Photodissociation of methanol at 193.3 nm: Translational energy release spectra

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Center-of-mass translational energy distributions of the dominant primary products resulting from 193.3 nm excitation of jet-cooled CH₃OH, CH₃OD, and CD₃OH were obtained by using the high-n Rydberg time-of-flight (HRTOF) technique. The appearance threshold in the HRTOF spectrum yields a bond dissociation energy, D₀(CH₃O-H), of 105±1 kcal mol⁻¹, in agreement with recent literature values. Translational energy release spectra from the three isotopomers exhibit progressions of 950±100 cm⁻¹, which are attributed to excitation in the ν₂ O–CH₃ stretch of the methoxy product. The progressions peak at ν = 1, with population out to at least ν = 5. This differs from the results of a recent wave packet dynamics study on a calculated excited state potential energy surface [Marston et al., J. Chem. Phys. 98, 4718 (1993)], which predicted no O–CH₃ stretch excitation in the methoxy fragment following photolysis of ground state methanol. The spatial anisotropy of the fragments (β = -0.7) implies a dissociation time ~1 ps. The impulsive model for rotational excitation is compared to the unresolved rotational contour of the vibrational peaks in the translational energy release spectra and is found to underestimate the extent of rotational excitation, though the model correctly predicts the increase in contour width observed for the O-deuterated species. The unresolved rotational contours are fit empirically. The inferred vibrational energy distributions are discussed in terms of a simple Franck-Condon model for the pseudodiatomic, Me-O–H. Implications of the vibrational and rotational photofragment distributions for the full 1 Å surface are discussed.

INTRODUCTION

The photochemistry of gaseous methanol, at both infrared and ultraviolet wavelengths, has received considerable attention. This is due in part to the practical significance of methanol, which ranks tenth in U.S. production of organic chemicals, with 3.8×10⁹ kg produced in 1992. In addition, the relatively small number of electrons in methanol makes it amenable to high-level computational studies. In infrared multiple photon dissociation (IRMPD) studies, it was found that, regardless of excitation frequency, the dominant primary products were the same as those seen in thermal excitation, namely, CH₃+OH. This is not surprising, since after the molecule has absorbed a number of infrared photons, further photoexcitation results in nonselective vibrational “heating” of the molecule within its ground electronic state until the lowest energy dissociation channel opens.

However, it is well established from end-product analyses as well as the more recent nascent hydrogen atom detection study by Satyapal et al., that the ultraviolet photolysis of methanol in its first absorption band (S₁→S₀) between 165 and 200 nm yields CH₃O+H as the dominant channel

\[ \text{CH}_3\text{OH} \rightarrow \text{CH}_3\text{O}+\text{H} \quad \Delta H_0 = 105 \text{ kcal mol}^{-1}. \]

Satyapal et al. estimated that reaction (1) accounts for 86% ±10% of the products at a photolysis wavelength of 193.3 nm, in agreement with the relative yield of this channel as determined by end-product analysis of methanol photolyzed with the 184.9 nm line of a Hg lamp. Reaction (1) is significantly more endothermic than the dominant thermal process

\[ \text{CH}_3\text{OH} \rightarrow \text{CH}_3+\text{OH} \quad \Delta H_0 = 90 \text{ kcal mol}^{-1}. \]

All primary products thought to be involved in the S₁ photochemistry of methanol are listed in Table I along with their respective reaction endothermicities. We note that the accurate determination of the bond dissociation energy, D₀(CH₃O-H), or equivalently the methoxy heat of formation, has been the subject of extensive investigation. This value is important because of the central role methoxy plays in combustion and atmospheric chemistry.

In addition to estimating the relative yield of the CH₃O+H channel, Satyapal et al. also reported that 82% of the available energy appeared as product translation from interpretation of Doppler line shapes of nascent H atoms. From the high translational energy release and the dominance of a thermodynamically disfavored process, it was concluded that the first absorption band of methanol accesses a potential energy surface (PES) that is repulsive in the O–H coordinate.

Prior to the study by Satyapal et al., several spectroscopic investigations assigned the broad, featureless first absorption band in simple alkyl alcohols to an n→π* transition in which a lone pair electron on the oxygen is excited into an antibonding orbital oriented in the plane defined by the carbon, oxygen, and hydroxy hydrogen. While this assignment is consistent with the small or absent C–H cleavage channel, it does not provide a rationale for the observed propensity for O–H bond scission over C–O bond scission. Moreover, other spectroscopic studies have assigned this ab...
TABLE I. Summary of enthalpy of reaction data for possible primary methanol photodecomposition channels.

