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Vacuum ultraviolet photodynamics of the methyl peroxy radical studied by double imaging photoelectron photoion coincidences

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The vacuum ultraviolet (VUV) photoionization of the methyl peroxy radical, CH₃O₂, and unimolecular dissociation of internal energy selected $CH_3O_2^+$ cations were investigated in the 9.7–12.0 eV energy range by synchrotron-based double imaging photoelectron photoion coincidence (i²PEPICO). A microwave discharge flow tube was employed to produce CH_3O_2 via the reaction of methyl radicals (CH_3) with oxygen gas. After identifying and separating the different sources of CH₃⁺ from photoionization of CH₃ or dissociative photoionization of CH_3O_2 , the high resolution slow photoelectron spectrum (SPES) of CH_3O_2 was obtained exhibiting two broad bands superimposed with a complex vibrational structure. The first band of the SPES is attributed to the $X^{3}A''$ and $a^{1}A'$ overlapped electronic states of $CH_3O_2^+$ and the second is assigned to the b^1A' electronic state with the help of theoretical calculations. The adiabatic ionization energy (AIE) of CH_3O_2 is derived as 10.215 \pm 0.015 eV, in good agreement with high-accuracy theoretical data from the literature. The vertical ionization energy of the b¹A' electronic state is measured to be 11.5 eV and this state fully dissociates into CH_3^+ and O_2 fragments. The 0 K adiabatic appearance energy (AE_{0K}) of the CH₃⁺ fragment ion is determined to be 11.1548 ± 0.020 eV.

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I. INTRODUCTION

Organic peroxy radicals (RO₂) are key reaction intermediates in the low temperature oxidation of hydrocarbons and play an essential role in combustion and atmospheric chemistry.^{1, 2} In the atmosphere they are involved in the tropospheric "self-cleansing" mechanism, controlling the cycling of radicals and the formation of secondary pollutants such as ozone and secondary organic aerosols (SOA). The structure, thermochemistry and kinetics of RO₂ are of considerable importance in order to fully understand their chemistry in the atmosphere. In the past decades many advances have been made on the detection and analysis of RO₂ by the constant development of analytical methods such as absorption spectroscopy,^{1, 3} cavity ring down spectroscopy (CRDS),⁴ chemical ionization mass spectrometry (CIMS)⁵ and photoionization mass spectrometry (PIMS)^{6, 7}.

The methyl peroxy radical, CH_3O_2 , is the simplest and most abundant alkyl peroxy radical in the atmosphere which has attracted a great deal of attention. The structure of CH₃O₂ and its thermochemistry have been studied extensively both experimentally and with quantum-chemical methods. The neutral CH_3O_2 at the X^2A'' ground state has a Cs symmetry with electronic configuration of $(1a')^2 \dots (9a')^2 (2a'')^2 (10a')^2 (3a'')^{1.8}$ Removing an electron from the 10a' orbital in photoionization can lead to two spin cationic states, the $X^{3}A''$ ground electronic state and the $a^{1}A'$ first excited state. Theoretical calculations with a complete basis set method using the atomic pair natural orbital expansion (CBS-APNO) at the QCISD(T)/CBS level of theory were performed by Meloni et al. and predicted the adiabatic ionization energy (AIE) of CH₃O₂ at 10.21 eV.⁶ Voronova et al. structure calculations at the CCSD(T)/aug-ccalso carried out electronic pVQZ//CCSD/aug-cc-pVTZ level and predicted the same theoretical value of AIE = 10.21 eV.⁹

 $CH_3O_2^+$ cation and its kinetics were studied experimentally by mass spectrometry with electron impact ionization¹⁰ and vacuum ultraviolet (VUV) lamp photoionization.¹¹ Meloni *et al.* investigated the synchrotron VUV photoionization of small alkyl peroxy radicals using the multiplexed photoionization mass spectrometer (MPIMS) at the Advanced Light Source, and found that most of these cations are unstable, whereas $CH_3O_2^+$ is among the very few bound alkyl peroxy cations.^{6, 12} Using the photoionization

efficiency curve (PIE) of CH₃O₂ they determined the AIE to be 10.33 ± 0.05 eV, and the appearance energy of the CH₃⁺ fragment ion from the dissociative photoionization (DPI) of CH₃O₂ was reported as 11.16 ± 0.05 eV. The PIE of CH₃O₂ was also measured at Hefei synchrotron by using a VUV photoionization time-of-flight (TOF) mass spectrometer and its AIE was reported to be 10.32 ± 0.03 eV.⁷ Bernstein *et al.* also observed CH₃O₂⁺ and obtained the rotationally resolved infrared (IR) spectrum of CH₃O₂ by IR+VUV laser photoionization mass spectrometery.¹³

