Jahn–Teller effect<sup>41,42</sup> in which nuclear motion along one (or more) of the e symmetry vibrations splits the electronic degeneracy. In

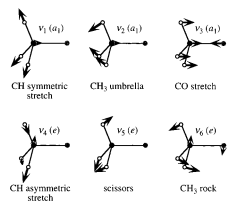


Figure 1. Vibrational normal modes of CH<sub>3</sub>O.

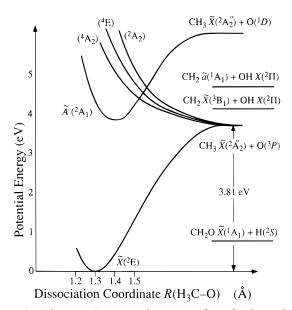
addition, the unpaired electron spin can couple to the magnetic field generated by the electronic orbital motion around the symmetry axis. As a result, methoxy is a prototypical example of the complications that arise when both Jahn—Teller and spin—orbit effects must be treated together.<sup>43</sup>

The spectroscopy of CH<sub>3</sub>O is far from straightforward, with a substantial amount of controversy surrounding the proper values of the spectroscopic parameters. As noted by Foster *et al.*,<sup>23</sup> there has been a great deal of confusion in determining the true location of the  $0_0^0$  band in the  $\tilde{A}(^2A_1) \leftarrow \tilde{X}(^2E)$  transition. These difficulties arise in part from the fact that all three non-totally symmetric (e) vibrations ( $\nu_4$ ,  $\nu_5$ , and  $\nu_6$ ) are active in the  $\tilde{A}$  state, in addition to the three totally symmetric (a<sub>1</sub>) vibrations ( $\nu_1$ ,  $\nu_2$ , and  $\nu_3$ ); the normal modes for the six vibrations are shown in Figure 1. The resulting spectra are rather complex, even at the very low rotational and vibrational temperature achieved in a supersonic jet expansion. Extensive vibrational<sup>30</sup> and rotational<sup>25</sup>, analyses of the  $\tilde{A} \leftarrow \tilde{X}$  laserinduced fluorescence (LIF) spectrum give the accepted value of the rovibronic band origin  $T_0 = 31\ 614.51\ \text{cm}^{-1}$ .

All experimental and theoretical investigations agree that the main progression in the  $\tilde{A} \leftarrow \tilde{X}$  transition is the 662 cm<sup>-1</sup>  $\nu_3$  mode, the C–O stretch. The extended Franck–Condon progression in this mode arises from the large change in C–O bond length between ground and excited states ( $r_{\rm CO}' = 1.58~{\rm \AA} \leftarrow r_{\rm CO}'' = 1.37~{\rm \AA}$ ).<sup>25</sup> The increase in  $r_{\rm CO}$  results from excitation of a CO  $\sigma$ -bonding electron to a nonbonding  $p_{\pi}$  orbital localized

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**Figure 2.** Schematic  $C_{3v}$  potential energy surfaces for the methoxy radical as a function of CO bond length.

on the oxygen atom, leaving essentially half a CO bond in the  $\tilde{A}(^2A_1)$  state. However, due to the complexity of the spectrum, the vibrational frequencies for several of the other modes have proved difficult to evaluate, although two recent investigations appear to have settled all the major discrepancies.<sup>30,31</sup> There has also been considerable variation in the reported values of the  $\tilde{A}$  state fluorescence lifetime.<sup>12,17–19,22</sup> The most recent measurements (obtained under collisionless conditions) for the  $\nu_3$  progression give  $\tau_{\rm FL}(3^n_0, n < 6) \approx 2~\mu{\rm s},~\tau_{\rm FL}(3^6_0) = 0.35-0.38~\mu{\rm s},~{\rm and}~\tau_{\rm FL}(3^7_0) < 0.02~\mu{\rm s}.^{44}$ 

Jackels performed the first systematic theoretical investigation of the many excited electronic states of  $CH_3O.^{33}$  In addition to the bound  $\tilde{X}(^2E)$  and  $\tilde{A}(^2A_1)$  states, he reported three electronic states of  $^4A_2$ ,  $^4E$ , and  $^2A_2$  symmetry which are purely repulsive along the C–O coordinate, shown schematically in Figure 2. The curve crossing between the lowest of these states and the  $\tilde{A}$  state was predicted to lie at 36 500 cm $^{-1}$ , slightly above the energy of the  $3_0^7$  transition. More detailed *ab initio* investigations, including the coupling strengths between different excited electronic states, have recently been undertaken by Pederson and Yarkony $^{39}$  and by Cui and Morokuma. $^{40}$ 

There are at least four photodissociation product channels open to methoxy at the excitation energies used in this study:

$$CH_{3}O(\tilde{X}(^{2}E)) \xrightarrow{h\nu} CH_{3}O(\tilde{A}(^{2}A_{1})) \rightarrow$$

$$CH_{3}(\tilde{X}(^{2}A_{2}'')) + O(^{3}P) \qquad \Delta H_{rxn,0} = 3.807 \pm 0.013 \text{ eV}$$

$$(I)$$

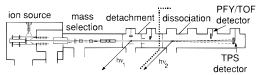
$$CH_{2}(\tilde{X}(^{3}B_{1})) + OH(\tilde{X}(^{2}\Pi)) \qquad \Delta H_{rxn,0} = 4.10 \pm 0.05 \text{ eV}$$

$$(II)$$

$$\text{CH}_2(\tilde{a}(^1\text{A}_1)) + \text{OH}(\tilde{X}(^2\Pi))$$
  $\Delta H_{\text{rxn},0} = 4.49 \pm 0.05 \text{ eV}$  (III)

$$CH_2O(\tilde{X}(^1A_1)) + H(^2S)$$
  $\Delta H_{rxn,0} = 0.78 \pm 0.07 \text{ eV}$  (IV)

The energetics are based on the heat of formation of methoxy  $\Delta H^{\circ}_{f,0}(\text{CH}_3\text{O}) = 0.29 \pm 0.02 \text{ eV}$  (6.8  $\pm$  0.4 kcal/mol) found in our preliminary investigation<sup>7</sup> and the literature values for other species.<sup>45</sup> Channel I is the major product channel for CH<sub>3</sub>O at



**Figure 3.** The fast radical beam translational spectrometer. The dotted line separates the radical production section on the left from the photodissociation section on the right. The flight distance L between the dissociation laser and a detector is 0.68 m for the TOF detector and either 1.00 or 2.00 meters for the TPS detector.

all photon energies. The formation of CH<sub>2</sub> + OH via channels II and/or III is observed as a minor product channel in CH<sub>3</sub>O photodissociation. The corresponding channels are also observed with CD<sub>3</sub>O. As discussed elsewhere,<sup>46</sup> detection of H atom loss is difficult though not impossible with our apparatus. For methoxy we have evidence that channel IV plays a very minor role in methoxy dissociation at high photon energies.

Our experimental results for both isotopes are presented in section III and analyzed in section IV. In section V we discuss the dissociation dynamics of methoxy in light of the experimental distributions and comment on the possibilities for further investigations.

#### **II. Experimental Section**

The experimental apparatus used in these studies, the fast beam photofragment translational spectrometer (Figure 3), has been described in detail elsewhere, <sup>47,48</sup> and only a brief description is given here. In order to produce a well-characterized sample of free radicals, we rely on the fact that open-shell species have positive electron affinities and form stable negative ions. The neutral free radical of interest is generated by laser photodetachment of the mass-selected negative ion precursor and subsequently photodissociated by a second laser:

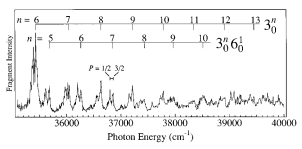
$$CH_3O^{-} \xrightarrow{h\nu_1} CH_3O + e^{-} \xrightarrow{h\nu_2} fragments$$
 (1)

Consequently, the apparatus in Figure 3 is divided into two main regions. In the first region ions are generated and photodetached, while radical photodissociation occurs in the second region.