<table>
<thead>
<tr>
<th>Reaction products</th>
<th>$\Delta H_0$(kcal mol$^{-1}$)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_3$O+2H</td>
<td>123.6</td>
<td>9</td>
</tr>
<tr>
<td>CH$_3$O+H</td>
<td>105</td>
<td>10-12</td>
</tr>
<tr>
<td>CH$_3$OH+H</td>
<td>95</td>
<td>11-13</td>
</tr>
<tr>
<td>CH$_3$+OH</td>
<td>90</td>
<td>14</td>
</tr>
<tr>
<td>CH$_3$O+H$_2$</td>
<td>20.3</td>
<td>9</td>
</tr>
</tbody>
</table>

*Estimated uncertainties are $\pm 2$ kcal mol$^{-1}$. 

The experimental work on methanol photochemistry has been complemented by several theoretical studies. There is agreement that irradiation in the first absorption band excites methanol into the $3s$ Rydberg level, and that the corresponding $1^1A''$ hypersurface is purely repulsive along the O–H coordinate, with no minima from the valence region to products. Moreover, there is no inconsistency between Rydberg excitation and repulsion along the O–H coordinate because the $3s$ Rydberg character in the valence region correlates with $\sigma^*$ character in the product channel through a process termed deRydbergization. In addition, there is consensus that the $1^1A''$ surface does not correlate to ground state CH$_3$OH+H products, and therefore this channel is thought to be unimportant. A similar situation occurs for the molecular elimination channel that yields CH$_3$O+H$_2$, in that the $1^1A''$ surface correlates to excited formaldehyde, though a dissociation path with a relatively small barrier was calculated by Buenker et al. 

Though all calculations show that the $1^1A''$ surface correlates with ground state CH$_3$+OH, there is less agreement on whether there is a barrier for this process. The most recent calculation of this surface by Marston et al. found that though both the C–O and O–HII coordinates are repulsive, the Franck–Condon region lies on the CH$_3$O+H side of the saddle point separating the two product valleys, as shown in Fig. 1. Following vertical excitation, methanol molecules near the equilibrium geometry for the ground electronic state therefore face a barrier to C–O bond scission while encountering no barrier to O–H bond breaking. Indeed, wave packet calculations on this surface indicate that excitation from the vibronic ground state leads exclusively to formation of CH$_3$O+H, while significant production of CH$_3$+OH requires at least 5 quanta of C–O stretch to overcome the barrier in the C–O coordinate. As noted above, experimental evidence on the role of the CH$_3$+OH channel is somewhat equivocal.

Furthermore, the wave packet dynamics calculations of Marston et al. indicate that the dissociation of methanol via the first excited PES “proceeds in a vibrationally adiabatic way.” In other words, $S'_1\rightarrow S_0$ excitation of methanol from its vibronic ground state should lead to formation of the vibrational ground state of methoxy, while excitation of metha-
the TOF distance was 33.9 cm for all except some of the
nol having one quanta of HO–CH₃ stretch should yield a
methoxy radical with one quanta of O CH₃ stretch, and so
forth. To test this prediction, we measured the translational
energy release of H atoms from the photodissociation of jet-
cooled methanol in its first absorption band, utilizing the high-n Rydberg time-of-flight (HRTOF) technique. We note
that translational energy spectra, if obtained with sufficient
resolution, carry useful information about the PES upon
which the dissociation proceeds, and therefore experimental
results can provide benchmarks for comparisons with theory.

EXPERIMENT

Our present experimental implementation of the HRTOF
 technique, which was pioneered by Welge and co-workers
 has been described previously. Briefly, a
molecular beam containing the sample was photolyzed with the
output from a 193 nm ArF excimer laser (Questek 2820),
which was focused by a 1 m, lens and collimated by an iris
before entering the vacuum chamber. Polarization was
achieved by using eight quartz plates placed in the path of the
ArF beam at Brewster’s angle. Radiation from a XeCl
excimer laser (Lambda Physik EMG 201 MSC) was split
equally to pump two dye layers. The 364.70 nm radiation
from one dye laser (Lambda Physik 3002) was focused into a
Kr cell to produce 121.57 nm Lyman-α radiation via fre-
quency tripling. The 121.57 nm radiation is focused into
the interaction region by a MgF₂ lens. H and D atoms were
excited by tuning the vuv radiation to the Lyman-α transi-
tions for H and D atoms, which are separated by 21.4 cm⁻¹.
To reduce ionization caused by residual 364.70 nm radiation,
the combination of lenses was arranged in such a way that
the 121.57 nm radiation was focused in the interaction re-
region, while the 364.70 nm fundamental was defocused. De-
tection and alignment of the vuv radiation were accom-
plished by using a photoionization cell with an adjustable
entrance iris and xyz positioner. This cell was connected to
the reaction chamber with flexible bellows. The vuv radi-
ad was estimated to have 10⁸ photons/pulse.

