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Investigation of nonadiabatic dynamics in the photolysis of methyl nitrate (CH_3ONO_2) by on-the-fly surface hopping simulation†

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The photolysis mechanism of methyl nitrate (CH_3ONO_2) was studied using the on-the-fly surface hopping dynamics at the XMS-CASPT2 level. Several critical geometries, including electronic state minima and conical intersections, were obtained, which play essential roles in the nonadiabatic dynamics of CH_3ONO_2 . The ultrafast nonadiabatic decay dynamics to the ground state were simulated, which gives a proper explanation on the broad and structureless absorption spectra of CH_3ONO_2 . The photodissociation channels, including $\text{CH}_3\text{O} + \text{NO}_2$, $\text{CH}_3\text{O} + \text{NO} + \text{O}$, and others, as well as their branching ratios, were identified. When the dynamics starts from the lowest two electronic states (S_1 and S_2), the $\text{CH}_3\text{O} + \text{NO}_2$ channel is the dominant photolysis pathway, although we observed the minor contributions of other channels. In contrast, when the trajectories start from the third excited state S_3 , both $\text{CH}_3\text{O} + \text{NO}_2$ and $\text{CH}_3\text{O} + \text{NO} + \text{O}$ channels become important. Here the $\text{CH}_3\text{O}-\text{NO}_2$ bond dissociation takes place first, and then for some trajectories, the N–O bond of the NO_2 part breaks successively. The quasi-degeneracy of electronic states may exist in the dissociation limits of both $\text{CH}_3\text{O} + \text{NO}_2$ and $\text{CH}_3\text{O} + \text{NO} + \text{O}$ channels. The current work provides valuable information in the understanding of experimental findings of the wavelength-dependent photolysis mechanism of CH_3ONO_2 .

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1. Introduction

Alkyl nitrates (RONO_2 , $\text{R} = \text{CH}_3$, C_2H_5 , etc) have received considerable research attention because they play an essential role in the NO_x -involved chemical reactions in the atmosphere.^{1–11} The oxidation and photoinduced reactions of reactive nitrogen oxides cause significant impacts on our atmospheric environments, such as affecting air quality, changing climates, and modifying nutrient cycles.^{1–4,7–13} A group of stable organic nitrates involved in the atmospheric NO_x reactions are RONO_2 . On one side, this series of compounds are generated by the photo-oxidation of hydrocarbons in the presence of NO_x or reactions of NO_3 with alkenes.^{1,2,6,11–16} Due to the thermal stability of RONO_2 , they may survive in the

atmosphere for a sufficiently long time and may be transferred to regions far from where they are produced. Therefore, several studies tried to explore their potential roles as NO_x reservoirs in the troposphere.² On the other side, RONO_2 experience photolysis and give NO_2 again after UV radiation.^{1,2,17–35} Therefore, the study of RONO_2 photoreaction at the atomic level is certainly important.

As the simplest alkyl nitrate, methyl nitrate (CH_3ONO_2) was found in marine boundary layer of the South Pacific, which was related to 20–80% of the reactive nitrogen (NO_x).¹⁶ The UV absorption spectra of CH_3ONO_2 ^{6,19,24,27,28,33,35–37} display an intense band around 190–220 nm, along with a much weaker diffusion band extending to around 330 nm. In fact, both are rather broad and structureless, namely, lack of vibronic progression. The broadness of the absorption band strongly indicates that the lifetime of the excited states should be extremely short.^{38,39} Therefore, after UV absorption, the wavepacket should quickly move out of the Franck–Condon (FC) region on the excited-state surfaces, and even decay *via* conical intersections (CIs). This implies that ultrafast excited-state dynamics in principle should take place.

Several experimental studies focused on the photolysis dynamics of CH_3ONO_2 and other alkyl nitrates.^{6,20,22–24,27,28,31,32,35,40} In particular, these works demonstrated that different photolysis products may be formed with laser excitation at different wavelengths. Yang *et al.*²⁰ found that the photolysis of CH_3ONO_2 at

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† Electronic supplementary information (ESI) available: I: Relevant information: the active space and transition orbitals, the geometrical parameters and state energies at S_0 -min, S_1 -min, and all CIs mentioned in this work, CI_{10} -seams, typical trajectories *via* different decay channels, and distribution of geometric parameters at S_1 to S_0 hops. II: The Cartesian coordinates of all critical geometries. See DOI: 10.1039/d1cp03226g

248 and 193 nm results in different dissociation products. The photolysis at 248 nm mainly gives $\text{CH}_3\text{O} + \text{NO}_2$ with a rather high quantum yield. At 193 nm two different channels were observed after the $\text{CH}_3\text{O}-\text{NO}_2$ bond breaking, in which the low-energy channel gives $\text{CH}_3\text{O} + \text{NO}_2$ products and the high-energy channel induces a further dissociation of the N–O bond in NO_2 , giving the $\text{CH}_3\text{O} + \text{NO} + \text{O}$ reaction. An additional channel $\text{CH}_3\text{ONO} + \text{O}$ was also identified. The UV photolysis of CH_3ONO_2 was examined by Talukdar *et al.*,³⁵ who suggested that the CH_3ONO_2 photodissociation at long wavelengths, namely 248 nm and 308 nm, is dominated by the $\text{CH}_3\text{O} + \text{NO}_2$ channel. At a much shorter wavelength of 193 nm, a visible quantum yield of the O atom was observed, which may come from the further dissociation of the NO_2 fragment. The photolysis of CH_3ONO_2 at a long wavelength (>300 nm) closing to the low-energy absorption tail always results in the $\text{CH}_3\text{O} + \text{NO}_2$ photoproducts. Derro *et al.*²⁴ examined the photolysis dynamics of CH_3ONO_2 at 193 nm. They suggested that the excitation of CH_3ONO_2 at 193 nm causes the local excitation of the $-\text{NO}_2$ moiety, and most available energy is thus partitioned to the NO_2 fragment during dissociation. As a consequence, after the $\text{CH}_3\text{O} + \text{NO}_2$ photoproducts are generated by the laser excitation at 193 nm, the highly excited NO_2 fragment may experience the further N–O bond dissociation, resulting in the $\text{CH}_3\text{O} + \text{NO} + \text{O}$ channel. Other alkyl nitrates and derivatives were also studied experimentally,^{6,17,21–24,27,28,31,32,35,40} which indicates a similar tendency.

Several theoretical efforts were made to examine the reactions of alkyl nitrates and similar compounds.^{5,18,25,26,41–49} The ground-state reactivity and the dissociation dynamics of CH_3ONO_2 were examined by electronic structure calculations and Born–Oppenheimer dynamics simulations.²⁵ One of the early theoretical studies on the excited states of alkyl nitrates was performed by Harris^{18,49} at the semi-empirical level, who provided the excitation energies and oscillator strengths of the low-lying excited states for the initial assignment of the absorption spectra of these species. Recently, Soto, Arenas and coworkers performed systematic studies on the photoreactions of CH_3ONO_2 and derivatives.^{26,41,43,48,50–52} At the multi-state complete active space second order perturbation theory (MS-CASPT2) level, they built the highly accurate potential energy surfaces (PESSs) of several excited states along the $\text{CH}_3\text{O}-\text{NO}_2$ dissociative coordinate.²⁶ The pathways towards different products were constructed. Several key geometries including the minimum-energy geometries and CIs were located. We noticed that the quasi-degeneracy of the low-lying electronic states exists in the dissociation limit due to the symmetry property of the photoproducts. This provides an important starting point to address the photolysis mechanism of CH_3ONO_2 .