Methoxide ions are formed in a 60 Hz pulsed supersonic expansion of CH<sub>3</sub>OH seeded in 5 atm Ne at room temperature. A pulsed electric discharge in the throat of the expansion creates negative ions,<sup>49</sup> which cool both rotationally and vibrationally during the expansion. The ions are accelerated to 8000 eV laboratory energy and mass selected by the Bakker time-of-flight (TOF) method.<sup>50</sup> The output of a pulsed dye laser intersects the ion beam at the appropriate time such that only methoxide ions are photodetached. Furthermore, the detachment energy is just above threshold<sup>9</sup> ( $h\nu = 1.71$  eV for CH<sub>3</sub>O,  $h\nu = 1.66$  eV for CD<sub>3</sub>O), so that methoxy is created only in its ground vibrational state.

In the second region of the apparatus, a frequency-doubled pulsed dye laser, operating between 35 000–40 500 cm<sup>-1</sup> with a bandwidth of 0.3 cm<sup>-1</sup>, intersects the packet of methoxy radicals. The dissociation laser is calibrated against the absorption spectrum of I<sub>2</sub>,<sup>51</sup> with an absolute accuracy of 1 cm<sup>-1</sup>. Fragments from a photodissociation event are detected directly, without an ionization step, using one of two microchannel plate (MCP) detectors. An aluminum strip across the center of each detector prevents undissociated radicals from striking the MCPs, so that all signal observed is due to recoiling photofragments.

Three types of experiments are performed to characterize the photodissociation of methoxy. First, photofragment yield (PFY) spectra are obtained by monitoring the total flux of fragments



**Figure 4.** Photofragment yield spectrum of CH<sub>3</sub>O. The 60 cm<sup>-1</sup> spin-orbit splitting between the  ${}^{2}E_{P}$ ,  $P = {}^{3}/{}_{2}$ ,  ${}^{1}/{}_{2}$  levels of the  $\tilde{X}$  state is shown.

at the TOF detector, located 0.68 m from the dissociation laser, as a function of laser wavelength. The resulting spectrum is complementary to absorption and fluorescence measurements.

Second, after the PFY spectrum is acquired, the photodissociation laser is fixed at a specific energy, and both fragments arising from a single parent radical are detected in coincidence using a time- and position-sensitive (TPS) wedge- and stripanode detector,  $^{52,53}$  located 2.0 m from the dissociation laser. Due to the favorable kinematics of this fast beam experiment, the data may be directly inverted to produce the two-dimensional-coupled translational energy and angular distribution  $P(E_{\rm T},\theta)$  where  $E_{\rm T}$  is the center-of-mass relative translational energy of the recoiling fragments and  $\theta$  is their scattering angle with respect to the electric field of the linearly polarized dissociation laser. This distribution can be separated according to

$$P(E_{\mathrm{T}},\theta) = P(E_{\mathrm{T}})\{1 + \beta(E_{\mathrm{T}})P_{2}(\cos\theta)\}$$
 (2)

into the angle-independent translational energy distribution  $P(E_T)$  and the energy-dependent anisotropy parameter<sup>54</sup>  $\beta(E_T)$ , which describes the angular distribution of fragments. The high laboratory kinetic energy of the fragments affords an MCP detection efficiency of  $\approx 50\%$ , which is independent of the identity of the fragment (except for H or D atoms, as noted below). For the experiments presented here, the translational energy resolution is given by  $\Delta E_T/E_T = 2.2\%$ . This coincidence detection scheme is only feasible when the mass ratio of the two fragments  $m_1/m_2 \leq 5$ .

Because of this restriction, detection of channel IV requires a different approach. The third mode of operation is a noncoincidence experiment, in which the time-of-flight of the photofragments is recorded by digitizing the signal from the TOF detector in 1 ns time bins.<sup>46</sup> The observable is a projection of the 3-D velocity distribution onto the radical beam axis, averaging out much of the detail which is present in the full  $P(E_{\rm T},\theta)$  distribution. In exchange for this sacrifice of detail, hydrogen atom loss channels are particularly distinct because of the large spread in flight times for H (or D) atoms scattered parallel and antiparallel to the ion beam direction. Given sufficient data, a Monte Carlo forward convolution routine can be implemented to determine translational energy and angular distributions which are consistent with the TOF data. In practice, the experiment is much more sensitive to D atoms than H atoms, and this experiment has only been performed for CD<sub>3</sub>O. Even for D atoms, the detection efficiency of  $\approx$ 8% is significantly reduced from that of "heavy" fragments such as  $CH_3$ .

# III. Results

A. Photofragment Yield Spectra of CH<sub>3</sub>O and CD<sub>3</sub>O. The PFY spectrum shown in Figure 4 (first reported in ref 7) is a direct observation of photodissociation in the  $\tilde{A}(^2A_1) \leftarrow \tilde{X}$ -( $^2E$ ) transition. The spectrum is composed of two prominent

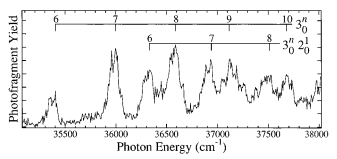


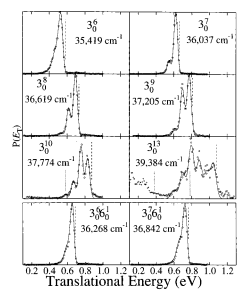
Figure 5. Photofragment yield spectrum of CD<sub>3</sub>O.

vibrational progressions superimposed on a small but finite continuum of photofragment signal. No signal was observed below 35 300 cm $^{-1}$ . Rotational resolution was not a priority in this study, and although it has been resolved, such spectra are very congested, in part due to the estimated rotational temperatures of 35–50 K in the radical beam.<sup>52,55</sup> Therefore, a relatively large laser step size of  $\approx 6$  cm $^{-1}$  was adopted to allow increased signal averaging at each photon energy. However, because of the underlying rotational structure, each individual scan was offset slightly in energy from the previous data in a given scan range in order to average over rotational features, which might otherwise provide spurious structure since the laser step size is larger than the laser bandwidth (0.3 cm $^{-1}$ ).

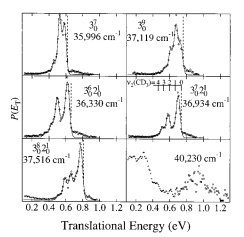
The two progressions in Figure 4 have a spacing of  $\approx 600$ cm<sup>-1</sup>, implicating progressions involving  $\nu_3$ , the CO stretch (see Figure 1). The CO stretching mode also dominates the absorption and LIF spectra, where it has a fundamental frequency of  $v_3 = 662 \text{ cm}^{-1.30}$  The two progressions we observe are assigned as the pure CO stretch,  $3_0^n$  (n = 6-13), and a combination band  $3_0^n 6_0^1$  (n = 5-10), which involves one quantum of the nontotally symmetric methyl rock vibration. The assignment of the combination band differs from that presented in our preliminary investigation<sup>7</sup> and is discussed in section IV.A. On a finer scale, each peak shows a splitting of approximately 60 cm<sup>-1</sup> (see Figure 4), corresponding to the spin-orbit splitting between the  ${}^{2}E_{3/2}$  and  ${}^{2}E_{1/2}$  components of the ground electronic state. Both spin components are present because each represents an allowed transition in the photodetachment of CH<sub>3</sub>O<sup>-</sup>(<sup>1</sup>A<sub>1</sub>). However, the "hot" spin-orbit distribution does not imply a hot vibrational distribution. In fact, the photoelectron spectrum of CH<sub>3</sub>O<sup>-</sup> (acquired on a different apparatus but with the same ion source) shows no hot bands, implying that all the ions are in their vibrational ground state.<sup>56</sup> To aid in comparison with LIF spectra<sup>13,16,23–28,30</sup> of jet-cooled methoxy, which probed only transitions from the lower <sup>2</sup>E<sub>3/2</sub> component of the ground state, the vibrational combs in Figure 4 are centered on those transitions originating in the <sup>2</sup>E<sub>3/2</sub> component.