The output from the second probe laser (Lambda Physik
2001), which counterpropagates with the vuv radiation, fur-
ter excites the hydrogen electron from the 2p orbital to a
high-n Rydberg orbit. These high lying Rydberg states are
optically metastable and thus drift with their nascent veloci-
ties until they pass through a wire mesh into a region with a
moderate electric field (~1800 V/cm). The excited H atoms
are rapidly and efficiently field ionized, then counted. The
detector in these experiments was a Chevron MCP (Galileo
Electro-Optics). The reception angle of the detector was 10⁻⁴
steradians with respect to the laser interaction region, while
the TOF distance was 33.9 cm for all except some of the
related H/D studies, where it was 24 cm. Ion signals were
determined from the FWHM of the
peaks in HRTOF spectrum from 193 nm HBr photolysis.

A 3% CH₃OH/He mixture was obtained by bubbling He
through methanol at a total pressure of 400 Torr. The low
stagnation pressure was chosen to avoid formation of weakly

bound complexes, which are known to affect translational
energy release spectra. This mixture was expanded into
the source chamber through a 500 μm diameter pulsed
nozzle (General Valve), operating at a repetition rate of 10
Hz with a pulse width of 500 μs. The molecular beam was
collimated by a 1 mm diameter skimmer located 2 cm from
the orifice; the laser beams crossed the molecular beam at a
point 5 cm downstream from the skimmer. The molecular
beam was monitored by a quadrupole mass spectrometer
(UIT100C). CH₃OH, CH₃OD, and CD₂OH (Alrich
were used without purification. For deuterated samples, the man-
ufacturer specifies an isotopic purity of 99.5+ percent.

RESULTS AND DISCUSSION

TOF spectra of nascent H and D atom photofragments
obtained from photolysis of CH₃OD are shown in Fig. 2.
Because the D atom signal is structured, we are able to de-
finitively assign it to the primary photodissociation channel
yielding CH₃O+D, as discussed below. The small H atom
signal, however, is broad and structureless except for the
small number of H atoms that appear at early arrival times
(i.e., 18–20 μs). This part of the spectrum has the same
qualitative structure as the D atoms, indicating that this sig-
nal derives from isotopic exchange of the chemically labile
hydroxyl D in the sample line. Apart from the signal that
results from isotopic contamination, the potential sources for
the H atom TOF spectrum from CH₃OD include the two
primary processes

\[ \text{CH}_3\text{OD} \rightarrow \text{CH}_3\text{OD} + \text{H} \quad \Delta H_0 = 95 \text{ kcal mol}^{-1}, \quad (3) \]
\[
\begin{align*}
\text{CH}_3\text{OD} & \rightarrow \text{CH}_3\text{O} + \text{D} \\
\text{CH}_3\text{O} + \text{H} & \rightarrow \text{CH}_2\text{O} + \text{H} + \text{D} \\
\Delta H_f &= 124 \text{ kcal mol}^{-1}.
\end{align*}
\]

Energy conservation eliminates channel (4) for the portion of the H atom signal arriving before 24.5 μs (i.e., this arrival time corresponds to \(E_{\text{c.m.}} = 24\) kcal mol\(^{-1}\) for \(\text{CH}_3\text{O} \rightarrow \text{CH}_2\text{O} + \text{H}\)). Moreover, the H atom signal does not change markedly in this region, as would be expected if the threshold of an important channel is reached. Furthermore, as discussed below, few if any of the nascent methoxy have sufficient internal energy to decompose to formaldehyde plus hydrogen. From this, we speculate that channel (4) does not contribute appreciably to the H atom signal. The energetic threshold for channel (3) does not correspond to the onset of the observed H atom signal. However, as noted above, theoretical studies on this system,\(^7\)\(^{21}\)\(^{22}\) as well as some of the experiments,\(^4\)\(^7\) indicate that this process is unlikely.