From the theoretical point of view, the direct simulation of the excited-state dynamics of CH_3ONO_2 should provide valuable information for the understanding of the CH_3ONO_2 photolysis mechanism. Therefore, we tried to employ the on-the-fly trajectory surface hopping (TSH) dynamics with the extended multi-state complete active space second order perturbation theory (XMS-CASPT2)^{53–55} to study the excited-state nonadiabatic dynamics of CH_3ONO_2 . Several key geometries, including the

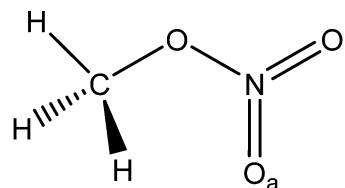
minima of electronic states and CIs, were located. The nonadiabatic dynamics including low-lying excited states was simulated. The results showed that the ultrafast nonadiabatic dynamics takes place after photoexcitation, followed by the successive dissociation dynamics. Different photoproducts were identified when dynamics starts from different excited states. The simulated results were in qualitative agreement with available experimental studies. This work deepens our understanding of the photochemistry of alkyl nitrates. In addition, we clearly demonstrate that both nonadiabatic internal conversion and photodissociation dynamics of CH_3ONO_2 take place within the ultrafast time scale. This indicates that such a system may become an interesting candidate in future experimental studies of ultrafast spectroscopy. At the same time, the theoretical treatment of photodynamics of volatile organic compounds (VOCs) becomes more and more attractive in recent years.^{1,2,56–60} We believe that the on-the-fly nonadiabatic dynamics^{56,58–67} may provide an indispensable simulation tool in the explanation of the photochemistry of VOCs.^{56,58–60,68,69}

2. Computational details

Scheme 1 shows the molecular structure of CH_3ONO_2 and atomic labels. The critical internal coordinates are listed in Table 1, in which the O–N, O–O_a, and O–O_b distances are described by r_a , r_b , and r_c , respectively. The distances of N–O_a and N–O_b are shown as r_d and r_e , respectively. The angle O_a–N–O_b (θ_a) describes the bending motion of the NO_2 fragment. The dihedral angle O–O_a–O_b–N (τ_a) describes the pyramidalization at the N atom.

The geometry optimization of the ground-state minimum (S_0 -min) and the vibrational analysis of methyl nitrate were performed at the B3LYP/6-311+G* level with the Gaussian 16 package.⁷⁰ The optimized S_0 -min geometry was used to calculate excited states using the XMS-CASPT2 with the def2-SVPD basis set. The XMS-CASPT2 calculations employed the active space of 12 electrons in 9 orbitals and four states were averaged, namely, XMS(4)-CASPT2(12,9). The active space contains three π , one σ and one n orbital, as well as one antibonding π^* orbital, two antibonding σ^* orbitals, one n/ σ mixed orbital. The orbitals in the active space are given in Fig. S1 in ESI.† The optimization of the S_1 minimum (S_1 -min) was also performed by the XMS(4)-CASPT2/def2-SVPD method with the same active space.

The ultrafast nonadiabatic decay dynamics simulations of CH_3ONO_2 were studied by the on-the-fly TSH method with Tully's fewest-switches algorithm⁷¹ at the XMS(4)-CASPT2(12,9)/def2-SVPD level. The initial geometries and velocities were



Scheme 1 Molecular structure of CH_3ONO_2 .

Table 1 Labels of important internal coordinates of CH₃ONO₂

Label	Internal coordinate
r_a	Distance O–N
r_b	Distance O–O _a
r_c	Distance O–O _b
r_d	Distance N–O _a
r_e	Distance N–O _b
θ_a	Angle O _a –N–O _b
τ_a	Dihedral angle O–O _a –O _b –N

sampled from the Wigner distribution function⁷² of the lowest vibrational state at the S₀-min. After the generation of many geometries by the Wigner sampling, their vertical excitation energies and oscillator strengths were calculated at the XMS(4)-CASPT2(12,9)/def2-SVPD level. The absorption spectrum was calculated by the nuclear ensemble approach (NEA),⁷³ in which 500 snapshots were taken, and the stick spectra were broadened by the Gaussian line shape with the phenomenological broadening parameter as 0.10 eV.

In the on-the-fly nonadiabatic dynamics simulation, the nuclear motion was propagated with a time step of 0.5 fs with the velocity-Verlet algorithm, and 100 steps of electronic motion were integrated within each step of nuclear motion. The decoherence correction approach proposed by Granucci *et al.*⁷⁴ was taken, and the parameter was set to 0.1 hartree.⁷⁵ Finally, 100, 99, and 98 trajectories were performed for the dynamics initialized from S₁, S₂, and S₃, respectively. All dynamics calculations were performed with the JADE-NAMD package.^{67,76,77} We developed the Python interface between the TSH dynamics in the JADE-NAMD package and the XMS-CASPT2 calculations in the BAGEL package.^{78,79} In the dynamics propagation process, the PESSs, nuclear gradients, and nonadiabatic coupling vectors were calculated directly at the XMS-CASPT2 level using the BAGEL package.

The S₁–S₀ (CI₁₀), S₂–S₁ (CI₂₁), and S₃–S₂ (CI₃₂) minimum-energy CIs (MECIs) were optimized at the state-averaged complete active space self-consistent field,^{80,81} SA(4)-CASSCF(12,9)/cc-pVDZ using the OpenMolcas package,⁸² in which the initial geometries were taken from hopping geometries in nonadiabatic dynamics. The single-point calculations were performed at those optimized CIs using the XMS(4)-CASPT2(12,9)/def2-SVPD level with the same setup as other XMS-CASPT2 calculations. We constructed the excited-state potential energy curves by the linear interpolation in internal coordinates (LIIC) approach from S₀-min to S₁–S₀ MECIs at the XMS(4)-CASPT2(12,9)/def2-SVPD level.

Furthermore, the S₁–S₀ CI seams (CI₁₀-seams) along the O–N distance (r_a) were also built at the SA(2)-CASSCF(12,9)/def2-SVPD level by the constrained optimization using the MOLPRO package.^{83–85} For convergence, only two-state averaged calculations were employed in the CI₁₀-seam construction. Although this only provided a preliminary view, the obtained CI-seam structure information is reasonable enough for our analysis purpose because we mainly focused on the CI₁₀ here.

3. Results

3.1. S₀-min and S₁-min

The molecular structures at the S₀ and S₁ minima of CH₃ONO₂ are shown in Fig. 1. The important geometrical parameters and excited-state energies at S₀-min and S₁-min are displayed in Tables S1 and S2 in ESI.† The geometrical structure of S₀-min shows C_s symmetry, where C, O, N, O_a, and O_b atoms are located in the same plane ($\tau_a = 0^\circ$). The distances of N–O_a (r_d) and N–O_b (r_e) are 1.21 Å and 1.20 Å, respectively. This geometry is similar to previous work.⁸⁶ At S₀-min, the XMS(4)-CASPT2(12,9)/def2-SVPD calculations show that the vertical energies of S₁ and S₂ are 4.84 and 5.76 eV, respectively. S₁ and S₂ correspond to the n (HOMO–1) → π* (LUMO) and n/σ (HOMO) → π* (LUMO) transition, respectively. Here the electronic character of the HOMO displays the mixture of n and σ, see Fig. S1 in ESI.† The orbitals involved in the electronic transition of S₁ are shown in Fig. S2(a) in ESI.† Both S₁ and S₂ show very weak oscillator strengths, while S₃ is a bright state. The vertical excitation energy of S₃ is 6.54 eV, which is dominated by the π (HOMO–2) → π* (LUMO) transition. The excitation energies of these states are similar to those of previous theoretical studies.^{26,28}