Building on the capabilities of photoionization mass spectrometry (PIMS) and photoelectron spectroscopy (PES), photoelectron photoion coincidence spectroscopy (PEPICO) analyzes both the photoelectrons and photoions in coincidence, and offers richer and more accurate information on energetics.¹⁴⁻¹⁶ Particularly, mass-selected threshold PES (ms-TPES) can be measured, which provides high-resolution thermochemical information with structure specific insights.^{9, 17, 18} Recently, Voronova *et al.* performed imaging PEPICO experiments at the Swiss Light Source (SLS) and reported the AIE of CH₃O₂ and the adiabatic appearance energy (AE_{0K}) of the CH₃⁺ fragment ion as 10.265 \pm 0.025 and 11.164 \pm 0.010 eV, respectively, by fitting the envelope of the TPES and the breakdown diagram.⁹

In this work, we present a detailed VUV photoionization and dissociative photoionization study of CH_3O_2 by using the high energy resolution of the VUV beamline DESIRS at synchrotron SOLEIL and the state-of-the-art method of double imaging photoelectron photoion coincidence (i²PEPICO).^{16, 19} The full slow photoelectron spectrum (SPES) of CH_3O_2 is acquired in the 9.7–12.0 eV energy range and assigned with the help of theoretical calculations, which are indispensable as the unfavorable Franck–Condon factors (FCF) combined with the proximity of the first electronically excited state render the measurement of the AIE difficult. The high photon energy resolution combined with the slow photoelectron technique provide a more precise comparison to the existing calculations leading to an increased accuracy of the AIE. In the following sections, we will first describe the experimental setup and methodology to generate CH_3O_2 using a microwave discharge flow tube. Then a procedure to separate the cold molecular beam and the thermal background gas

components within the photoionization region is introduced and used to get positionfiltered and mass-selected SPES.

II. EXPERIMENTAL SETUP

The experiments were performed at the VUV beamline DESIRS with the i²PEPICO spectrometer DELICIOUS III at synchrotron SOLEIL, France. A flow tube combined with a microwave discharge generator was employed as reactor to produce radicals and initiate their reactions. The configurations of the beamline, the i²PEPICO spectrometer and the flow tube have already been introduced before¹⁶⁻²¹ and only a brief description is given here.

On the DESIRS beamline, synchrotron photons emitted from a variable polarization undulator were dispersed by a 6.65 m normal incidence monochromator equipped with four gratings to cover the energy range of 5–40 eV.²⁰ In our experiments the 200 l.mm⁻¹ grating providing high photon flux was used and the photon energy resolution was set at \sim 3 meV. A gas filter filled with Ar was adopted to suppress high-order harmonics and the absolute photon energy was calibrated on-line with an accuracy of ±1 meV-using the resonant absorption lines of Ar.²²

The flow tube was composed of a main tube with an inner diameter of 2.5 cm and a collinear sliding injector, installed inside the source chamber of SAPHIRS, the permanent molecular beam endstation on the beamline.^{19, 21} The microwave discharge generator running at 2.45 GHz (Sairem GMP 03 KSM) was mounted perpendicularly to the gas flow and initiated radical reactions in the flow tube. The pressure inside the flow tube was measured by a gauge (Varian CDG 500) and maintained at 1–10 Torr by a closed-loop feedback throttle valve.²¹ After passing through the flow tube, the gas mixture was sampled and expanded through two skimmers (1 mm diameter) into the ionization chamber of SAPHIRS.