This leads us to consider possible secondary photolysis channels, whereby a photofragment absorbs another photon from the same laser pulse in which it was created. This photofragmented reaction product then fragments further, leading to production of more H atoms. Photolysis fluence dependences were not measured in these experiments. We note that while observation of a nonlinear photolysis fluence dependence can confirm that a process is not a “single photon process,” the converse is not true. That is, observation of a linear dependence does not prove that the observed signal arises from a one-photon process.

From what is known about the photochemistry of methanol, the following secondary photolysis channels seem plausible:

\[
\begin{align*}
\text{CH}_3\text{O} & \rightarrow \text{CH}_2\text{O} + \text{H} \quad \Delta H_f = 20 \text{ kcal mol}^{-1} \quad (5) \\
\text{CH}_2\text{O} & \rightarrow \text{CHO} + \text{H} \quad \Delta H_f = 87 \text{ kcal mol}^{-1} \quad (6) \\
\text{CH}_3 & \rightarrow \text{CH}_2 + \text{H} \quad \Delta H_f = 109 \text{ kcal mol}^{-1} \quad (7)
\end{align*}
\]

Nascent photofragments may contain appreciable rovibronic excitation and thus energy from the first photon may appear as translational energy in a secondary photolysis channel. Because of this, any combination of these processes could in principle account for the observed H atom signal. Satyapal et al.\(^8\) also observed D atoms from \(\text{CD}_3\text{OH}\) and found that this signal had a nonlinear photolysis fluence dependence. They tentatively attributed the D atoms to photolysis of \(\text{CD}_3\).

We now turn to the structured part of the spectra. Arrival time spectra of H atom photofragments from \(\text{CH}_3\text{OH}\) photodissociation are shown in Fig. 3. Spectrum (a) was taken with unpolarized 193 nm radiation, while (b) and (c) were obtained with horizontally and vertically polarized radiations, respectively. The spatial arrangement of the molecular beam, polarized photolysis radiation, and the detector is indicated in the inset. We estimate that the polarization ratio achieved with the Brewster plates is \(\sim 10:1\), and therefore part of spectrum (c) is due to residual horizontally polarized radiation. This is consistent with the transition dipole moment being perpendicular to the line of departure of the hydrogen that derives from O–I\(\text{I}\) bond rupture. The molecule dissociates on a time scale which is rapid compared to that of molecular rotation (\(\sim 1\) ps). We estimate a spatial anisotropy parameter \(\beta = -0.7\), in reasonable agreement with Satyapal et al.,\(^8\) who reported \(\beta = -0.60 \pm 0.03\).

Translational energy distributions derived from HRTOF spectra of the three methanol isotopomers are shown in Fig. 4; structure is due to methoxy internal excitation. The maximum translational energies are 42.5 ± 1.42.5 ± 1, and 41.2 ± 1 kcal mol\(^{-1}\) for \(\text{CH}_3\text{O} + \text{H}, \text{CD}_3\text{O} + \text{H}, \text{and CH}_3\text{O} + \text{D}\), respectively. Figure 4 displays probability density versus the quantity \((1 + m_H/m_{\text{CH}_3\text{O}})E_H\), where \(E_H\) is the lab frame hydrogen translational energy (deuterium replaces hydrogen where appropriate). Values for c.m. translatinal energies are obtained by adding to this the quantity \((m_H/m_{\text{CH}_3\text{O}})E_{\text{parent}}\), where

\[
\begin{align*}
\text{H} & \rightarrow \text{OCH}_3 \quad (a) \\
\text{H} & \rightarrow \text{OCD}_3 \quad (b) \\
\text{D} & \rightarrow \text{OCH}_3 \quad (c)
\end{align*}
\]
$E_{\text{trans}}$ in the lab frame; again, deuterium replaces hydrogen where appropriate. This term accounts for the velocity of the molecular beam. The resulting c.m. values are $42.9 \pm 1$, $42.9 \pm 1$, and $42.0 \pm 1$ kcal mol$^{-1}$ for CH$_3$O+H, CD$_3$O+H, and CH$_3$O+D, respectively.