Figure 5 shows the corresponding PFY spectrum for the  $CD_3O$  radical. The general features of the spectrum are analogous to  $CH_3O$ ,  $3_0^n$  is the main progression and is accompanied by a combination band. However, the continuum signal underneath the vibrational structure is more intense at lower energies for  $CD_3O$  than for  $CH_3O$ . In addition, the combination band observed for  $CD_3O$  is assigned to a different progression involving umbrella motion,  $3_0^n 2_0^1$ , as discussed in section IV.A.

**B.** Translational Energy and Angular Distributions. The photofragment translational energy distributions  $P(E_T)$  arising from CH<sub>3</sub>O and CD<sub>3</sub>O, are given in Figures 6 and 7 for excitation of several vibrational levels shown in the PFY spectra of Figures 4 and 5, respectively. The translational energy is binned in intervals of 10 meV. For photon energies  $h\nu < 37\,500$ 



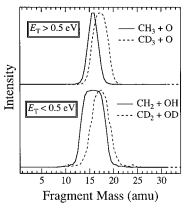
**Figure 6.**  $P(E_T)$  distributions for CH<sub>3</sub>O. Circles show CH<sub>3</sub> + O data, crosses show CH<sub>2</sub> + OH data, while the line represents the fit to CH<sub>3</sub> + O discussed in the text. Thermodynamic thresholds are given for CH<sub>3</sub> + O (--), CH<sub>2</sub>  $\tilde{X}(^3B_1)$  + OH ( $\cdots$ ), and CH<sub>2</sub>  $\tilde{\alpha}(^1A_1)$  + OH ( $\cdot-$ ).



**Figure 7.**  $P(E_T)$  distributions for CD<sub>3</sub>O. The legend is identical to Figure 6.

cm<sup>-1</sup>, all signal arises from simple C–O bond fission, *i.e.*, channel I. Most of the available energy is deposited into translation of the recoiling fragments, because the distributions peak near the maximum translational energy available (see section IV.B) as denoted by the dashed vertical line in each distribution of Figures 6 and 7. The most striking feature in the CH<sub>3</sub>O  $P(E_T)$  distributions is the progression with a spacing of  $\approx$ 74 meV (600 cm<sup>-1</sup>), which is easily seen in the  $3_0^n$  data sets ( $n \ge 7$ ). As detailed in section IV.B, these features arise from excitation of the  $\nu_2$  umbrella mode of the CH<sub>3</sub> fragment. Note that the vibrational features are much less distinct in the  $P(E_T)$  distributions from the  $3_0^6$  and  $3_0^8 6_0^1$  bands.

In the CD<sub>3</sub>O  $P(E_{\rm T})$  distributions (Figure 7), most of the available energy again goes into product translation, with a product vibrational progression of  $\approx$ 57 meV (460 cm<sup>-1</sup>) observed, corresponding to excitation of the  $\nu_2$  umbrella mode in CD<sub>3</sub> photofragment. The most surprising results in Figure 7 are the product state distributions observed when the  $3_0^n 2_0^1$  combination band is excited. In section IV.B, we will show that these  $P(E_{\rm T})$  distributions are the result of a bimodal distribution in the  $\nu_2$  mode of CD<sub>3</sub>, with the two most intense peaks corresponding to  $\nu_2 = 0$  and  $\nu_2 = 2$ . Note the stark contrast between the distributions arising from  $3_{0.0}^{n-1}$  excitation of CD<sub>3</sub>O and those arising from  $3_{0.0}^{n-1}$  excitation of CH<sub>3</sub>O.



**Figure 8.** Photofragment mass spectrum exciting  $3_0^{13}$  of CH<sub>3</sub>O (—) and 40 230 cm<sup>-1</sup> of CD<sub>3</sub>O (···). Note that for  $E_T > 0.5$  eV, the deuterium isotope has the broader fwhm, implying fragmentation to channel I, while for  $E_T < 0.5$  eV the hydrogen isotope has the broader fwhm, implying fragmentation to channels II and/or III. The peak heights are normalized to unity and do not represent branching ratios.

All of the  $P(E_T)$  distributions presented arise from excitation of the  ${}^2E_{3/2}$  level of the  $\tilde{X}$  state. Excitation from the  ${}^2E_{1/2}$  levels exhibits identical behavior, which is to be expected since the level excited in the  $\tilde{A}({}^2A_1)$  state is identical. In addition,  $P(E_T)$  distributions acquired at three different photon energies within the rotational envelope of the  $3_0^8$  band of CH<sub>3</sub>O show no significant differences. Therefore, further investigations of the dependence of the photodissociation dynamics on the excited state rotational level were not pursued.

For  $hv > 37\,500$  cm<sup>-1</sup>, the structured features in the  $P(E_{\rm T})$  distributions arising from C–O bond fission remain, but a new feature is observed in both isotopes for  $E_{\rm T} < 0.5$  eV which is due to production of CH<sub>2</sub> + OH (CD<sub>2</sub> + OD). The evidence for this new channel is shown in Figure 8, which gives the product mass spectra of CH<sub>3</sub>O and CD<sub>3</sub>O for  $3_0^{13}$  and 40 230 cm<sup>-1</sup> excitation respectively, divided into contributions for  $E_{\rm T} > 0.5$  eV and  $E_{\rm T} < 0.5$  eV. The comparison between the two isotopes is useful because the product mass resolution is not sufficient to unambiguously resolve peaks arising from CH<sub>2</sub>, CH<sub>3</sub>, O, and OH. Although only a single peak appears in the mass spectra, the width of this peak depends on the product masses.

For CD<sub>3</sub>O the two fragments have masses 16 and 18 regardless of whether channel I (CD<sub>3</sub> + O) or channel II/III (CD<sub>2</sub> + OD) is produced, resulting in an identical product mass spectrum regardless of the branching ratio between these two channels. In CH<sub>3</sub>O, the analogous decay pathways have product masses 15 and 16, or 14 and 17, respectively. Therefore, formation of channel I should give rise to a narrower peak for CH<sub>3</sub>O than for CD<sub>3</sub>O, while channel II/III products will give a broader peak for CH<sub>3</sub>O than for CD<sub>3</sub>O. From these considerations, the only conclusion consistent with the results in Figure 8 is that channel I products dominate for  $E_T > 0.5$  eV, while channel II/III products dominate when  $E_T < 0.5$  eV.

The branching ratio of  $CH_3 + O:CH_2 + OH$  decreases with photon energy and is 3:1 for  $3_0^{13}$ . The same trend occurs in  $CD_3O$ , where the extreme case of excitation at  $h\nu = 40$  230 cm<sup>-1</sup> results in a branching ration  $CD_3 + O:CD_2 + OD$  of  $\approx 1:$  1, as shown in Figure 7.

In all the data sets acquired for methoxy, the anisotropy parameter  $\beta(E_{\rm T})$  is independent of  $E_{\rm T}$ . For CH<sub>3</sub>O, the  $3_0^n$  transitions show angular distributions which are isotropic to within experimental error, while the  $3_0^n 6_0^1$  transitions are slightly anisotropic, described by  $\beta = 0.4 \pm 0.1$  for  $3_0^6 6_0^1$ , and  $\beta = 0.3 \pm 0.1$  for  $3_0^7 6_0^1$ . These positive values of  $\beta$  are

unexpected for a perpendicular electronic transition and are discussed further in section V.B. In CD<sub>3</sub>O, all the transitions give isotropic angular distributions within experimental error.