The gas mixture crossed the photon beam at the center of the i²PEPICO spectrometer.¹⁶ Electrons and ions produced in photoionization were extracted and accelerated in opposite directions towards an electron velocity map imaging (VMI)²³ device and an ion modified Wiley-McLaren TOF 3D-momentum imaging analyzer. A

position sensitive detector (PSD, DLD 80, Roentdek) was mounted to collect electron images which were filtered with ion mass- and momentum-selection in coincidence. An Abel inversion algorithm, pBasex, was used to analyze electron images to get PES or SPES.^{24, 25} A custom-made PSD (40 mm diameter) was used for ion detection whose 3D momentum distribution was determined by TOF and transverse position measurements.

III. RESULTS AND DISCUSSION

Two kinds of experiments have been performed to unravel the double origin of CH_3^+ observed in the experiment: photoionization of the methyl radical, CH_3 , and dissociative photoionization of CH_3O_2 . The first experiment was performed without adding oxygen into the flow tube. The methyl radicals were produced through the hydrogen-abstraction reaction (1) of fluorine atoms with methane (30 sccm, Air Liquide), where fluorine atoms were generated in the microwave discharge of a mixture of 5% F₂ diluted in helium (10 sccm, Air Liquide).

$$F + CH_4 \rightarrow CH_3 + HF \tag{1}$$

The second experiment was carried out by adding a large amount of oxygen into the flow tube (600 sccm, Air Liquide) to enhance the production of CH_3O_2 via the reaction⁷

$$CH_3 + O_2 + M \rightarrow CH_3O_2 + M$$
 (2)

where M represents molecules removing the internal energy of the nascent CH_3O_2 by collisions. For both experiments, helium was added as carrier gas and the overall flow rate was fixed at 1740 sccm with the pressure inside the flow tube at 1.6 Torr.

A. Time-of-Flight Mass Spectra and Ion Images

The photoionization mass spectrum in the absence of oxygen was measured at hv = 12.0 eV and is presented in Fig. 1(a). The most intense peak is observed at m/z = 15 and assigned to the methyl radical, produced from reaction (1). Minor peaks are also observed at m/z = 14, 26, 28 and 30 in the mass spectrum and are assigned to CH₂, C₂H₂, C₂H₄ and C₂H₆, respectively, generated from secondary reactions in the flow tube.

After the addition of oxygen, the photoionization mass spectrum at hv = 12.0 eV was measured and is presented in Fig. 1(b). Besides the above-mentioned peaks, another intense peak at m/z = 47 appeared which is assigned to CH₃O₂ produced from reaction (2). The more intense peak at m/z = 30 is attributed to two isobaric products, C_2H_6 and H_2CO by the associated mass-selected SPES (ms-SPES). The peak of m/z = 32 corresponds to O_2^+ from the residual high-order/high-harmonic photoionization of oxygen presented in high concentration in the flow tube.



FIG. 1. Synchrotron photoionization time-of-flight mass spectra measured at hv = 12.0 eV (a) without oxygen and (b) adding oxygen into the flow tube.

The ion images of CH₃⁺ and CH₃O₂⁺ were acquired with the time-focusing mode of DELICIOUS III¹⁶ and are displayed in Fig. 2, together with their intensity projections along the y-axis direction. In this mode, the final position on the detector is a convolution of the initial position and particle velocity. Parent ions, such as the ones shown in Fig. 2, possess narrow velocity distributions, especially for the ions formed in the molecular beam (MB), so that the dispersion seen in the ion positions is mainly related to the size of the interaction region. Parent ions formed in the MB will also have a net velocity along the MB's direction, as seen in Fig. 2, where MB propagates from east to west along the x-axis direction, and the propagation direction of synchrotron radiation (SR) is along the y-axis. Therefore, unlike VMI,²³ the time-focusing mode is utilized to map the photoionization region allowing the separation of ions from MB and the thermal background, plus the selection of on-axis ions to increase the ion mass and ion and electron kinetic energy resolution, which can be viewed as an adjustable virtual skimmer.^{16, 19}