If these maximum translational energies correspond to ground state methoxy, the corresponding bond dissociation energies can be determined. With this assumption, our results yield bond dissociation energies for CH$_3$O–H, CD$_3$O–H, and CH$_3$O–D of 105$\pm$1, 105$\pm$1, and 106$\pm$1 kcal mol$^{-1}$, respectively, in accord with recent measurements and calculations. The larger CH$_3$O–D bond energy is due to the change in zero-point energies upon deuteration.

The data presented in Fig. 4 show that translation accounts for a large fraction of the available energy. This is in agreement with the results of Satyapal et al., who reported that the average c.m. translational energy in this channel is $82\%$ of the available energy. However, our results reveal information on how the CH$_3$O internal energy is distributed.

The spectra in Figs. 4(a) and 4(b) evince a progression, with the first three maxima spaced by 950$\pm$100 cm$^{-1}$. We assign this to the methoxy v$_3$–CH$_3$ stretch based on the following: (i) literature values for v$_3$ include assignments in fluorescence at 1045$\pm$10 cm$^{-1}$ and a calculated value of 1064 cm$^{-1}$; (ii) the spacing of the observed progression is changed very little by deuteration of the methyl group, ruling out significant methyl hydrogen motion; and (iii) previous studies indicate that initial excitation is localized on the oxygen and dissociation proceeds rapidly on an excited PES. We note that the S$_1$$\rightarrow$S$_0$ photophysics of methanol is reminiscent of the first absorption band of CH$_3$SH, studied previously in our laboratory. In both cases, dissociation of the X–H (X=O,S) bond occurs on a time scale $\leq 1$ ps. However, methoxy O–CH$_3$ stretch excitation following 193 nm photolysis is more extensive than thiomethoxy S–CH$_3$ stretch excitation following 248 nm photolysis of methyl mercaptan.

The O–CH$_3$ vibrational progression seen in Figs. 4(a) and 4(b) is also apparent in the photodissociation of CH$_3$OD, as shown in Fig. 4(c), but the peaks are more poorly resolved. This is attributable to the greater impulse force imparted to the methoxy by a recoiling deuterium as compared to hydrogen. An estimate of the rotational excitation caused by the departing atom can be made by using the "impulsive model." Briefly, the CH$_3$ moiety is taken as a structureless particle and the Me–O–H bond angle, $\alpha_0$, is assigned the ground state equilibrium value of 108$^\circ$. This yields

$$\frac{E_{\text{rot}}}{E_{\text{avail}}} = \frac{M_HM_{\text{CH}_3}}{(M_H+M_O)(M_{\text{CH}_3}+M_O)-M_HM_{\text{CH}_3}} \sin^2 \alpha_0 - \cos^2 \alpha_0.$$

Equation (8) gives values for $E_{\text{rot}}/E_{\text{avail}}$ of 0.026 and 0.049 for H and D, respectively. Inspection of Figs. 4(a) and 4(b) reveals that the energy spacing from threshold to maximum for the first peak in the progression is roughly 600 cm$^{-1}$ or 0.04 of the available energy. For the case of CH$_3$O+D shown in Fig. 4(c), the first peak does not display a distinct maximum, but it appears to be shifted from the origin by $\approx 1100$ cm$^{-1}$, or 0.08 of the available energy.

The impulsive model predicts the relative increase in rotational excitation upon D/H substitution at the oxygen, but underestimates the amount of rotational excitation. Though choosing $\alpha_0=108^\circ$ was arbitrary, setting $\alpha_0=90^\circ$ to give the maximum rotational excitation results in only a modest increase in the predicted $E_{\text{rot}}/E_{\text{avail}}$, i.e., from 0.026 to 0.028 and from 0.049 to 0.053 for H and D, respectively. Agreement cannot be obtained by adjusting $\alpha_0$. One explanation for the underestimate of methoxy rotational excitation is that torques deriving from the angular part of the excited PESs are neglected. This is a known limitation of the impulsive model. Such torque can either decrease or increase product rotation, as has been demonstrated for the cases of CINO and CICN, respectively. Thus the 1$^1\Lambda^+$ PES may exert torque that increases methoxy rotation.