We calculated the absolute crossing section of the theoretical absorption spectra, including all individual contributions to different excited states using the NEA.⁷³ The absorption spectra in the broad energy range (from 180 nm to 320 nm) and in the long-wavelength domain (from 240 nm to 320 nm) are given in Fig. 2(a) and (b), respectively, along with the experimental ones.^{6,27} Please notice that all available experimental works only gave the absorption spectra above 190 nm.⁸⁷

As shown in Fig. 2(a), the theoretical and experimental absorption spectra agree well in the whole energy domain. The short-wavelength absorption band is dominant by the S₃ transition with very high oscillator strength. The profiles of theoretical and experimental bands near 190 nm are highly consistent, except that the theoretical peak intensity is a little lower. The electronic transitions to both S₁ and S₂ are much weaker, and their absorption intensities are several magnitudes lower than the transition to S₃. Thus, the contribution of the former two components cannot be seen directly in Fig. 2(a). If we examine the long-wavelength region (from 240 nm to 320 nm) in Fig. 2(b), the very weak long tail of the whole

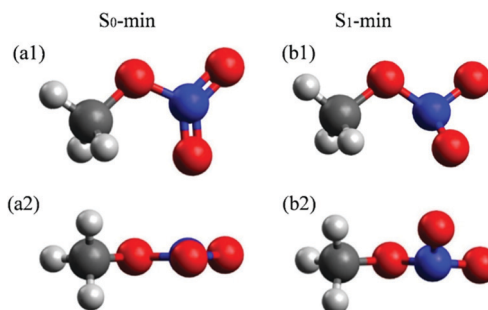


Fig. 1 Optimized S₀-min (B3LYP/6-311+G*) and S₁-min [XMS(4)-CASPT2(12,9)/def2-SVPD] geometries of CH₃ONO₂. The top and side views are shown up and down, respectively.

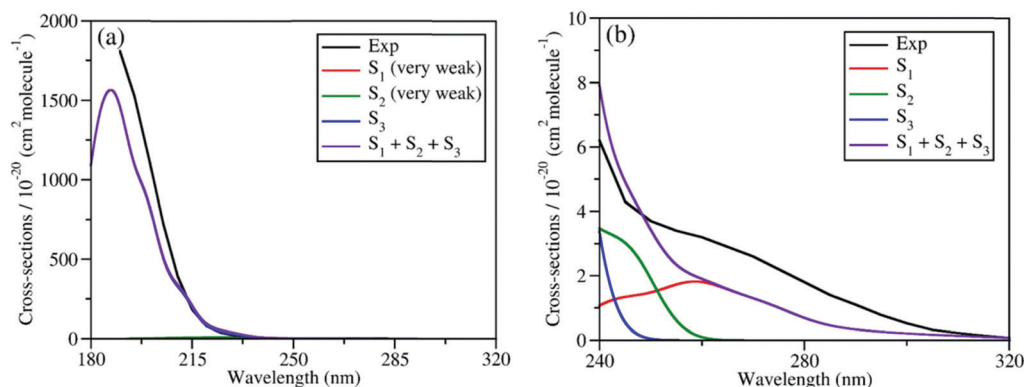


Fig. 2 Comparison of experimental and simulated UV absorption spectra of CH_3ONO_2 : (a) from 180 nm to 320 nm; (b) from 240 nm to 320 nm. The theoretical spectra were simulated at XMS(4)-CASPT2(12,9)/def2-SVPD level. The experimental spectra in (a) and (b) were taken from previous works.^{6,27} The intensities of absorption bands shown in (a) and (b) differ by several magnitudes.

absorption band is attributed to the summation of the transitions to both S_1 and S_2 . Their overall contribution basically agrees well with the experimental data, except that the theoretical band is slightly weaker.

Overall, the simulated spectra agree well with the experimental one in the broad energy domain, including peak position, intensity, and profile. In particular, the transition amplitudes to different excited states differ by several magnitudes, while the absolute cross sections in theoretical spectra agree well with available experimental data in all relevant energy domains. This confirms that the XMS-CASPT2 gives a reliable description of these low-lying excited states of the current system in the FC region.

In addition, this also indicates that the initial sampling of nuclear conditions used in the further TSH dynamics are reasonable.

In contrast to the S_0 -min geometry, the S_1 -min geometry displays a significant pyramidalization at the N atom ($\tau_a = 26.8^\circ$). With the pyramidal movement of the N atom, we observed the elongation of the $\text{N}-\text{O}_a$ ($r_d = 1.27 \text{ \AA}$) bond and the $\text{N}-\text{O}_b$ ($r_e = 1.32 \text{ \AA}$) bond, as well as the bending motion of the $\text{O}_a-\text{N}-\text{O}_b$ ($\theta_a = 109.0^\circ$) angle. This geometry deformation implies that the hybridization status of the N atom changes from sp^2 to sp^3 when the geometry changes from S_0 -min to S_1 -min. At S_1 -min, the relative energies of S_0 , S_1 , S_2 , and S_3 with

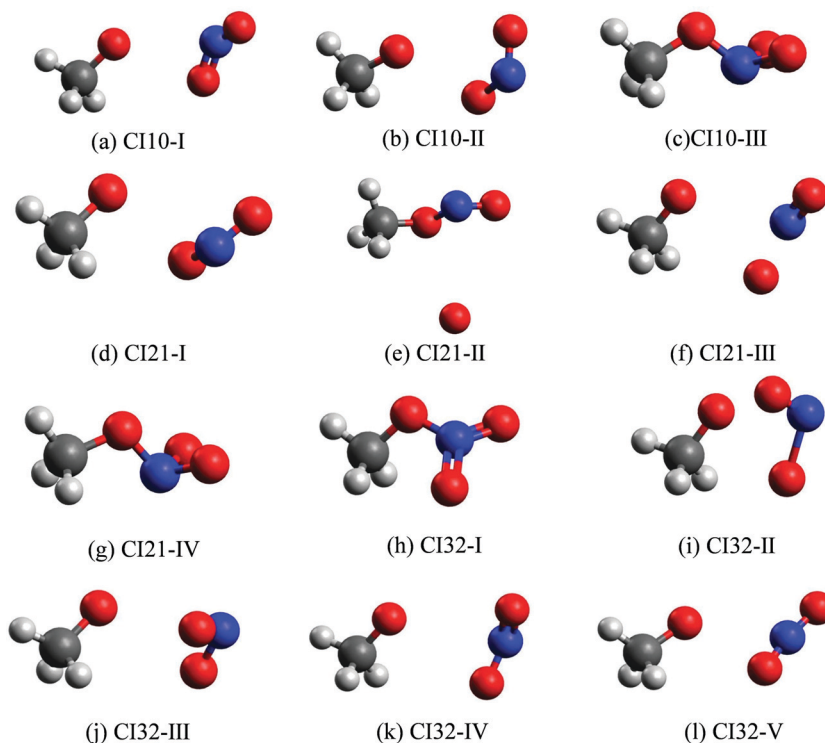


Fig. 3 The MEClIs optimized at the SA(4)-CASSCF(12,9)/cc-pVDZ level: (a)–(c): S_1 – S_0 Cls; (d)–(g): S_2 – S_1 Cls; (h)–(l): S_3 – S_2 Cls.

respect to the S_0 energy of S_0 -min are 2.06, 3.74, 5.54, and 7.27 eV, respectively. The transition of the S_1 state is n (HOMO) $\rightarrow \pi^*$ (LUMO), and the orbitals are shown in Fig. S2(b) in ESI.[†] We noticed that HOMO-1 at S_0 -min becomes HOMO at S_1 -min, and the same occupied orbital is always involved in the electronic transitions of S_1 , indicating that its electronic character remains unchanged from S_0 -min to S_1 -min.

