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A Multireference Configuration Interaction Study of the Photodynamics of Nitroethylene

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Supporting Information

ABSTRACT: Extended multireference configuration interaction with singles and doubles (MR-CISD) calculations of nitroethylene ($H_2C=CHNO_2$) were carried out to investigate the photodynamical deactivation paths to the ground state. The ground (S_0) and the first five valence excited electronic states (S_1 - S_5) were investigated. In the first step, vertical excitations and potential energy curves for CH₂ and NO₂ torsions and CH₂ out-of-plane bending starting from the ground state geometry were computed. Afterward, five conical intersections, one between each pair of adjacent states, were located. The vertical calculations mostly confirm the previous assignment of experimental spectrum and theoretical results using lower-level calculations. The conical intersections have as main features the torsion of the CH₂ moiety, different distortions of the NO₂ group and CC, CN, and NO bond stretchings. In these



conical intersections, the NO₂ group plays an important role, also seen in excited state investigations of other nitro molecules. Based on the conical intersections found, a photochemical nonradiative deactivation process after a $\pi - \pi^*$ excitation to the bright S₅ state is proposed. In particular, the possibility of NO₂ release in the ground state, an important property in nitro explosives, was found to be possible.

1. INTRODUCTION

The explosophore NO₂ group is an essential component in many propellants and explosives of practical relevance.¹ These nitro compounds are usually thermally unstable, composed of easily broken N–N, O–C and C–N covalent bonds, have significant biological activity, and may contribute to atmospheric pollution.² Furthermore, decomposition processes of nitro compounds are especially important because they occur at relatively low temperatures and are involved in the detonation of explosives.^{3,4}

Most of the energetic molecules have diffuse electronic spectra and propensity to dissociate and rearrange.⁵ In the detonation initiation of these materials, photochemical processes are very important.^{6–12} In particular, as has been increasingly recognized in the field of photochemical reactions, ^{13,14} nonradiative decomposition processes of energetic molecules through conical intersections play an important and even dominant role.^{15–17} Therefore, investigations of nonadiabatic processes occurring in energetic molecules are especially interesting.

The study of gas phase energetic molecules can unveil their properties and reactions at the molecular level, and determine decomposition features as a function of the electronic and vibrational excitations.⁷ However, the experimental measurements of decomposition processes and their interpretation are

very complex, a picture asking for theoretical investigations. Moreover, in spite of the importance of excited states in energetic materials research, only few studies have been carried out in this field. 6,17

We recently investigated the electronic excitation spectra of energetic molecules containing nitro groups, namely, nitramide (H_2NNO_2) ,¹⁸ *N*,*N*-dimethylnitramine, $((CH_3)_2NNO_2)$,¹⁹ 1,1-diamino-2,2-dinitroethylene (FOX-7 or DADNE),²⁰ and hexahydro-1,3,5-trinitro-1,3,5-triazine (RDX).²¹ These molecules are building blocks of energetic materials and bear easily broken N–NO₂ bonds. Nitroethylene (H₂C=CHNO₂) and FOX-7, in contrast with the other studied molecules, have a C–NO₂ bond instead of a N–NO₂. These C–NO₂ bonds are also relatively weak and have its NO₂ group adjacent to a C=C bond.

Nitroethylene is the simplest member of the nitro-olefin series. This molecule displays conjugation effects between the ethylenic π electrons and those on the nitro group²² that affect the electronic spectrum. Molecules of this class are potentially synthetic reagents, powerful electron acceptors due to the

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presence of NO₂ groups, and can also behave as precursors to useful polymers and copolymers for explosives technologies.³

In spite of the variety of aforementioned interests, previous work on the nitroethylene molecule is scarce. Karle et al. discussed its unimolecular decomposition using MP2, MP4 and G2 methods⁴ and Zhang and co-workers focused on ground-state isomerization reactions and vertical calculations of the first five electronic transitions, respectively at the DFT/B3LYP/6-31G* and TD-DFT-B3LYP/6-31G* levels of theory.²³ Shamov and Khrapkovskii considered the pyrolysis mechanism of nitroethylene using the B3LYP/6-31G(d) method.²⁴ Khrapkovskii and co-workers reviewed quantum chemical data on gas-phase decomposition mechanisms of C-nitro compounds including nitroethylene.²⁵