C. Noncoincidence Time-of-Flight Distributions. As mentioned in the experimental section, a noncoincidence experiment must be performed in order to assess the importance of the hydrogen atom loss pathway, channel IV. In our preliminary work on methoxy, we reported a branching ratio for channel IV in CD<sub>3</sub>O of  $2\%_{-1}^{+5}$ , and  $10\%_{-5}^{+10}$  for the  $3_0^8$  and  $3_0^{10}$  bands, respectively.<sup>7</sup> The experiments were repeated recently, and the data again show evidence of D atom loss from CD<sub>3</sub>O, but with less intensity than our earlier data. On the basis of our experience with photodissociation of the vinoxy radical, in which D atom loss is a major channel, 46 it appears that the branching ratio for CD<sub>2</sub>O + D is smaller than the values reported previously, representing no more than 5% of the branching ratio at  $3_0^{10}$ . In any event, this decay channel seems to be of minor importance in methoxy, at least for the photon energy range used in this experiment.

#### IV. Analysis

A. Photofragment Yield Spectra of CH<sub>3</sub>O and CD<sub>3</sub>O. The rovibronic term value  $T_0$  for the  $\tilde{\rm A}(^2{\rm A}_1) \leftarrow \tilde{\rm X}(^2{\rm E})$  transition in CH<sub>3</sub>O was given by Liu *et al.*<sup>25</sup> as 31 614.51 cm<sup>-1</sup> from rotational analysis of the combined LIF and microwave<sup>14</sup> data. The effective  $0_0^0$  transition, *i.e.*, including spin—orbit splitting, measured from the lowest level of the  $^2{\rm E}_{3/2}$  component of the ground state, is 31 644.6 cm<sup>-1</sup>. It is convenient to use the value when comparing the vibronically resolved  $\tilde{\rm A}(^2{\rm A}_1) \leftarrow \tilde{\rm X}(^2{\rm E}_{3/2})$  components in our PFY spectra with jet-cooled LIF experiments, in which all transitions originate from the  $^2{\rm E}_{3/2}$  component.

The significant increase in CO bond length for the  $\tilde{A} \leftarrow \tilde{X}$ transition gives rise to an extended Franck-Condon progression in the  $v_3$  mode for both absorption and LIF measurements, and the same progression should dominate the PFY spectrum. While fluorescence is observed in CH<sub>3</sub>O up to  $\approx$ 37 000 cm<sup>-1</sup>, spectroscopic assignments from LIF are complicated in this energy range due to spectral congestion and a Fermi resonance between  $v_2$  and  $2v_3$  in the  $\tilde{A}$  state. We assign the lowest energy dissociative transition as 3<sub>0</sub><sup>6</sup>, corresponding to an energy 3775  $cm^{-1}$  above the zero-point level of the  $\tilde{A}$  state. This transition represents the first dissociative member of the  $3_0^n$  progression, which may also contain contributions from  $3_0^{n-2}2_0^1$ ,  $3_0^{n-4}2_0^2$ , etc., due to the Fermi resonance. On the basis of the most recent rovibrationally resolved LIF data,  $^{30}$  the  $3_0^6$  transition occurs at 35 437 cm<sup>-1</sup>, compared with 35 419 cm<sup>-1</sup> in our PFY spectrum. The comparison of LIF with PFY transition frequencies agrees to within experimental uncertainty when one considers that our rotational temperature (35-50 K) is greater than in the LIF spectra (5 K), displacing the band maxima (which we measure) from the true rovibronic origin. In addition, our laser step size of  $\approx$ 6 cm<sup>-1</sup> prevents us from determining band maxima more precisely than this value.

In our previous analysis of the PFY spectrum of CH<sub>3</sub>O,<sup>7</sup> we assigned the combination band shown in Figure 4 as  $3_0^n 5_0^1$ , the only feasible assignment based on the accepted vibrational frequencies of the  $\tilde{A}$  state at that time.<sup>23</sup> Due to a recent reassignment<sup>30</sup> of the  $\nu_6$  fundamental frequency in the  $\tilde{A}$  state (previously 595 cm<sup>-1</sup>, currently 929.5 cm<sup>-1</sup>), it is unclear from the peak positions alone whether the combination band we observe in the PFY spectrum of CH<sub>3</sub>O arises from the  $\nu_5$  (1403 cm<sup>-1</sup>) or the  $\nu_6$  mode. The ambiguity arises because the  $3_0^n 5_0^1$  and  $3_0^{n+1} 6_0^1$  progressions have nearly identical frequencies.

However, our dynamics results discussed in section V.B.1, together with recent fluorescence depletion spectroscopy results, 44 suggest that the combination band observed in dissociation involves the  $\nu_6$  methyl rocking mode rather than  $\nu_5$ . The  $3_0^56_0^1$  transition measured by LIF<sup>30</sup> occurs at 35 673 cm<sup>-1</sup>, compared with 35 666 cm<sup>-1</sup> in the PFY spectrum.

The PFY spectrum of CD<sub>3</sub>O (Figure 5) shows some interesting differences from that of CH<sub>3</sub>O. Assignment of the  $3_0^n$ progression is straightforward, due to the small isotope shift in this mode ( $v'_3 = 663 \text{ cm}^{-1}$ ).<sup>23</sup> However, the combination band we observe is more difficult to assign. While one might expect that the same  $3_0^n 6_0^1$  progression found in CH<sub>3</sub>O will be active in CD<sub>3</sub>O, this assignment gives poor agreement with the data. Specifically, the  $v_5$  fundamental in CD<sub>3</sub>O is reported as 1047 cm $^{-1}$ , <sup>23</sup> and we estimate the  $\nu_6$  fundamental as 693 cm $^{-1}$  (based on the measured isotope shift in  $\nu_5$ ). From our observation of the  $3_0^6$  transition at 35 404 cm<sup>-1</sup>, we expect the  $3_0^6 5_0^1$  transition at  $\approx$  36 451 cm<sup>-1</sup>, which is in fact a minimum between two peaks in Figure 5, and the  $3_0^6 6_0^1$  transition at  $\approx 36~097~\text{cm}^{-1}$ , which lies on the blue edge of the  $3_0^7$  transition. By contrast, the  $\nu_2$  fundamental (CD<sub>3</sub>O umbrella) is 971 cm<sup>-1</sup>, from which the  $3_0^6 2_0^1$  transition is predicted at  $\approx 36 \text{ } 375 \text{ } \text{cm}^{-1}$ , in better agreement with the first peak at 36 330 cm<sup>-1</sup> in the strong combination band of CD<sub>3</sub>O. Therefore, we assign the combination band in the PFY spectrum of CD<sub>3</sub>O as  $3_0^n 2_0^1$ , an assignment corroborated by the dissociation dynamics discussed in section

**B.** Translational Energy Distributions. 1.  $CH_3O$ . Analysis of the  $P(E_T)$  distributions will focus primarily on the  $CH_3$  + O product channel, for which the most detailed information is obtained. The main goals of the analysis are to determine (i) the best value of the bond dissociation energy  $D_0(CH_3-O)$ , (ii) the  $CH_3$  fragment vibrational distribution, and (iii) a measure of rotational excitation of the  $CH_3$  fragment. The major analysis results are unchanged from those presented previously,  $^7$  but a fuller description of the analysis is given here.

The balance of energy for photodissociation of methoxy is given by

$$h\nu + E_{\text{int}}(\text{CH}_3\text{O}) = D_0(\text{CH}_3-\text{O}) + E_{\text{T}} + E_{\text{V}}(\text{CH}_3) + E_{\text{R}}(\text{CH}_3) + E_{\text{SO}}(\text{O}^3\text{P}_i)$$
 (3)

where  $E_{\rm int}$  is the most probable rotational energy of the parent radical,  $E_V$  and  $E_R$  are the product vibrational and rotational energy, and  $E_{SO}$  is the spin-orbit energy of the oxygen atom. On the basis of a rotational temperature of 50 K, we estimate  $E_{\rm int}({\rm CH_3O}) = 35~{\rm cm^{-1}}$ . We can determine  $D_0$  since we measure  $E_{\rm T}$  directly, but only if we locate an energy in the  $P(E_{\rm T})$ distribution for which the last three terms of eq 3 are zero. The steep falloff in intensity on the high-energy side of each distribution in Figure 6 is evidence for this thermodynamic limit, in which all the available energy goes into product translation, forming  $CH_3(v = 0, J = 0) + O(^3P_2)$ . For each data set we obtain an independent estimate of  $D_0$  by extrapolating to the energy where the  $P(E_T)$  distribution reaches zero intensity. An average of these thresholds, deconvoluted from the experimental resolution, gives the best value of  $D_0(\text{CH}_3-\text{O}) = 3.807 \pm 0.013$ eV. From the known heats of formation of CH<sub>3</sub> and O, 45 we obtain the heat of formation  $\Delta H^{\circ}_{f,0}(\text{CH}_3\text{O}) = 0.29 \pm 0.02 \text{ eV}$  $(6.8 \pm 0.4 \, \text{kcal/mol})$ . Additional thermodynamic quantities are derived from this value in ref 7.