The ion images and their intensity projections can be divided into two parts: the region of interest (ROI) from MB and the "Back" from the background gas in the photoionization region. In the ROI and as mentioned above, the ions from MB have a defined velocity along the MB direction and are therefore shifted from the image center.^{19, 21} For the Back, the ions from the thermal background gas do not have velocity along the MB direction and appear at the image center. With the coincidence scheme we can get the position-filtered electron and ion signals correlating only to the cold ROI or the thermal ions of a given mass.^{16, 21} Note that this scheme also reduces the false coincidences, because for an event to be considered the ion needs to arrive at the expected TOF and at the expected position, similar to the scheme reported by Sztáray *et al.*²⁶



FIG. 2. Mass-selected ion images and their intensity projections along the y-axis corresponding to (a, c) CH_3^+ and (b, d) $CH_3O_2^+$, recorded at hv = 12.0 eV, where the left ROI correlates to the part of molecular beam with a distance away from the image center, and the right Back part to the background in the photoionization region.

B. CH₃⁺ from Photoionization or Dissociative Photoionization

Mass-selected photoelectron kinetic energy matrices show the photoelectron signal as a function of photon energy and electron kinetic energy for a particular mass (presently only including the ROI part in Fig. 2).^{18, 27} From these matrices the SPES is extracted by integrating the photoelectron signal along a constant ionic state (diagonal line), up to a maximum value, KE_{max} , according to the expression,

$$SPES(h\nu) = \int_0^{KE_{max}} A(h\nu + KE, KE) dKE$$
(3)

where A(hv, KE) is the photoelectron kinetic energy matrix as plotted in Figure 3.

This allows both an implicitly correct hot electron subtraction, and a more relaxed compromise between resolution and statistics since larger values of KE_{max} are typically used with respect to TPES. The photoelectron kinetic energy matrices corresponding to the m/z = 15 ions without and with the addition of oxygen gas into the flow tube were measured with a photon energy step size of 10 meV and are presented in Fig. 3(a) and (c). In both photoelectron matrices, the electron signal appears at ~9.8 eV, and the electron kinetic energy increases linearly with photon energy in accordance with a direct photoionization of the methyl radical. But, after the addition of oxygen gas into the flow tube, further weak signals can be identified with small electron kinetic energy at the lower-right corner of Fig. 3(c), in the 11.2–12.0 eV photon energy range.

The ms-SPES corresponding to the m/z = 15 ions are obtained from the photoelectron matrices and presented in Fig. 3(b) and (d), with 10-times-magnified data in blue.²⁵ The ms-SPES records the signal of photoelectrons with near-zero kinetic energy (KE_{max} = 100 meV) and its energy resolution can be comparable to the traditional TPES.^{18, 27-29} As shown in Fig. 3, the two ms-SPES mainly exhibits the same structures and their intense peaks are located at hv = 9.838 eV, corresponding to the AIE of the methyl radical. In addition, several minor peaks can be observed close to the main peak which are ascribed to the v₂ umbrella vibrational excitations in the photoionization.^{30, 31} In Fig. 3(b), the electron signal at the high energy part of the ms-SPES is almost zero, as seen in the 10 times magnified data in blue, due to the minute resonant photoionization possibility of the methyl radical within this energy range³⁰.



FIG. 3. Mass-selected photoelectron kinetic energy matrices corresponding to CH_3^+ ions acquired (a) without and (c) with adding oxygen gas into the flow tube; (b) and (d) are the corresponding mass-selected slow photoelectron spectra (ms-SPES), with 10 times magnified data in blue.

However, as shown as the blue magnified data in Fig. 3(d), the ms-SPES acquired with adding oxygen gas is different from the one without oxygen in Fig. 3(b), with additional signal appearing with the addition of oxygen. Therefore, electrons corresponding to the mass of m/z = 15 in Fig. 3(d) have a double origin: below 11.1 eV CH₃⁺ and the corresponding electrons are produced by direct photoionization (PI) of the methyl radical, while above 11.1 eV CH₃⁺ is created due to DPI of CH₃O₂.^{6, 9}

C. SPES of CH₃O₂

The photoelectron kinetic energy matrix and the ms-SPES corresponding to the mass of m/z = 47 are presented in Fig. 4. The electron signal appears at ~10.2 eV and exhibits a broad band in the ms-SPES in the 10.2–11.1 eV energy range. The electron signal is almost zero at hv > 11.1 eV, due to the dissociative character of CH₃O₂⁺ prepared at high energy, as described above.