3.2. Conical intersections

Fig. 3 shows all optimized MECIs, and Table S2 (ESI[†]) gives their corresponding energies. Normally, it is better to perform the dynamics simulation and the CI optimization at the same level of theories. However, several test CI optimization tasks at the XMS-CASPT2 level did not give us the converged results. To give a comprehensive picture of all involved MECIs and CI-seams, we took the CASSCF method to perform the CI optimization, since it is easier to obtain the converged results. To make sure that all obtained MECIs are reasonable, the single-point calculations at the XMS-CASPT2 level were conducted again. The results in Table S2 (ESI[†]) show that these MECIs optimized at the CASSCF level are reasonable because the energy gaps of electronic states involved in the state degeneracy are very small at the XMS-CASPT2 level.

The first minimum-energy CI_{10} is CI_{10} -I (1.91 eV) characterized by the significant CH_3O-NO_2 stretching motion ($r_a = 3.11$ Å) with respect to the S_0 -min geometry. The O_a-N-O_b bond angle ($\theta_a = 133.5^\circ$) at this CI is very close to that (130.0°) at the S_0 -min geometry. At CI_{10} -I, the system is roughly viewed as two radical fragments, CH_3O and NO_2 . The important frontier orbitals are composed of the p orbital located at the O atom of the CH_3O radical and the π orbital at the NO_2 side. Here the later π orbital is a delocalized one, which is the linear combination of p orbitals at O_a and O_b atoms in the NO_2 part, as shown in Fig. 4(a).

The second minimum-energy CI_{10} (CI_{10} -II) lies at a higher energy domain (3.92 eV). At CI_{10} -II, both the large CH_3O-NO_2 stretching and the O_b-N-O_a bond angle bending motion were observed, which are characterized by 3.19 Å and 110.3° , respectively. Here the frontier orbitals are still composed of the p orbital located at the O atom of the CH_3O radical and the p orbital at the O_b atom of the NO_2 side. Different from the CI_{10} -I,

the frontier orbitals of the NO_2 side become fully localized at the O_b atom, shown in Fig. 4(b). The underlying reason is easy to understand; the distances of $N-O_a$ and $N-O_b$ are very similar at CI_{10} -I. Two localized p orbitals associated with the O_a and O_b atoms show similar energies; thus, their electronic coupling results in the delocalized molecular orbitals, see Fig. 4(a). In contrast, the bond r_d and r_e of the NO_2 radical are quite different at CI_{10} -II. Therefore, the frontier orbitals become localized at the O_b atom, see Fig. 4(b).

At the third minimum-energy CI_{10} , labeled as CI_{10} -III, the O-N bond is in a bonded status and the strong pyramidalization at the N atom exists ($\tau_a = 34.8^\circ$). Different from CI_{10} -I and CI_{10} -II, as shown in Fig. 4(c), the frontier orbitals are localized in the NO_2 fragment.

We noticed that a previous study once localized a CI_{10} that is close to the CI_{10} -II here.²⁶ Another study showed that a similar system CH_3NO_2 has a CI that displays the bonded O-N bond and the pyramidalization at the N atom.⁵² This is similar to the CI_{10} -III in the current work.

We also located the MECIs between different excited states, namely CI_{21} and CI_{32} , and all obtained CI geometries are given in Fig. 3. The CI_{21} -IV and CI_{32} -I display the obvious pyramidalization at the N atom, and no bond breaking is observed at these two CIs, while the bond cleavage is observed at other CI_{21} and CI_{32} . More information about other CIs is listed in Tables S2 and S3 (ESI[†]).

3.3. Nonadiabatic dynamics

Nonadiabatic excited-state dynamics of CH_3ONO_2 were simulated using the on-the-fly TSH dynamics at the XMS-CASPT2 level. We first considered the dynamics starting from S_1 . After excitation to S_1 , the ultrafast $S_1 \rightarrow S_0$ population decay was observed, see Fig. 5(a). Within the first 18 fs, almost all trajectories were propagating on S_1 , and no decay was observed. After 53 fs, 50% of the trajectories jumped back to S_0 . At 120 fs, the S_1 population decayed to 23%. In the whole decay dynamics, both S_2 and S_3 did not play important roles here.

The current population dynamics indicates that most trajectories go back to the ground state and only a few trajectories stay on the excited state. Thus it is necessary to identify

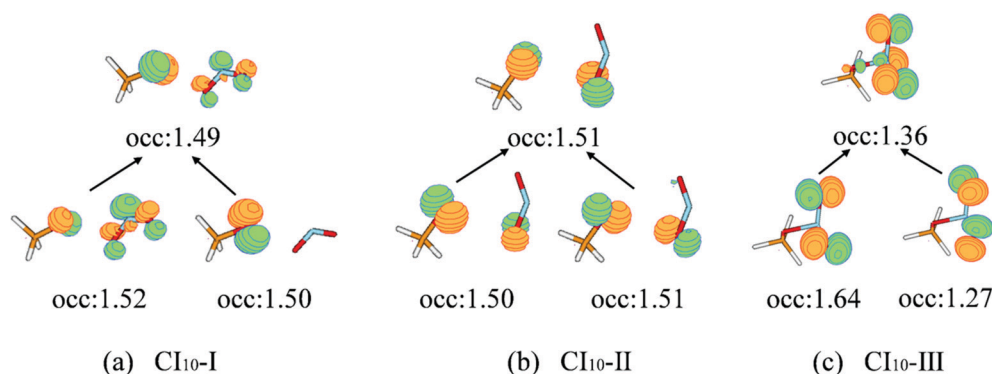


Fig. 4 The natural transition orbitals along with their fractional occupation numbers at the $S_1 \rightarrow S_0$ MECIs optimized at the SA(4)-CASSCF(12,9)/cc-pVDZ level.

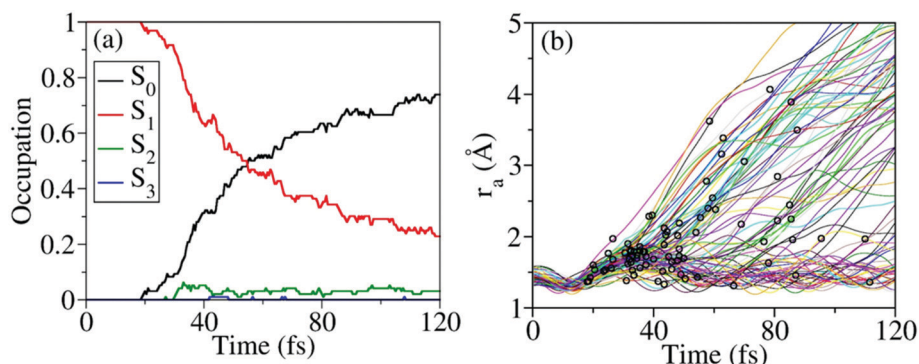


Fig. 5 Time-dependent (a) average fractional occupation and (b) r_a value of several trajectories simulated in the nonadiabatic dynamics starting from S_1 at the XMS(4)-CASPT2(12,9)/def2-SVPD level. Black circles mark the $S_1 \rightarrow S_0$ hops. After 120 fs, the electronic structure calculations of several trajectories may not be converged at random steps. Thus the overall population dynamics is only shown up to 120 fs.

which molecular motion is essential to drive the $S_1 \rightarrow S_0$ decay dynamics. From the trajectory propagation from the FC region to the $S_1 \rightarrow S_0$ hops, the obvious elongation of the $\text{CH}_3\text{O}-\text{NO}_2$ ($r_a = 1.31\text{--}4.24$ Å) bond and the significant pyramidalization at the N atom ($\tau_a = 7.2\text{--}170.1^\circ$) were noticed. After the $S_1 \rightarrow S_0$ internal conversion, most trajectories continue moving towards the dissociation limit, which results in the $\text{CH}_3\text{O}-\text{NO}_2$ bond cleavage, as shown in Fig. 5(b).