In order to determine the product vibrational distributions, the data are fit to a set of rotational/spin—orbit distribution functions separated by the accurately known term energies of the CH<sub>3</sub> umbrella mode.<sup>57</sup> The distribution functions  $f_n(E_T)$  are

**TABLE 1: Product Branching Ratio for CH<sub>3</sub>O → CH<sub>3</sub> + O** 

transition	$hv^a$ (cm <sup>-1</sup> )	fwhm <sup>a</sup> (meV)	$\langle E_{\rm V} \rangle^b  ({\rm meV})$	v = 0	v = 1	v = 2	v = 3	v = 4	v = 5
36	35,419	68/63	12	84	14	1	0	0	0
$3_0^{7}$	36,037	47/39	14	82	16	1	0	0	0
38	36,619	50/41	26	67	28	3	1	0	0
$3_0^9$	37,205	53/43	44	52	40	7	1	0	0
$3_0^{10}$	37,774	53/43	73	35	45	16	4	1	0
$3_0^{13}$	39,384	73/70	179	19	15	20	28	14	4
$3_0^6 6_0^1$	36,268	53/45	10	87	11	1	0	0	0
$3_0^{7}6_0^{1}$	36,842	58/50	14	80	17	1	0	0	0

<sup>&</sup>lt;sup>a</sup> fwhm of vibrational peaks after deconvolution of experimental resolution. The left value in this column assumes production of only  $O(^3P_2)$ , while the right value assumes a statistical  $O(^3P_j)$  distribution (see text). <sup>b</sup> Average vibrational energy.

**TABLE 2:** Product Branching ratio For  $CD_3O \rightarrow CD_3 + O$ 

transition	hv (cm <sup>-1</sup> )	$fwhm^a (meV)$	$\langle E_{\rm V} \rangle^b  ({\rm meV})$	v = 0	v = 1	v = 2	v = 3	v = 4	v = 5
36	35,404	80	62	37	38	14	7	4	0
37	35,996	47	48	39	46	10	4	1	0
38	36,590	51	65	28	47	17	5	2	1
$3_0^9$	37,119	52	70	25	43	23	7	2	0
$3_0^{10}$	37,679	63	88	18	39	28	10	4	1
$3_0^6 2_0^1$	36,330	57	74	43	11	31	11	4	0
$3_0^{7} 2_0^{1}$	36,934	51	69	47	11	27	12	3	0
$3_0^8 2_0^{1}$	37,516	52	79	41	17	20	17	4	1

<sup>&</sup>lt;sup>a</sup> fwhm of vibrational peaks after deconvolution of experimental resolution. <sup>b</sup> Average vibrational energy.

nearly Gaussian in shape,<sup>58</sup> with an asymmetric tail extending to lower translational energy, *i.e.*, toward higher rotational energy. The shape of this function is identical for every vibrational peak in a given spectrum. The model distribution,  $F(E_{\rm T})$  is given by

$$F(E_{\rm T}) = \sum_{n=0}^{n'} \alpha_n f_n \{ E_{\rm T} - (h\nu - D_0 - n\omega_2 - \Delta); \Gamma \}$$
 (4)

Here n labels the number of quanta in the CH<sub>3</sub>  $\nu_2$  umbrella mode,  $\Gamma$  is the fwhm of the combined rotational/spin—orbit envelope and instrumental resolution, and  $\Delta$  is the energy difference between the maximum possible  $E_{\rm T}$  for each n { $E_{\rm T}^{\rm max}$ - $(n) = h\nu - D_0 - n\omega_2$ } and the peak of the nth rotational distribution function. The coefficients  $\alpha_n$  give the CH<sub>3</sub> product vibrational distribution, while the instrumental resolution can be deconvoluted from the width  $\Gamma$  to give an indication of the energy balance between the last two terms of eq 3.59

The solid line in Figure 6 shows the best nonlinear least-squares fit to each data set, with the results given in Table 1. We note that the fits are sensitive to the strong negative anharmonicity in the  $\nu_2$  mode of CH<sub>3</sub>, confirming our assignment that the highest energy peak in each spectrum corresponds to  $\nu_2 = 0$ . The third column of Table 1 shows the fwhm of the rotational envelope in meV for two limiting cases. The larger values fit the data under the assumption that all O atoms are produced in their  $^3P_2$  ground state, while the smaller values result from assuming a statistical 5:3:1 population of the  $^3P_{2:1:0}$  spin—orbit states (which have energies of 0:20:28 meV,  $^{60}$  respectively). Both limits reproduce the data well, implying that this data is not sufficiently sensitive to distinguish between the two cases.

Finally, we consider product state distributions for channels II and III, producing  $CH_2 + OH$ , which are observed in the energy range  $E_T < 0.5$  eV upon excitation of  $3_0^{10}$  and  $3_0^{13}$ . While there is no reproducible vibrational structure in this data, we can speculate on the relative importance of  $CH_2(\tilde{X}(^3B_1))$  vs  $\tilde{a}(^1A_1)$ . Referring to Figure 6, any data with  $E_T > E_{\max}^T$  III can only be due to fragmentation via channel II. For  $E_T < E_{\max}^T$  III, both channels can contribute, but the significant rise in the

signal concurrent with the opening of the electronically excited methylene channel implies that channel III is the major source of  $CH_2 + OH$ . While it is energetically possible that the data with  $0.6 \le E_T \le 0.78$  eV for  $3_0^{13}$  is due to channel II, the mass spectrum for this energy range implies that this data arises from  $CH_3 + O$ , as denoted by the symbols in Figure 6.

2. CD<sub>3</sub>O. A similar data analysis is undertaken for the CD<sub>3</sub> + O product channel in the deuterated species, with the results given in Table 2. Deuteration lowers the frequency of the methyl radical umbrella mode from 606 to 458 cm<sup>-1</sup>,61 which explains the less-resolved vibrational structure observed in the top two panels of Figure 7. Two important differences are observed in comparison to CH<sub>3</sub>O dissociation. First, for a given level of excitation in the  $3_0^n$  progression, the CD<sub>3</sub> vibrational distributions are substantially hotter than those observed for CH<sub>3</sub>. Second, the vibrational distributions for the  $3_0^n 2_0^1$  series are bimodal, with the population of  $v_2 = 0$  and  $v_2 = 2$  always greater than that for  $v_2 = 1$ . The origin of these effects will be discussed in the next section. Fitting all the data sets for CD<sub>3</sub>O gives the best value of  $D_0(\text{CD}_3-\text{O}) = 3.85 \pm 0.02 \text{ eV}$ , and therefore  $\Delta H^{\circ}_{f,0}(CD_3O) = 0.16 \pm 0.04 \text{ eV } (3.7 \pm 0.9 \text{kcal/mol}).^{62}$ Note the significant difference of 0.13 eV (3.1 kcal/mol) compared to the heat of formation of CH<sub>3</sub>O, reflecting differences in vibrational frequencies between the H and D isotopes.