FIG. 4. (a) Photoelectron kinetic energy matrix and (b) mass-selected slow photoelectron spectrum corresponding to the mass of m/z = 47.

By combining the ms-SPES of m/z = 15 and 47 (in Fig. 5a), the full SPES of CH₃O₂ is retrieved, for the first time, in the 9.7–12.0 eV energy range and presented as a black solid line in Fig. 5(b). Previous high-accuracy calculations showed that the AIEs of the X³A" ground electronic state and the a¹A' first excited state of CH₃O₂⁺ are located at 10.21 and 10.43 eV, respectively,^{6, 9} and contribute to the first band of the SPES in the 10.2–11.1 eV photon energy range. However, due to the small energy difference between these two cationic states and their vibrational congestion, they are overlapped and cannot be separated in the SPES even with higher energy resolution.

Voronova *et al.* calculated the Franck–Condon factors involved in the photoionization and found that the strongest active vibrational modes are the OOC bend (320 cm^{-1}) and the CO stretch (505 cm^{-1}) for the X³A" ground electronic state, and the OOC bend (497 cm^{-1}) and the OO stretch (1578 cm^{-1}) for the a¹A' excited state.⁹ Voronova *et al.* used detailed Franck–Condon simulations to fit the TPES, in which the CH₃–OO torsional mode was treated as a hindered rotor and obtained an AIE of CH₃O₂ at $10.265 \pm 0.025 \text{ eV}$.⁹ However, as their experimental spectrum was of lower resolution and lacked a clear–vibrational structures, the fitting was mainly based on the overall envelope of the TPES. The main uncertainty in the AIE was attributed to the relative ratio of the singlet and triplet transition intensities and they found that Franck–Condon simulations with AIEs between 10.24 and 10.29 eV provided acceptable fits to the data.



FIG. 5. (a) Mass-selected slow photoelectron spectra (SPES) corresponding to m/z = 47 and 15 ions; (b) SPES of CH₃O₂ (black) and its partial Franck–Condon fitting (purple).

In comparison to the literature $TPES^9$ (see Fig. S1), owing to the high energy resolution of the DESIRS beamline and of the DELICIOUS III setup,^{16, 19, 20} the present SPES exhibits a defined vibrational structure, especially at the beginning of the first band. We have also tried to fit the overall SPES with the calculated Franck-Condon factors⁹ and cannot get a satisfying result, due to the two overlapped X^3A'' and a^1A' states with a complex congested profile. So, we focused on the fitting of the vibrational peaks at the beginning of the SPES and a satisfactory Franck-Condon fitting, taking into account the contributions of both overlapped states, is obtained for the well-resolved and intense vibrational peaks below 10.6 eV, as shown in Fig. 5(b). The vibrational peaks in this region, corresponding to the beginning of the first band, are mainly attributed to the CO stretching and the OOC bending modes of the X² A" ground state, and the OOC bending mode of the a⁺A' excited state. The best match between experimental and simulated spectra is achieved assuming an AIE of CH_3O_2 of 10.215 \pm 0.015 eV, in excellent agreement with the literature high-accuracy theoretical data of 10.21 $eV_{,-}^{6,9}$ and the error bar mainly includes the contributions of the energy uncertainty in the Franck-Condon fitting and the synchrotron photon energy resolution. The vibrational peaks in this <u>region</u>below 10.6 eV, corresponding to the beginning of the first band, are mainly attributed to the CO stretching and the OOC bending modes of the X^3A'' ground state, and the OOC bending mode of the a^1A' excited state, as shown the assignments in Fig. 5(b), whereas the two peaks with very weak intensities at hv = 10.11 and 10.17 eV in the SPES are ascribed to the hot bands involved in the photoionization.