According to Fig. 5(b), most hops take place at geometries with a long $\text{CH}_3\text{O}-\text{NO}_2$ distance, while the system is still not fully broken. Thus the $\text{CH}_3\text{O}-\text{NO}_2$ bond strengthening motion should be one of the key degrees of freedom in the nonadiabatic dynamics. The S_1 - S_0 CI seams along the $\text{CH}_3\text{O}-\text{NO}_2$ bond length were built with the constrained optimization by freezing the $\text{CH}_3\text{O}-\text{NO}_2$ distance, as shown in Fig. 6.

The energies of three CI_{10} -seams as a function of r_a are shown in Fig. S3 (ESI†).

Both the CI_{10} -seam-I and CI_{10} -seam-II exist extensively in both short $\text{CH}_3\text{O}-\text{NO}_2$ distance and long $\text{CH}_3\text{O}-\text{NO}_2$ distance regions. We only obtained a small part of the CI_{10} -seam-III, and the other optimization jobs did not give the converged results. From the geometrical distribution at hops within 120 fs, many hops are governed by the CI_{10} -seam-III ($\sim 72\%$ of hops). Therefore, these hopping geometries do not show the full cleavage of the $\text{CH}_3\text{O}-\text{NO}_2$ bond. Instead, the strong pyramidalization at the N atom is observed. The other two CI_{10} seams result in the hop geometries displaying the longer $\text{CH}_3\text{O}-\text{NO}_2$ distances, and the ratio of hops around CI_{10} -seam-I and CI_{10} -seam-II is about 13% and 11% of all hops, respectively, assigned by the $\text{O}_a-\text{N}-\text{O}_b$ bond angles of these hops.

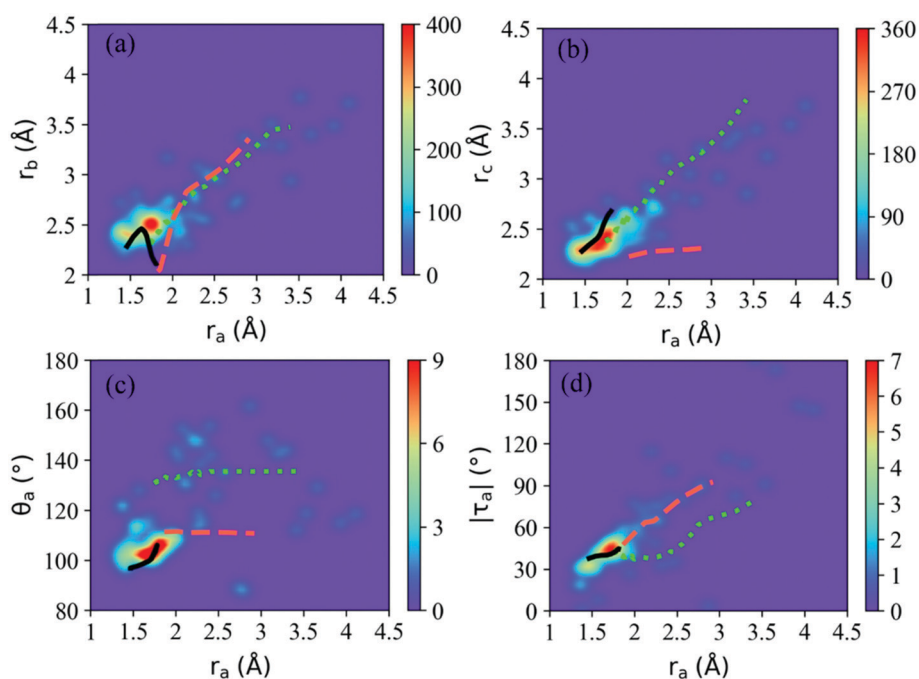


Fig. 6 Distribution of the important geometric parameters at the first S_1 to S_0 hops within 120 fs. In all panels, the green dotted line, orange dashed line, and black solid line mark the CI_{10} -seam-I, CI_{10} -seam-II, and CI_{10} -seam-III, respectively.

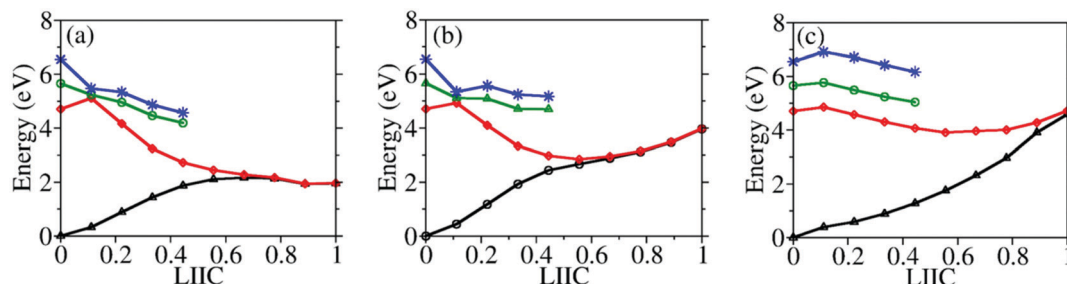


Fig. 7 Linear-interpolated PESs between S_0 -min and S_1 - S_0 MECIs in the FC region at the XMS-CASPT2 levels: (a) S_0 -min to CI_{10-I} ; (b) S_0 -min to CI_{10-II} ; (c) S_0 -min to CI_{10-III} . In all panels, the black line, the red line, the green line, and the blue line represent the S_0 , S_1 , S_2 , and S_3 states, respectively.

To understand why CI_{10-III} is the leading channel in the S_1 to S_0 nonadiabatic decay dynamics, the linear-interpolated PESs from S_0 -min to S_1 - S_0 CIs were constructed, as shown in Fig. 7. At the SA(4)-CASSCF(12,9) optimized MECI geometries, the single point calculations at the XMS-CASPT2 level also indicated that the lowest two states remain nearly degenerated. Thus the SA(4)-CASSCF(12,9) optimized CI geometries are reasonable. Although the energy of CI_{10-III} is higher, the reaction pathway from S_0 -min to it shows a minor barrier on the S_1 PES. Although CI_{10-I} and CI_{10-II} lie in the low-energy region, the pathways towards them display obvious barriers that may be caused by the existence of the CIs between different excited-states. In addition, the CI_{10-III} geometry is close to the FC region; this also means that the system may easily access the CI_{10-III} without undergoing drastic structural deformation.

After the analysis of the first $S_1 \rightarrow S_0$ hops, the next task is to examine the final photoreaction products. In fact, most trajectories follow the $CH_3O + NO_2$ dissociation pathway, no matter *via* which CI. Only for a few of the trajectories passing CI_{10-III} , the $CH_3O + NO + O$ channel was found, as shown in Table 2.

The typical trajectories towards different dissociation channels are given in Fig. S4–S7(a) (ESI[†]), along with the state degeneracy in the dissociation limit. These trajectories start from S_1 , experience the S_1 - S_0 hop *via* CI_{10-I} (Fig. S4, ESI[†]), CI_{10-II} (Fig. S5, ESI[†]), and CI_{10-III} (Fig. S6, ESI[†]), and move to the $CH_3O + NO_2$ channel at the end. No matter which CI is passed, the $CH_3O + NO_2$ dissociation limit always displays the quasi-degeneracy of two lowest states S_0 and S_1 . Although the trajectory starting from S_1 may generate the $CH_3O + NO + O$

channel, this channel becomes important in the dynamics initialized from the S_3 state. Thus the typical trajectory towards this channel in Fig. S7 (ESI[†]) starts from S_3 , in which all four lowest electronic states included in the state-averaged calculations become nearly degenerated in the $CH_3O + NO + O$ dissociation limit. The underlying reasons for the state degeneracy are relevant to the symmetries of molecular geometries and the degeneracy of frontier orbitals, and more details are given in the below section.