## V. Discussion

### A. Photofragment Yield Spectra of CH<sub>3</sub>O and CD<sub>3</sub>O.

One of the important conclusions from the PFY spectrum is that, within our detection limits,  $CH_3O$  does not dissociate for photon energies below the  $3_0^6$  transition. This threshold implies that the crossing of the  $\tilde{A}$  state by the lowest of the three repulsive curves shown in Figure 2 occurs in the vicinity of  $3775~\rm cm^{-1}$  above the  $\tilde{A}$  state zero-point level. The onset of dissociation and the energy of the curve crossing has been the subject of much debate among both experimentalists and theorists, with our value being the lowest reported. Our threshold is unambiguous because we detect dissociation directly by measuring the presence of primary photofragments. A very recent determination of the dissociation threshold by Powers et

*al.*,<sup>44</sup> via LIF lifetime measurements and fluorescence depletion spectroscopy (FDS), gives 3793 cm<sup>-1</sup>, in good agreement with our value

One of the main complexities in the  $A(^2A_1)$  state of CH<sub>3</sub>O arises because two quanta of  $\nu_3$  establish a Fermi resonance with  $\nu_2$  (CH<sub>3</sub>O umbrella, 1289 cm<sup>-1</sup>), giving rise to a series of multiplets at each overtone in the  $3_0^n$  series. By fitting the observed multiplet spectra to an anharmonic oscillator Hamiltonian for  $\nu_2$  and  $\nu_3$ , Powers *et al.* Obtained the dominant character of each member of the Fermi multiplet. An interesting question is whether the competition between predissociation and fluorescence depends on which member of the multiplet is excited, in which case the PFY and LIF spectra would differ. If mixing between  $\nu_2$  and  $\nu_3$  is not too strong, one might expect that components of predominantly  $\nu_3$  character would dissociate more readily than those with more  $\nu_2$  character.

The splittings between Fermi multiplet components are significant. For example, the  $3_0^5$ ,  $3_0^12_0^2$ , and  $3_0^32_0^1$  bands are separated by 35 and 43 cm<sup>-1</sup>, respectively,  $^{30}$  giving a total width to this feature of 78 cm<sup>-1</sup>. However, the peak widths in the PFY spectrum (Figure 4) are also broadened due to the width of the rotational bands (at least 20 cm<sup>-1</sup>) and the fact that we observe transitions from both  $^2E_{3/2}$  and  $^2E_{1/2}$  components ( $\Delta E \approx 60 \text{ cm}^{-1}$ ) of the ground state. It is therefore difficult to determine whether different multiplet components decay with different rates from the PFY spectrum. In fact, the LIF and FDS experiments of Powers *et al.*<sup>44</sup> show no significant differences in lifetimes for different members of a particular multiplet, indicating that  $\nu_2$  and  $\nu_3$  are indeed strongly mixed.

In CD<sub>3</sub>O, the Fermi resonance between  $\nu_2$  and  $\nu_3$  is lifted, and the vibrational frequencies of all modes except  $\nu_3$  are substantially lowered. As a consequence, the  $3_0^n 2_0^1$  combination band now appears as a distinct progression. The  $3_0^6 2_0^1$ transition is the first member of this band seen in the PFY spectrum. While dissociation is energetically possible from excitation of the  $3_0^5 2_0^1$  transition (expected at 35 670 cm<sup>-1</sup>), this transition is not observed in Figure 5, implying that for CD<sub>3</sub>O, energy deposited in  $v_2$  is not available to facilitate fragmentation. In contrast, CH<sub>3</sub>O dissociates when the  $3_0^56_0^1$  transition is excited, i.e., with five quanta in  $v_3$ . The isotopic comparison implies that excitation of the e symmetry vibration  $\nu_6$  (observed for CH<sub>3</sub>O) promotes dissociation, while energy deposited in the  $a_1$  symmetry  $v_2$  mode (observed for CD<sub>3</sub>O) is not strongly coupled to the reaction coordinate. We give further evidence for these conclusions in section V.B., based on the translational energy and angular distribution data.

**B.** Channel I Dissociation Dynamics. Unimolecular photodissociation is a powerful technique because the disposal of energy among translation, vibrational, rotational, and electronic degrees of freedom in the products is a direct consequence of the potential energy surfaces on which this half-collision occurs. Consequently, the main information we obtain on the dissociation dynamics of the methoxy radical is derived from the  $P(E_T)$  distributions presented graphically in Figures 6 and 7 and numerically in Tables 1 and 2. In this section we discuss the dissociation mechanism for channels I—III in light of the experimental product state distributions.

1.  $CH_3 + O$  Products. It is evident from Figure 6 that most of the available energy in the  $\tilde{A} \leftarrow \tilde{X}$  photodissociation of  $CH_3O$  goes into product translation for channel I. Although radicals similar to methoxy, such as  $CH_2CHO$ , dissociate in the ultraviolet by internal conversion (IC) to the ground state potential energy surface (PES), the  $P(E_T)$  distributions for methoxy are inconsistent with such a mechanism. Indeed, SEP studies<sup>6</sup> on  $CH_3O$  imply that, if IC were facile,  $H + CH_2O$ 

would be the dominant product channel. Instead, the large recoil energies observed in our data are indicative of dynamics on a purely repulsive PES, in which potential energy is converted efficiently into translational kinetic energy. The results of *ab initio* calculations by Jackels,  $^{33}$  represented schematically in Figure 2, predict three excited electronic states of  $^4A_2$ ,  $^4E$ , and  $^2A_2$  symmetry, which are repulsive along the C–O bond and correlate asymptotically to  $CH_3(\tilde{X}(^2A_2^{\prime\prime}))+O(^3P)$ . All our results for this channel are consistent with a mechanism in which non-adiabatic coupling between these surfaces and the optically prepared levels of the  $\tilde{A}(^2A_1)$  state leads to predissociation of  $CH_3O$ . Given this basic description of the dissociation mechanism, we can explore further mechanistic details in the experimental data.

First, we consider the flow of energy into product vibration. Based on the fitting procedure presented in section IV.B., the  $P(E_{\rm T})$  distribution is described accurately by excitation in a single CH<sub>3</sub> mode, the  $\nu_2$  umbrella vibration. Focusing on the  $3_0^n$  progression, and ignoring for the moment n=6, a smooth increase in the average CH<sub>3</sub> vibrational excitation  $\langle E_{\rm V} \rangle$  is observed with increasing photon energy (Table 1). Very little product excitation arises from the  $3_0^7$  level, in contrast to the inverted distributions for  $3_0^{10}$  and  $3_0^{13}$ .

The sudden approximation is often invoked to explain dissociation dynamics on purely repulsive surfaces. In this model the fragments are assumed to dissociate sufficiently rapidly that there is negligible coupling between translational and vibrational degrees of freedom. In the sudden limit, the CH<sub>3</sub> vibrational distribution is obtained by a Franck–Condon projection of the CH<sub>3</sub> moiety in the  $\tilde{A}$  state onto the asymptotic  $\nu_2$  vibrational levels of the free methyl radical. Specifically,  $\angle$ HCO in the  $\tilde{A}$  state is 106°, while asymptotically the methyl radical is planar, corresponding to  $\angle$ HCO = 90° within  $C_{3\nu}$  symmetry. Using *ab initio* force constants for CH<sub>3</sub>, the sudden approximation predicts a most probable value of  $\nu_2$  = 5 for the CH<sub>3</sub>  $\nu_2$  distribution, somewhat hotter than the distribution observed at  $3_0^{13}$  and clearly incorrect for all lower photon energies.

The failure of the sudden approximation implies substantial coupling of translational and vibrational degrees of freedom as the dissociation proceeds on the repulsive surface, allowing ∠HCO to relax smoothly with increasing CO bond length. In other words, at low photon energies, e.g.,  $3_0^7$  excitation, CH<sub>3</sub>O appears to evolve adiabatically to products. This situation is entirely reasonable when one considers that the light H atoms can respond quickly to motion of the heavy C and O atoms as  $r_{\rm CO}$  increases, maintaining an equilibrium value of  $\angle$ HCO at each value of  $r_{CO}$ . This picture also explains the behavior of the  $3_0^n P(E_T)$  distributions with increasing photon energy. The recoil velocity between the C and O atoms increases with increasing available energy, such that the H atoms cannot respond as accurately to the changing potential along the ∠HCO coordinate, resulting in increased asymptotic CH3 vibrational excitation.