Fig. S1 shows a comparison between the TPES of Ref. 9 and our SPES. In the region above ~10.7 eV the TPES band is larger which is somewhat attributed to the temperature differences between the two measurements as determined from the breakdown diagrams (*vide infra*). The internal temperature of the free radical was determined to be 330 ± 30 K at the SLS,⁹ whereas it was 160 ± 40 K in our experiment (see Section III.D).

The second band of the ms-SPES locatesed in the 11.1–12.0 eV energy range and its vertical ionization energy (VIE) lies at ~11.5 eV, as seen in Fig. 5. The second band is assigned with the aid of theoretical calculations as the second excited electronic state, $b^{1}A'$. Concretely, the geometry optimization and energy calculation of the ground electronic state of the neutral CH₃O₂ radical are performed at the DFT B3LYP/6-311G* level of theory, whereas those of the excited electronic states of the $CH_3O_2^+$ cation are at the TDDFT B3LYP/6-311G* level of theory. All these calculations are carried out by using the Gaussian 09 program package. The theoretical calculations show that the electronic configuration of the b¹A' cationic state is $(1a')^2 ... (9a')^2 (2a'')^2 (10a')^2$ and its VIE is theoretically calculated to belocates at 11.56 eV at the TDDFT B3LYP/6-311G* level,³² agreeing well with the experimental value of 11.5 eV. We also calculated the AIE (VIE) of the third excited electronic state $A^{3}A''$ at ~12.4 (14.612.9) eV with the same level of theory, *i.e.* beyond the present photon energy range, and hence it does not contribute to the second band. It is worth noting that we had also tried to calculate the AIE of the b^1A' state, but theour theoretical calculations aimed at optimizing the-its equilibrium structure of the $b^{+}A'$ state did not converge. This may indicate that the $b^{1}A'$ electronic state is not stable, agreeing well with our experimental results, since the $CH_3O_2^+$ cations prepared in the b¹A' state fully dissociate into the CH_3^+ and $O_2^$ fragments, as shown in Fig. 5(a).

Interestingly, unlike the literature TPES,⁹ in the present SPES there is almost no population in the Franck–Condon gap (~11.1 eV) between the two bands (see Fig. S1). This discrepancy could be explained by the temperature difference involved in the two experiments. The temperature difference could come from the fact that in our case we separate the weak expansion from the thermal gas using the ion ROI, as shown in Fig. 2, and only take into account the cold part of the molecular beam in the SPES. In addition, the different methods to produce the CH₃ radical could also contribute somewhat to the temperature difference. We used the reaction $F + CH_4 \longrightarrow CH_3 + HF$ to generate the CH₃ radical, since-whereas the literature TPES was obtained using photolysis to initiate the reaction.generate the precursor methyl radical, followed by low pressure expansion into vacuum which, in the absence of supersonic cooling, yielded a thermalized sample. Therefore, the internal temperature was nearly two times higher than with the present hydrogen-abstraction method. Another explanation for this discrepancy could be the different sensitivity to autoionization resonances of both methods, which are known to affect threshold spectroscopy in general, and leads to significant transition intensities even in wide Franck-Condon gaps. This latter feature of threshold photoionization (SPES and TPES) is routinely utilized in recording breakdown curves even in photon energy regions with no appreciable peaks in the photoelectron spectrum, or to expand the spectroscopic window into the Franck-Condon gap^{27, 28}.

D. Breakdown Diagram and the Adiabatic Appearance Energy

The X³A" ground electronic state and the a¹A' first excited electronic state are bound, as demonstrated in the ms-SPES of the m/z = 47 ions in Fig. 5(a). Earlier theoretical calculations showed that these two states adiabatically correlate to the CH₃⁺(X¹A₁) + O₂(X³ Σ_g^-) and the CH₃(X²A₂) + O₂⁺(X² Δ_g) dissociation limits, respectively.⁶

The fractional ion abundances obtained from the ms-SPES of m/z = 47 and 15 plotted as a function of the photon energy yield the breakdown diagram, which are shown in Fig. 6. As the photon energy increases, the energy distribution of the parent cation is shifted to higher energies, thereby leading to more dissociation. Finally, when the photon energy equals the appearance energy of the fragment ion, AE_{0K} of CH_3^+ here, the whole energy distribution is above the dissociation limit, and the ratio of the parent $CH_3O_2^+$ cation goes to zero.³³



FIG. 6. Breakdown diagram for $CH_3O_2^+$. Open symbols are experimentally measured ion abundances and lines are the statistical rate theory modeling of the data. The solid gray area represents the uncertainty in the ion temperature (160 ± 40 K), and the patterned area represents the uncertainty in the appearance energy of CH_3^+ fragment ions (11.148 ± 0.020 eV).