In a short summary, the ultrafast decay dynamics (the lifetime ~ 53 fs) was observed for CH_3ONO_2 when the trajectories start from S_1 , which is governed by three S_1 - S_0 CIs (CI_{10-I} , CI_{10-II} , and CI_{10-III}). Among them, CI_{10-III} plays a crucial role in the S_1 to S_0 hops. Most trajectories passing CI_{10-III} finally move towards two dissociation channels, in which the major channel is $CH_3O + NO_2$, and the minor channel is $CH_3O + NO + O$. The trajectories decaying *via* the other S_1 - S_0 CIs (CI_{10-I} and CI_{10-II}) give dissociation products $CH_3O + NO_2$. Overall, the primary photolysis channel is $CH_3O + NO_2$ and the ratio towards this channel is 83%. The minor channel gives the $CH_3O + NO + O$ products with a ratio $\sim 5\%$.

Please notice that the above ratio is only a rough estimation of the probabilities of photolysis product channels. The reason is as follows. When trajectories jump back to the ground state, the high kinetic energy results in distorted geometries. At some of these geometries, the electronic structure calculations become unstable, giving either the non-convergence results or the sudden flip of state energies. Therefore, we only considered the trajectory propagation before these questionable events. However, the geometries at these events may not be easily assigned as the dissociative or bonded status. For instance, the N–O bond in the NO_2 fragment may become quite long while it is still not fully dissociated. In this case, our assignment of the photolysis channels according to the N–O bond distance was highly approximated. In this sense, the current estimation of the photolysis products ratio is rather qualitative, while such information should still be quite useful to understand the CH_3ONO_2 photodynamics.

When the dynamics starts from S_2 and S_3 , the time-dependent occupations of electronic states are shown in Fig. 8(a) and (b). No matter whether the trajectory starts from S_2 or S_3 , the ultrafast nonadiabatic dynamics was observed, as shown in Fig. 8(a) and (b). Even in the early time of dynamics,

Table 2 Final photoproducts and reaction ratios

Initial state	Final products	Ratio (%)
S_1	$CH_3O + NO_2$	83
	$CH_3O + NO + O$	5
	CH_3ONO_2 (no breaking)	12
S_2	$CH_3O + NO_2$	87
	$CH_3O + NO + O$	3
	$CH_3ONO + O$	3
	CH_3ONO_2 (no breaking)	7
S_3	$CH_3O + NO_2$	53
	$CH_3O + NO + O$	42
	$CH_3ONO + O$	5

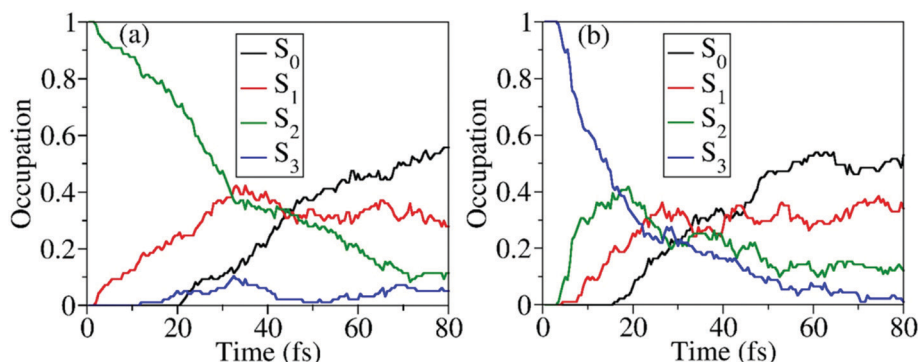


Fig. 8 Time-dependent average fractional occupations of adiabatic electronic states in nonadiabatic dynamics: (a) initiated from S_2 ; (b) initiated from S_3 . After 80 fs, the electronic structure calculations of several trajectories may not be converged at random steps. Thus the overall population dynamics is only shown up to 80 fs.

the excited-state population decays were observed, and the S_0 population started to increase quickly. As expected, when the higher excited states are prepared, the S_0 population rises more quickly.

Starting from the S_2 , all three S_1 – S_0 CIs play essential roles in the nonadiabatic decay to the ground state, as shown in Fig. S8 (ESI[†]). It is hard to distinguish their precise contributions here because more distorted hopping geometries appear. More than 93% trajectories move to the dissociation limit, giving three different reaction channels, $\text{CH}_3\text{O} + \text{NO}_2$ ($\sim 87\%$), $\text{CH}_3\text{O} + \text{NO} + \text{O}$ ($\sim 3\%$), and $\text{CH}_3\text{ONO} + \text{O}$ ($\sim 3\%$). When the dynamics starts from S_3 , the final photoproducts are $\text{CH}_3\text{O} + \text{NO}_2$ ($\sim 53\%$), $\text{CH}_3\text{O} + \text{NO} + \text{O}$ ($\sim 42\%$), and $\text{CH}_3\text{ONO} + \text{O}$ ($\sim 5\%$), as shown in Table 2.

4. Discussion

The theoretical study of the nonadiabatic dynamics of CH_3ONO_2 represents a great challenging task due to its rather complicated electronic configurations in the photolysis dynamics. First, the different types of complicated multi-state quasi-degeneracy may appear from time to time along the CH_3O – NO_2 dissociation pathway, which brings difficulty in electronic treatments. Second, the photochemistry of CH_3ONO_2 involves high-lying excited states, while their theoretical description by currently available electronic structure approaches may not be accurate enough. Third, after the internal conversion, the excessive energy could lead to highly distorted geometries at which the electronic structure calculations become non-converged or state energies may change suddenly. Because we can only trust the trajectory propagation before these improper events, the estimation of the branching ratio of the final photoproducts is rather qualitative. One question is whether it is possible to use some tricks to improve the numerical stability in the trajectory propagation after hops, for instance using some more robust theories, such as the algebraic diagrammatic construction method to the second order [ADC(2)] or density functional theory (DFT), to restart our propagation after S_1 – S_0 hops. However, we do not prefer this approach because the quasi-degeneracy of the two-lowest electronic states (S_1 and S_0), even the

multi-state degeneracy, exists in the dissociation limit. At the same time, the shorter time step may also not solve the numerical instability problem. In the current system, many orbitals may show very similar energies and start to highly mix due to orbital symmetry in the dissociation limit. In this case, the orbitals inside and outside of the active space are easily mixed, and this brings the discontinuities of the electronic wavefunctions. When the strong geometry distortion exists, this orbital mixture becomes even more pronounced. Therefore, the employment of the smaller-time step does not alleviate this problem.

However, considering the computational facilities and available theoretical approaches currently, the current treatment represents a state-of-the-art approach that in principle provides valuable information to address the major reaction channels and relevant key molecular motion of the excited-state dynamics of CH_3ONO_2 . Previous computational work demonstrated the existence of visible barriers along the direct CH_3O – NO_2 bond-breaking pathway when no additional nuclear motions are involved.²⁶ In the current work, we demonstrated that the pyramidalization at the N atom along with the CH_3O – NO_2 bond stretching can drive the system accessing an S_1 – S_0 CI (CI_{10} -III) and induce the ultrafast internal conversion dynamics. Even when the inclusion of other relevant degrees of freedom modifies the barrier height for the direct excited-state CH_3O – NO_2 dissociation channel (Fig. 7), the pathways towards the CI_{10} -I and CI_{10} -II still display visible barriers. Thus these two CIs only act as the secondary channels in the internal conversion dynamics of CH_3ONO_2 starting from S_1 . When the trajectories start from high-lying excited states (S_2 and S_3), the barriers towards CI_{10} -I and CI_{10} -II may be overcome more easily, and thus they start to play more important roles here.