The photodissociation dynamics of methyl iodide show a strong similarity to our results on methoxy. For the  $CH_3I \rightarrow CH_3 + I^*$  channel, which occurs on a purely repulsive surface, the methyl fragment is produced with little vibrational or rotational excitation. An adiabatic mechanism very similar to that described above for methoxy governs the dynamics of this channel in  $CH_3I$ , as elucidated by classical trajectory calculations on *ab initio* potential energy surfaces for this system.<sup>64</sup>

We next consider partitioning of energy between the CH<sub>3</sub> rotational energy and the spin-orbit energy of  $O(^3P_j)$  (eq 3). As shown in column 3 of Table 1, the peak widths of each

vibrational component increase monotonically in the  $3_0^n$  progression for  $n \geq 7$ , denoting increasing rotational and/or spin—orbit product excitation. Although we present the rotational envelope widths for the two limiting cases in which no energy or a statistical distribution of energy is deposited in the oxygen atom, there is no reason to believe that either limit reflects reality, or that the spin—orbit state distribution is independent of photon energy, as our two limiting cases assume. Indeed, by analogy with the spin—orbit distributions measured in  $O_2$  dissociation<sup>65</sup> it is likely that the j distributions of the oxygen atoms from CH<sub>3</sub>O are not statistical but fluctuate in a seemingly random fashion depending the value of n. Subject to this assumption, we conclude that the increasing peak widths in Table 1 reflect an increase of rotational excitation as a function of photon energy.

However, the amount of the available energy deposited in product rotation is still relatively small. For example, in the  $3_0^{10}$   $P(E_{\rm T})$  distribution (Figure 6), the energy difference between  ${\rm CH}_3(v=0,J=0,K=0)$  and the peak of the  ${\rm CH}_3(v=0)$  distribution is 50 meV (6% of the available energy). This energy corresponds to the rotational energy of a  ${\rm CH}_3(v=0,J=6,K=0)$  fragment when the cofragment is  ${\rm O}(^3{\rm P}_2)$ ; an excited O atom would require even smaller rotational quantum numbers for  ${\rm CH}_3$ . Simply put, the fact that we resolve vibrational structure at all in the  $P(E_{\rm T})$  distributions means that product rotational excitation is not extensive. We propose that the small amount of product rotation indicates a dissociation pathway which deviates little from  $C_{3v}$  symmetry for  $3_0^n$  excitation.

In contrast to the  $P(E_T)$  distributions just discussed, the distributions for  $3_0^6$  and  $3_0^n 6_0^1$  lack the well-resolved vibrational features present in the other spectra in Figure 6. For  $3_0^6$ , the much broader vibrational peak is likely due to an experimental effect rather than an increase in rotational excitation with respect to  $3_0^7$ . The most recent literature value for the fluorescence lifetime of  $3_0^6$  is 0.37  $\mu$ s,<sup>44</sup> a substantial fraction of the 9  $\mu$ s flight time required for a CH<sub>3</sub>O radical with 8 keV energy to travel from the dissociation laser to the TPS detector. Excited state lifetimes in this range degrade the  $P(E_T)$  distributions to lower  $E_{\rm T}$  because long-lived radicals remain intact for a portion of the flight time, resulting in a smaller recoil vector measured at the detector, which translates to an artificially small value of  $E_{\rm T}$  for that event. Using the fluorescence lifetime given above, this effect causes a 1/e broadening of 51 meV.66 When this width is deconvoluted from the 68 meV width given in Table 1, the remaining rotational envelope has a width of 45 meV, slightly narrower than the 47 meV width measured for  $3_0^7$ . Therefore, rotational excitation of the CH<sub>3</sub> + O products following  $3_0^6$  excitation follows the same trend as the other members in the  $3_0^n$  progression despite the broad peak observed in Figure 6.

The  $P(E_{\rm T})$  distributions following  $3_0^6 6_0^1$  and  $3_0^7 6_0^1$  excitation show broader peaks than their quasi-isoenergetic counterparts in the  $3_0^n$  progression. Powers  $et~al.^{44}$  found that the fluorescence lifetime for  $3_0^6 6_0^1$  lies between the limits 20 ns  $\geq \tau_{\rm FL} \geq$  20 ps. As a result of this short lifetime, the broadening effect described above for  $3_0^6$  is negligible compared to the experimental resolution, and the increased width of the peaks is likely a result of greater product rotational excitation. The observation of rotational broadening is reasonable when one considers that the  $\nu_6$  methyl rocking motion would naturally develop into rotational excitation of the methyl radical about the b-axis, and supports the assignment that the combination band in Figure 4 is principally  $3_0^n 0_0^1$  rather than  $3_0^n 5_0^1$ .

The product angular distributions for  $3_0^6 6_0^1$  and  $3_0^7 6_0^1$  excitation are described by anisotropy parameters of  $\beta = 0.4$  and  $\beta$ = 0.3, respectively, while the angular distributions of the  $3_0^n$ progression are isotropic ( $\beta = 0$ ). However, the  $\tilde{A}(^2A_1) \leftarrow \tilde{X}$ -(2E) transition has a perpendicular transition dipole moment, which should result in an anisotropy parameter  $\beta \leq 0$ . Although the vibronic symmetry of  $3_0^n 6_0^1$  is parallel (e  $\leftarrow$  e), the electronic part of the transition dipole remains (a  $\leftarrow$  e) in the absence of perturbations by other electronic states,<sup>30</sup> and hence should still give  $\beta \leq 0$ . Therefore the positive values of  $\beta$ measured for  $3_0^6 6_0^1$  and  $3_0^7 6_0^1$  excitation implies that the  $\tilde{A}$  state is perturbed by another electronic state of e symmetry (such as the nearby <sup>4</sup>E state shown in Figure 2), which mixes an electronic e ← e component into the transition dipole. These results are consistent with the observation of both parallel and perpendicular rotational structure resolved in CH<sub>3</sub>O bands containing fundamentals of e vibrations.<sup>30</sup>

2.  $CD_3 + O$  *Products*. The  $P(E_T)$  distributions for the C-O bond fission channel in  $CD_3O$  show two main differences from the  $CH_3O$  distributions. First, the  $CD_3$  vibrational distributions resulting from  $3_0^n$  excitation are substantially more excited than at similar photon energies in  $CH_3O$ , as verified by comparison of the average product vibrational energies  $\langle E_V \rangle$  in Tables 1 and 2. Second, the  $CD_3$  vibrational distributions resulting from  $3_0^n 2_0^1$  excitation are bimodal, as shown in Figure 7.

Increased vibrational excitation of the  $CD_3$  umbrella motion resulting from  $3_0^n$  excitation follows directly from the adiabatic dissociation mechanism proposed for channel I dynamics in  $CH_3O$ . When H atoms are replaced with D atoms in methoxy, the increased mass inhibits the ability of these atoms to track the motion of the C and O atoms during dissociation. As a result, the dissociation is less vibrationally adiabatic, resulting in more excitation of the  $\nu_2$  mode in  $CD_3$ .

A similar isotope effect is observed in CD<sub>3</sub>I dissociation, and the complete description of these effects found in Ref. 64 is most likely applicable to CD<sub>3</sub>O. However, for the present purposes, the simple explanation given above is sufficient.