Concerning the excited electronic states of nitroethylene up to the year 2006, apparently only three other theoretical studies and an older experimental work can be found in the literature. The spectrum of nitroethylene in gas phase was measured in the 1960s by Loos and co-workers,²⁶ who observed a very weak band at 4.2 eV and an intense band centered at 6.12 eV with a shoulder at 5.12 eV. Those authors used the semiempirical Pariser-Parr-Pople configuration interaction (PPP-CI) method to assign the intense band and its shoulder to $\pi - \pi^*$ transitions. In another work, Ha employed SCF-CI wave functions and a small Gaussian basis set to find also the character of the first band as a $n-\pi^*$ transition and of the intense band and its shoulder as two $\pi - \pi^*$ transitions.²⁷ In 2006, Zhang and co-workers²³ examined vertical excitations computed at the TDDFT-B3LYP/6-31G* level once more confirming the $\pi - \pi^*$ character of the intense band and its shoulder.

Recently, we carried out a systematic examination of the electronic spectrum of nitroethylene using a variety of methods:28 the approximate coupled-cluster singles and doubles wave function with the resolution of the identity (RI-CC2), the time dependent density functional theory with the CAMB3LYP functional (TDDFT/CAMB3LYP) and the DFT multireference configuration interaction (DFT/MRCI). Vertical transition energies and optical oscillator strengths of up to 20 singlet transitions were computed. Additionally, semiclassical simulations of ultraviolet (UV) spectra at the RI-CC2 and DFT/MRCI levels were performed. The main features in the UV spectrum were assigned to a weak $n-\pi^*$ transition, and to two higher energetic ones as $\pi - \pi^*$ bands. The simulated spectra is in good agreement with the experimental spectrum. These features of the nitroethylene spectrum are similar in molecules containing NO₂ groups. The transition energies of the bands in the DFT/MRCI simulation agreed quite well with experiment, although it overestimated the band intensities. The RI-CC2 calculation produced intensities comparable to the experiment, but the bands were blue-shifted by 0.5 eV. A strong $\pi - \pi^*$ band, not previously measured, was found in the 8-9 eV range.

The goal of this work is to discuss the photochemical pathways for the decay of nitroethylene to the ground state. In this discussion, the photodynamics of ethylene^{29–35} and fluoroethylene³⁶ will guide us, the latter serving as an example for a polar biradicoloid. Both cases can be qualitatively explored by a 3×3 CI analytical model as developed by Bonačić-Koutecký and Michl.^{37,38} This model predicts for non-symmetric biradicaloids like fluoroethylene the appearance of a degeneracy between the S₀ and S₁ states by simple torsion of the CC double bond, as was confirmed by quantum chemical

calculations.³⁶ In contrast, it is well documented that in ethylene the torsion is not sufficient for reaching an intersection with the ground state but that pyramidalization and hydrogen transfer modes play an important role.^{29,31,32} From this point of view, nitroethylene is an interesting example, since it polarizes the C=C bond due to electronegativity differences of the groups bonded to it, but it also includes contributions from π conjugation and lone pair orbitals. For this reason, a multitude of excited states already appears in the case of vertical excitations,²⁸ and conical intersections should involve CH₂ and NO₂ torsions. The calculation of excited-state energy surfaces and the search for conical intersections of nitroethylene is thus even more involved.

In this work, we employ the multireference configuration interaction with singles and doubles (MR-CISD) method³⁹ to calculate potential energy curves for selected coordinates, to determine minima on the crossing seam (MXS) and to characterize nonadiabatic interactions between different electronic states of nitroethylene. These features should allow us to develop an overall picture of the nitroethylene photochemistry through possible nonradiative decay mechanisms originating in a bright electronically excited molecule leading it to reach the ground state.