No matter which CI is passed, most trajectories finally move to dissociation limits. One major dissociation channel gives the products $\text{CH}_3\text{O} + \text{NO}_2$. At this dissociation limit, the ground state CH_3O radical geometry is very close to the C_{3v} symmetry and the two lowest electronic states of the CH_3O radical become almost degenerated. This explains the existence of the state degeneracy in the dissociation limit of this channel. In fact, such state degeneracy was also noticed by previous work.²⁶ In principle, the existence of the 3-fold rotational axis of the

CH₃O may give the high symmetry structure,^{88,89} and this is the origin of the state degeneracy.^{38,39} Such a symmetric geometry is easily destroyed by the Jahn–Teller effects.^{38,39} For CH₃O, previous theoretical discussions clearly demonstrated that such Jahn–Teller distortion is very weak.^{88,89} As a consequence, the energy gap between the two lowest electronic states remains extremely small after considering the geometry distortion due to the symmetry breaking. Therefore, when the trajectories following the CH₃O + NO₂ channel move to the dissociation limit, we observed the state degeneracy.

In fact, the state degeneracy may also exist for the NO₂ part.^{90–92} As shown in previous studies, the two lowest electronic states of the NO₂ part may become close to each other when the O–N–O bond angle approaches ~106° or ~180°. ^{90,92} In our calculations, we seldom saw that the O_a–N–O_b bond angle accesses these values in the CH₃O + NO₂ dissociation limit when the trajectory starts from S₁. When the higher excited electronic state is excited initially, the excessive energy may drive the strong bending motion to nearly 106° during the dynamics and push the system close to such degeneracy. If so, many states become nearly degenerated, and one of such typical trajectories is shown in Fig. S9 (ESI†).

Some trajectories give the CH₃O + NO + O photolysis channel. The resulting O atom has three p orbitals with the same energies. When two lowest quasi-degenerated states of the CH₃O radical were taken into account, a much higher state degeneracy situation should exist. In this situation, electronic characters become extremely complicated in the dissociation limit. Our current calculation setup cannot provide the perfect description of this highly degenerated situation involving many electronic states. However, the current calculations still roughly captured the existence of this channel. Here we also showed that the ratio of this channel increases when the high-lying electronic state is excited. This feature shows a qualitative agreement with experimental works.^{20,24,27,35}

Although the current calculations face many limitations, we still can get a valuable explanation of available experimental works. First, we provided a clear interpretation of the absorption spectra of CH₃ONO₂. The ultrafast nonadiabatic decay dynamics in the current simulation indicated that the excited-state lifetime is extremely short. This gives an excellent explanation of why the low-lying bands in the absorption spectra of CH₃ONO₂ are quite broad and diffusive.^{6,27} Considering the existence of ultrafast nonadiabatic dynamics, the further ultrafast-spectroscopy experimental studies of such dynamics of CH₃ONO₂ and derivatives should be highly interesting. Second, the current simulation gives some valuable data to explain the photolysis dynamics. In experimental works,^{20,27,35,37} the photolysis dynamics at the long wavelength, ~250 nm or > 300 nm, always gives the CH₃O + NO₂ channel. In our calculations, the dynamics starting from both S₁ and S₂ show that this channel is always dominant. Although we obtained other channels possibly due to the limitation of current calculations, their branching ratios of the minor channels are rather low. We also observed that some trajectories starting from S₁ do not show dissociation, and this may explain why the quantum yield of the CH₃O + NO₂

photolysis channel is not unity in low-pressure long-wavelength (>300 nm) experimental work. Only when the trajectories started from S₃, did we observe that both CH₃O + NO₂ and CH₃O + NO + O channels become important. The system first breaks into CH₃O + NO₂, and then the N–O bond of NO₂ breaks. These features are consistent with experimental works.^{20,24,30} In the end, the CH₃ONO + O is observed when the trajectories start from S₃. This channel was confirmed by previous experimental studies with short-wavelength laser excitation.^{20,24,35}

We analyzed the dependence of the ratios of dissociation channels on the excitation energy. In each energy window, the reaction probability of each channel is determined by the destination of trajectory propagation starting from such a window and the transition probability of each initial sample.⁵⁸ As shown in Fig. 9, only in the low-energy domain (4.0–4.5 eV and 4.5–5.0 eV), the non-dissociative channel appears while it vanishes quickly with the increase of photolysis excitation energy. At the same time, the CH₃O + NO₂ channel is always dominant when the low excitation energy is taken. Above 5.0–5.5 eV, the ratio of the CH₃O + NO₂ channel decreases monotonically with the higher photolysis excitation energy, 75% at 5.5–6.0 eV and 40% at 6.0–7.0 eV. As contrast, the CH₃O + NO + O channel does not appear at all in the very low-energy domain (4.0–4.5 eV) and is basically neglectable (≤2% at 4.5–5.0 eV). This channel starts to be important with the increase of the photolysis excitation energy, giving 22% at 5.5–6.0 eV and even more than half in the high-energy domain (52% at 6.0–7.0 eV). The CH₃ONO + O channel only appears in the high-energy domain. This observation is even more consistent with the experimental results,^{6,24,27,28} compared to the data obtained by considering the photolysis from different excited states only. All experimental observations indicated that the CH₃O + NO₂ channel is dominant in the low-energy excitation, while both CH₃O + NO₂ and CH₃O + NO + O channels may become equally important in the high-energy photolysis excitation situations.

As discussed in ref. 93 and 94, the excitations of sunlight, short-pulse laser fields, and continuum-wave laser fields in principle should give rather different initial conditions for photoreactions. Using more advanced theoretical approaches suggested by these works, the proper initial conditions can be defined in the nonadiabatic dynamics simulation. In principle, these theoretical treatments should be important because the dynamics simulation results may be dependent on the selection of initial conditions. In the current work, we still keep the original ways of the initial sampling in nonadiabatic dynamics simulations. As discussed in our manuscript several times, this work represents our initial efforts to treat the photolysis dynamics of alkyl nitrate systems. In the current stage, we only wish to provide some initial understanding of the photolysis dynamics of these compounds qualitatively, instead of the accurate description at the quantitative level. In particular, the nonadiabatic dynamics simulation of these systems is still rare. Our calculations of both PES and nonadiabatic dynamics indicate that the photolysis reaction of CH₃ONO₂ is extremely fast. We also get the branching ratios of photolysis channels *vs.*