By far the most surprising result from the CD<sub>3</sub>O  $P(E_T)$ distributions is the bimodal CD3 product state distribution observed following excitation of the  $3_0^n 2_0^1$  combination band (cf. Table 2). To the best of our knowledge, methoxy photodissociation represents the first observation of such an isotope effect on the *product* state distributions. A compelling hypothesis for the origin of this dynamical effect lies in the near equality between the umbrella mode frequency in the A state of CD<sub>3</sub>O (971 cm<sup>-1</sup>) and the corresponding first overtone frequency of the umbrella mode in the CD<sub>3</sub> fragment  $2\nu_2$ (CD<sub>3</sub>) = 966 cm<sup>-1</sup>.61 Therefore, if the single quantum of  $v_2(CD_3O)$ excited in the  $3_0^n 2_0^1$  combination band does not couple well to the reaction coordinate, the 971 cm<sup>-1</sup> of energy deposited into this motion will result in a propensity to populate the nearly degenerate first overtone of  $\nu_2(CD_3)$ , i.e., the amount of energy in umbrella motion will be conserved throughout the dissociation. This hypothesis is also supported by the fact that  $3_0^5 2_0^1$ does not dissociate (cf. Figure 5), even though this state has more energy than the  $3_0^6$  level, which does dissociate. This intriguing hypothesis calls for a comparison with reaction dynamics calculations based on the new potential energy surfaces which are now becoming available for methoxy.<sup>39,40</sup>

3. Curve-Crossing mechanism. Until now we have discussed the dissociation dynamics without consideration of the mechanism that couples the optically accessed  $\tilde{A}(^2A_1)$  state with the  $^4A_2$ ,  $^4E$ , and  $^2A_2$  repulsive states. The calculations of Jackels demonstrated that the  $^4E$  state is reached via a one-electron

transition from the  $\tilde{A}$  state, while the two  $A_2$  configurations require the rearrangement of two electrons.<sup>33</sup> He argued that the  ${}^{2}A_{1} \leftrightarrow {}^{4}E$  interaction should be the strongest, because these states have a first-order spin-orbit interaction, analogous to that of the  ${}^2\Sigma \leftrightarrow {}^4\Pi$  coupling in a diatomic radical. The  ${}^2A_1 \leftrightarrow {}^4A_2$ interaction can occur by a second-order spin-orbit interaction, while the  ${}^{2}A_{1} \leftrightarrow {}^{2}A_{2}$  interaction is most likely dominated by off-diagonal elements in the nuclear kinetic energy operator.

Recent ab initio calculations by Pederson and Yarkony, 39 and by Cui and Morokuma<sup>40</sup> have defined the minimum seam of crossings between the A state and each repulsive curve. The new calculations place the crossing points at lower energy than those obtained by Jackels, in better agreement with our photodissociation threshold. The coupling constants between the states are also calculated, along with the dependence of the coupling at geometries away from  $C_{3v}$  symmetry.<sup>39</sup>

Until now, the experimental data has been discussed for simplicity under the tacit assumption that one repulsive curve accounts for the entire dynamics of channel I. Two features in Figure 6 may shed some light on whether coupling with multiple curves contributes to the final product state distributions. First, the  $P(E_{\rm T})$  distribution following  $3_0^7$  excitation is much narrower than the  $3_0^6$  distribution, indicating a substantial increase in dissociation rate at the higher of these two photon energies (see section V.B.1.). One hypothesis for the increased dissociation rate is that the  $3_0^6$  distribution is mediated primarily by the <sup>4</sup>A<sub>2</sub> surface, while the 3<sub>0</sub><sup>7</sup> distribution contains a new contribution from the <sup>4</sup>E surface. This picture agrees with Jackel's argument that coupling with the <sup>4</sup>E surface is stronger than coupling to the other two repulsive surfaces. However, it is also possible that dissociation following  $3_0^6$  excitation may require tunneling through a small barrier, while at  $3_0^7$  dissociation occurs without a barrier. A second noteworthy feature from Figure 6 is that the CH<sub>3</sub> vibrational excitation resulting from  $3_0^{13}$  excitation is bimodal (cf. Table 1), which could plausibly result from the contribution of an additional repulsive curve at this high photon energy leading to vibrationally cold products. Detailed assessment of these issues should soon be possible in light of the new potential energy surfaces being developed.

C. Channel II-IV Dissociation Mechanisms. Our experimental information on the CH2 + OH product channel is limited. We find that this product channel is absent for transitions with  $h\nu < 37 500 \text{ cm}^{-1}$  and competes with the dominant CH<sub>3</sub> + O channel at higher photon energies. As pointed out in section IV.B on energetic grounds it appears that channel III, producing electronically excited methylene fragments, dominates production of ground state methylene, channel II. Concerning the mechanism for this channel, the recent ab initio study of the methoxy system by Cui and Morokuma<sup>40</sup> explored the excited state isomerization of  $CH_3O \rightarrow CH_2OH$ , finding a transition state 1.00 eV above the A state minimum. They point out that the A state of CH<sub>3</sub>O correlates through this transition state to channel III products, while the  $\tilde{X}$  state correlates to channel II products. Our data are consistent with the excited state mechanism, and the observed production of some ground state methylene could easily be explained by a transition to a lower surface along the isomerization or dissociation coordinate. A similar mechanism is probably responsible for  $CD_2 + OD$  dissociation in  $CD_3O$ .

Cui and Morokuma also found a substantial barrier to formation of H + CH<sub>2</sub>O on excited state surfaces, in contrast to the small barrier on the ground state for this channel. The fact that we observe very little H + CH<sub>2</sub>O is consistent with their result, indicating that all the dynamics in ultraviolet methoxy dissociation occur on excited state surfaces.

#### VI. Conclusions

The ultraviolet  $\tilde{A}(^2A_1) \leftarrow \tilde{X}(^2E)$  transition in both CH<sub>3</sub>O and CD<sub>3</sub>O is observed to predissociate due to curve crossings with one or more of the excited <sup>4</sup>A<sub>2</sub>, <sup>4</sup>E, and <sup>2</sup>A<sub>2</sub> repulsive states. Several interesting differences in the photodissociation dynamics have been observed between the two isotopes. The photofragment yield spectrum of CH<sub>3</sub>O shows a main progression in the C-O stretch and a combination band with one quantum of the e symmetry  $v_6$  methyl rock. The  $3_0^n$  progression also dominates in CD<sub>3</sub>O, but the  $3_0^n 2_0^1$  is the most prominent combination band involving the  $a_1$  symmetry  $v_2$  umbrella motion. The  $3_0^6$ transition is the lowest energy transition observed to dissociate in either isotope, placing the threshold for CH<sub>3</sub>O dissociation 3775 cm<sup>-1</sup> above the zero point level of the A state. In CH<sub>3</sub>O the combination band  $3_0^n 6_0^1$  dissociates for n = 5, while in  $CD_3O$  combination band dissociation is not observed unless n≥ 6. This mode-specific effect demonstrates that the way in which energy is distributed in the A state effects the competition between dissociation and fluorescence.

The major product channel is C-O bond fission giving a methyl radical and an oxygen atom. The translational energy distributions for both isotopes show that the  $v_2$  umbrella mode of the CH<sub>3</sub>(CD<sub>3</sub>) fragment is excited, although most of the available energy is channeled into product translation. The photofragment vibrational distributions lie much closer to the adiabatic limit than the sudden limit, indicating that the light H atoms can follow dissociation along the C-O bond distance; this picture is less accurate for CD<sub>3</sub>O than CH<sub>3</sub>O, as expected. Excitation of  $3_0^n 6_0^1$  in CH<sub>3</sub>O results in more rotational excitation of the products, consistent with sampling of non- $C_{3\nu}$ geometries in the  $\tilde{A}$  state. Excitation of  $3_0^n 2_0^1$  in  $CD_3O$  shows a surprising dynamical effect: the product state distribution of the CD<sub>3</sub> fragment is bimodal, with excitation of  $v_2 = 0$  and  $v_2$ = 2 always greater than that for  $v_2 = 1$ . This effect is clear evidence that the energy distribution among parent vibrational modes has a strong effect on the vibrational distribution in the products.

The results given in this article should enable a rigorous comparison between experiment and theory because the data comprise both translational and vibrational product state distributions, acquired at many different photon energies for two isotopes. New ab initio potential energy surfaces are now becoming available for methoxy, and it appears that the methoxy radical will continue its tradition as a benchmark molecule in the fields of spectroscopy and dynamics.

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