The present experimental resolution of $\sim 170$ cm$^{-1}$ is too low to provide detailed information about methoxy rotational distributions. Furthermore, we cannot rule out the possibility that unresolved features buried within the O–CH$_3$ progression conceal additional vibrational excitation in modes other than v$_3$. However, if we concern ourselves only with this vibration, the extent of v$_3$ excitation in nascent methoxy can be estimated by simulating the data presented in Fig. 4. Specifically, the following expression was used to represent the translational energy distributions of each v$_3$ level

$$P(E) = P_0 N_v (E-E_v)^3 \exp\left(-\gamma_v (E-E_v)\right),$$

where $N_v = 6 \gamma_v^{-4}$ is a normalization factor and $\gamma_v$ is an adjustable parameter that varies little from one vibrational level to another. It was found empirically that the $\gamma_v$ values for entries (a) and (b) are essentially the same for all v (i.e., $\gamma_v=1/200$ cm$^{-1}$), while the $\gamma_v$ values for entry (c) are the same for all v, but are smaller than those for (a) and (b) by a factor of 2/3 (i.e., $\gamma_v=1/300$ cm$^{-1}$). We attach no special significance to Eq. (9). It fits the data (see Fig. 5) and may reflect the CH$_3$O rotational distribution. However, it does not provide a unique fit.

The $P_v$ values obtained from the fits shown in Fig. 5 are presented in Table II. Despite the latitude in deriving $P_v$ values, the following features are robust: (i) the O–CH$_3$ stretch vibration is inverted, with v = 1 having the largest population and measurable population to at least v = 4; (ii) the average CH$_3$O vibrational energy is approximately 1700 cm$^{-1}$, independent of isotopic substitution. This amount of vibrational excitation is considerably greater than that seen following excitation of CH$_3$SH in its first absorption band at 248 nm. In that case, the thiomethoxy internal energy distribution peaked at $v=0$. We also note that at $v=3$, the CD$_3$O+H spectrum appears more distinct than that of CH$_3$O+H. This was fit in the simulation with a somewhat broader rotational distribution. Alternate explanations include slight excitation in modes other than v$_3$, e.g., CH stretch, umbrella, and bend modes. Some excitation in these modes is expected, since the C$-$H bonds are longer and H$-$C$-$O bond angle is larger in methanol than in methanol.

As discussed in the Introduction, the methoxy state distributions reflect forces in photoexcited methanol as it
evolves on the excited PES. From our data, the dominance of the O–CH₃ stretch progression suggests that this system may be modeled as pseudotriatomic, with CH₃ treated as structureless. This is the approach employed by Marston et al. in their theoretical study of methanol photofragmentation.²³

Our translational energy release spectrum resembles the one-dimensional Franck–Condon progression often observed in electronic transitions. This suggests that such a model might provide insight.²⁶ Briefly, ground state MeOH is promoted to the excited PES, where it evolves. In addition to the repulsive force that breaks the O–H bond, the excited PES exerts forces that change the bond distances and angles in the newly forming methoxy. Projecting the excited PES wave function onto a methoxy basis subsumes all fragmentation processes for the dominant photofragmentation channel of methanol, CH₃OH+hν→CH₃O+H, were recorded by using the HRTOF method. Though most of the available energy appears as c.m. translation, a methoxy O–CH₃ stretch progression is observed which is peaked at v = 1. This is consistent with a significant change in this coordinate on the excited PES, relative to either ground state methanol or methoxy. Studies with the isotopic variants CD₂OH and CH₃OD reveal that photofragmentation of the latter yields significantly broader features. The extent of this broadening can be rationalized by consideration of the greater exit channel impulse imparted by D relative to H. The impulsive model somewhat underestimates the magnitude of methoxy rotation, as inferred from the rotational contour, suggesting that a torque is exerted by the angular part of the excited PES. H atoms observed from CH₃OD may be attributable to a multiphoton process, such as secondary photolysis of a primary photoproduct.

CONCLUSIONS

Vibrationally resolved c.m. translational energy distributions for the dominant photofragmentation channel of methanol, CH₃OH+hν→CH₃O+H, were recorded by using the HRTOF method. Though most of the available energy appears as c.m. translation, a methoxy O–CH₃ stretch progression is observed which is peaked at v = 1. This is consistent with a significant change in this coordinate on the excited PES, relative to either ground state methanol or methoxy. Studies with the isotopic variants CD₂OH and CH₃OD reveal that photofragmentation of the latter yields significantly broader features. The extent of this broadening can be rationalized by consideration of the greater exit channel impulse imparted by D relative to H. The impulsive model somewhat underestimates the magnitude of methoxy rotation, as inferred from the rotational contour, suggesting that a torque is exerted by the angular part of the excited PES. H atoms observed from CH₃OD may be attributable to a multiphoton process, such as secondary photolysis of a primary photoproduct.