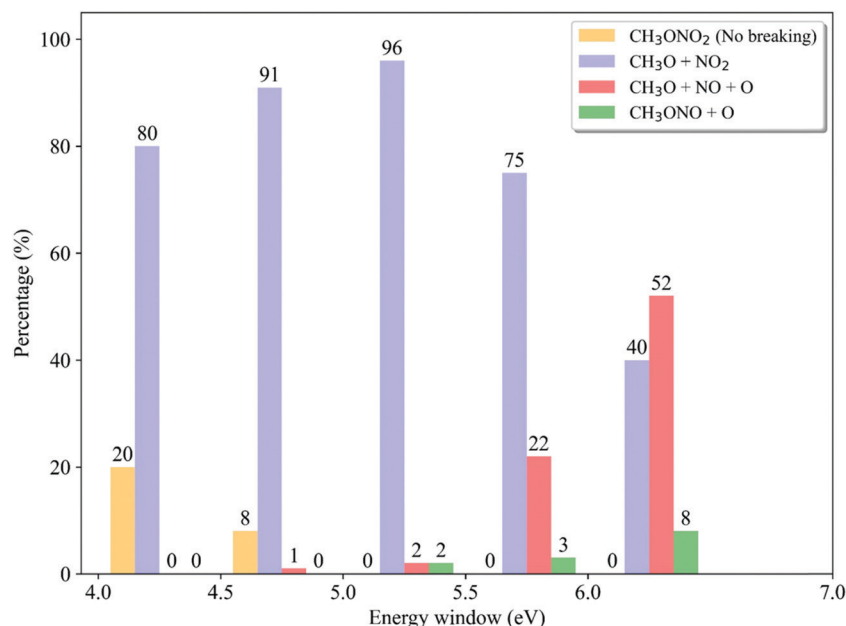


Fig. 9 The dissociation channel ratios of different initial energy windows.

the excitation wavelengths, which agrees well with the experimental observations. Therefore, the main purpose of the current research is basically achieved. For the precise lifetime, we still do not have any available experimental data for comparison. As our current simulation gives the primary results on the ultrafast excited-state lifetime, this result certainly should be verified by further experimental and theoretical works. The new time-resolved ultrafast pump-probe experiments may be helpful to address the ultrafast photolysis dynamics. At the same time, additional efforts should be made to give a better representation of the initial conditions in the nonadiabatic dynamics simulation of similar systems, in order to give better descriptions of the experimental setups and even sunlight. We believe that such a challenging study should be an important research topic in the future. On one side, this provides the direct bridge to link the experimental observations and theoretical simulations. On another side, this helps us to get a deep understanding of the realistic photochemistry of these alkyl nitrates in the atmosphere, which are environmentally significant.

It is well known that the nitro- and nitrate- compounds may undergo intersystem crossing, as shown by several previous works.^{95–97} In the current system, the intersystem crossing dynamics may not play an essential role here. The current photolysis dynamics takes place extremely fast, less than 60 fs. Within such a short time scale, the contribution of the intersystem crossing dynamics should not be significant, because in principle the weak spin-orbital coupling should exist in such molecules due to the missing heavy atom. On the other hand, the investigation of the possible role of the intersystem crossing dynamics should be an interesting topic in the future.

In the simulation of excited-state dynamics, the selection of suitable electronic state methods and dynamics approaches is not a trivial task. In the electronic-structure treatments, it is

well known that all excited-state electronic-structure approaches have their own advantages and shortcomings. Among them, the CASPT2 approaches display reasonable accuracy in many benchmark calculations.^{58,98,99} In this sense, this approach can more or less be viewed as the “accurate approach” in the excited-state electronic-structure calculations, while its shortcoming is also well known.¹⁰⁰ In the current work, the XMS-CASPT2 method is an extended MS-CASPT2 approach, while it essentially provides a similar computational accuracy with higher efficiency.^{54,55} On the other hand, some other high-level approaches, such as multireference configuration interaction (MRCI),¹⁰¹ may also be the possible choice. In recent years, we noticed that there is a huge development in other advanced excited-state calculation approaches, such as multireference coupled cluster (MRCC),¹⁰² and full configuration-interaction quantum Monte Carlo (FCI-QMC) methods.¹⁰³ If these methods can be used to simulate the excited state dynamics, we expect that more fruitful insights can be generated. However, there is still a long way to employ them in the excited-state dynamics.

In the choice of the dynamics approaches, a similar situation is held. The on-the-fly TSH is a practical choice that gives a reasonable description of the excited-state nonadiabatic dynamics with the balance of computational accuracy and efficiency. However, its shortcoming is also widely known, for instance, its overcoherence problems.^{74,104,105} Other more advanced approaches, such as *ab initio* multiple spawning,⁶² multi-configurational Ehrenfest,¹⁰⁶ variational multiconfiguration Gaussian (vMCG),^{107,108} different on-the-fly semiclassical dynamics based on the mapping Hamiltonian,^{77,109} and coupled-trajectory surface-hopping dynamics based on exact factorization,^{110–112} may become possible choices in the future. If we find a suitable electronic structure method that shows enough accuracy in the treatment of the

excited state, it would be highly interesting to take it and try to benchmark the performance of different on-the-fly mixed-quantum-classical and semiclassical dynamics approaches. This should provide important evidence to help us select the proper dynamics approaches in the treatment of realistic polyatomic systems with full dimensionality.

5. Conclusion

In this work, we studied the excited-state nonadiabatic dynamics of CH_3ONO_2 using the on-the-fly TSH simulation at the XMS(4)-CASPT2(12,9)/def2-SVPD level. We mainly paid close attention to the nonadiabatic decay dynamics and the successive photolysis channels.

Three $\text{S}_1\text{--S}_0$ MECIs were located at the SA(4)-CASSCF(12,9)/cc-pVDZ level. Both $\text{CI}_{10}\text{--I}$ and $\text{CI}_{10}\text{--II}$ display significant $\text{CH}_3\text{O--NO}_2$ elongation, while the $\text{O}_a\text{--N--O}_b$ bond angle bending motion is observed at $\text{CI}_{10}\text{--II}$. The $\text{CI}_{10}\text{--III}$ shows strong pyramidalization at the N atom, as well as a much shorter $\text{CH}_3\text{O--NO}_2$ distance with respect to $\text{CI}_{10}\text{--I}$ and $\text{CI}_{10}\text{--II}$.

The ultrafast decay processes from different initial excited states were simulated using the on-the-fly TSH simulations. A majority of trajectories starting from S_1 jump back to S_0 via the $\text{CI}_{10}\text{--III}$ geometries with the weakness of the $\text{CH}_3\text{O--NO}_2$ bond and the pyramidalization at the N atom. When the trajectories start from high-lying excited states (S_2 and S_3), the internal conversion dynamics becomes faster as expected, and all three CI_{10} become essential in the nonadiabatic decay.

Starting from both S_1 and S_2 , most trajectories show the photodissociation and follow the $\text{CH}_3\text{O} + \text{NO}_2$ photolysis channel. In the dissociation limit, the state degeneracy exists between S_1 and S_0 due to the 3-fold rotational symmetric geometry of CH_3O . The other channels only play a minor role here. When the trajectories start from S_3 , we still observed that the $\text{CH}_3\text{O} + \text{NO}_2$ channel is the major one, while the secondary channel $\text{CH}_3\text{O} + \text{NO} + \text{O}$ also becomes important. For the latter one, the $\text{CH}_3\text{O--NO}_2$ bond first breaks and the cleavage of the N–O bond in the NO_2 fragment then occurs. A third photolysis channel $\text{CH}_3\text{ONO} + \text{O}$ also appears. The dependence of the photoproduct channels on the UV excitation wavelength is qualitatively in agreement with available experimental studies.

The current work presents a primary study of our understanding of organic nitrate photochemistry. It should be highly interesting to simulate these processes with highly accurate electronic structure methods, treat the highly-excited state dynamics with more proper dynamics approaches, simulate the experimental signals and study the chemical substitution effects on the photolysis. At the same time, in recent years we witness that great experimental efforts were made to study the ultrafast nonadiabatic dissociation dynamics of VOC systems.^{113–118} On the basis of the current theoretical work, the CH_3ONO_2 and derivatives may be a group of interesting prototype VOC systems in experimental ultrafast spectroscopy studies. These works will help us understand more thoroughly

the photochemistry of VOCs and this represents a great challenging task in future studies.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

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