The CH₃O₂⁺ breakdown diagram is modeled as described by Voronova *et al.*,⁹ including a CH₃–OO hindered rotor in the energy distribution model for neutral CH₃O₂. In short, the 0 K appearance energy of CH₃⁺ and the internal temperature of CH₃O₂ were treated as fitting parameters. Since the dissociation is fast on the timescale of the experiment, the breakdown curves depend only on these two variables. Hence, the diagram can be used effectively as a molecular thermometer, similarly to the earlier work by Voronova *et al.*⁹ As these new experiments were optimized for a higher-quality photoelectron spectrum, the noise in the breakdown curve is more significant and the best fit model temperature and the 0 K appearance energy of CH₃⁺ is determined to be 160 ± 40 K and 11.1548 ± 0.020 eV, respectively. The AE_{0K} agrees with the 11.16 ± 0.05 eV determination of Meloni *et al.*⁶ and the 11.164 ± 0.010 eV by Voronova *et al.*⁹ Then, based on the thermochemical cycle, the bond dissociation energies of the parent cation CH₃O₂⁺ and the neutral radical CH₃O₂ are determined at D₀(CH₃⁺–OO) = 0.93 ± 0.02 eV and D₀(CH₃–OO) = 1.31 ± 0.02 eV, respectively, with the AIE of CH₃O₂ and the literature AIE of the methyl radical.

IV. CONCLUSIONS

The photoionization of CH₃O₂ and unimolecular dissociation of internal energy selected CH₃O₂⁺ cations were investigated in the 9.7–12.0 eV photon energy range by the state-of-the-art method of double imaging photoelectron photoion coincidence (i²PEPICO) at the DESIRS beamline of SOLEIL. The CH₃O₂ radicals were produced in a microwave discharge flow tube via the reaction of methyl radicals with oxygen molecules, where the methyl radicals were formed via the hydrogen-abstraction of methane. After identifying and separating the different sources of CH₃⁺ from photoionization of CH₃ or dissociative photoionization of CH₃O₂, the full SPES of CH₃O₂ was obtained with a resolution of ~10 meV and exhibits two broad bands superimposed with an intricate vibrational structure. The first band of the SPES is attributed to the X³A" and a¹A' overlapped electronic states while the second band is assigned as the b¹A' excited electronic states with the aid of theoretical calculations. The adiabatic ionization energy of CH₃O₂ is determined to be AIE = 10.215 ± 0.015 eV. The vertical ionization energy of the b¹A' electronic state is determined for the first time-and lies at around 11.5 eV.

 $CH_3O_2^+$ cations prepared at the b¹A' electronic state fully dissociate into CH_3^+ and O_2 fragments. The 0 K appearance energy of the CH_3^+ fragment ion was determined to be 11.1548 ± 0.020 eV by fitting the breakdown diagram with a CH_3O_2 internal temperature of 160 ± 40 K, which is significantly lower than the one obtained when CH_3 is produced by the hydrogen-abstraction of CH_4 with photolytically generated Cl. As demonstrated in this work, the i²PEPICO combined with the continuous microwave discharge flow tube has proven to be a valuable tool for providing high-accuracy thermochemical data together with rich dynamic information regarding the VUV photoionization and dissociative photoionization of radicals. Other peroxy radicals are being investigated on DESIRS and their high-resolution VUV photodynamics will be the subject of a forthcoming manuscript.

SUPPLEMENTARY MATERIAL

See the supplementary material for the slow photoelectron spectrum-SPES of CH₃O₂ obtained at SOLEIL and its threshold photoelectron spectrum-TPES at the Swiss Light Source (SLS) adapted from *J. Phys. Chem. Lett.* 9, 534–539 (2018).

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