2. COMPUTATIONAL DETAILS

The calculations were carried out in two steps. The first one consisted of a state-averaged multiconfiguration self-consistent field (SA-MCSCF) calculation followed by a multireference configuration interaction with single and doubles excitations (MR-CISD). The state averaging was performed with equal weights and included six states (SA-6), namely, the ground state $S_0(1^1A')$, three A" and two A' states in the case of vertical excitations. For describing these excitations, a complete active space-CAS(8,6)-including eight electrons and six orbitals, the four highest occupied orbitals (two a" and two a'), and two virtual orbitals of a" symmetry, was chosen. The SA-MCSCF character of the orbitals are $2a'' [\pi (NO_2)]$, $15a' [n\sigma]$, $16a' [n\sigma(O)]$, $3a'' [\pi (C=C+NO_2)]$, $4a'' [\pi^*]$ and $5a'' [\pi^*]$ (see Figure 1). The 15a' and 16a' orbitals are respectively n_{+} and n_{-} type-orbitals because the former corresponds to symmetric linear combinations of the oxygen lone pair orbital and the latter, the highest occupied molecular orbital (HOMO), represents the antisymmetric combination.²⁸ This SA-6-CASSCF calculation was followed by a MR-CISD with the reference configurations generated from the same CAS (8,6)



Figure 1. SA-MCSCF active space nitroethylene molecular orbitals.

Table 1. Computed MCSCF, MR-CISD, and MR-CISD+Q Vertical Excitation Energies ΔE (eV) in Comparison with Previous Results and Experimental (Exp.) Data^{*a*}

	10000	100		6	1	DTT () TO ork	
state	MCSCF	MRCI	MRCI+Q	f	character	DFT/MRCI ²	exp.
S_1-1^1A''	3.47	3.69	4.03	0.000	n <i>π</i> * (16a'-4a")	3.62(0.000)	4.20(0.002)
S_2-2^1A''	4.06	4.14	4.48	0.000	$n_{+}-\pi^{*}$ (15a'-4a")	4.11(0.000)	
S_3-2^1A'	6.08	6.33	6.05	0.034	$\pi - \pi^* (2a'' - 4a'')$	5.19(0.051)	5.12(0.012)
S_4-3^1A''	7.00	7.33	7.37	0.000	$n_{+}-\pi^{*}$ (15a'-5a")	6.25(0.000)	
S_5-3^1A'	7.18	7.58	6.00	0.375	$\pi - \pi^* (2a'' - 4a'')$	6.09(0.432)	6.12(0.304)
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^aThe computed MR-CISD oscillator strengths (*f*) are also reported. The DFT/MRCI and experimental oscillator strength values are shown in parentheses. ^bOur previous results.¹⁸ ^cExperimental data.¹²

space and applying the interacting space restriction.⁴⁰ This orbital space has been chosen to represent the reference space for the computation of the five lowest singlet valence excited states as discussed in our previous work.²⁸ Since the focus of this work is laid on the exploration of the photodeactivation mechanism of the valence states, Rydberg states, which appear above those in the upper range of the states chosen,²⁸ were not included in our calculations.

The carbon, oxygen, and nitrogen 1s orbitals were frozen in the CI calculations. The Davidson^{41,42} corrections for sizeextensivity were also computed. The $6-31++G^{**}$ Gaussian basis set⁴³ was used. This type of calculation is the standard one and will be denoted MRCI. It will be used for the calculation of the vertical spectra, potential curves, and MXS structures. Two significantly extended approaches will be described in the following paragraph. They will be used only for calculating vertical excitation energies.

For vertical excitations, two additional kinds of calculations were carried out. In the first, the basis set was extended significantly to $6-311+G(2d1f)^{44,45}$ on the heavy atoms. For the hydrogen atoms, a 6-31G** basis was chosen. The reference space was kept at the CAS(8,6) level. This calculation is denoted MRCI(extbas). In the second kind of calculation, the active space of the reference wave function was increased while keeping the basis set at the 6-31++G** level. This extended reference wave function is denoted RAS(2,1)CAS(8,6)AUX-(2,1). The acronym RAS stands for a restricted active space and AUX for auxiliary space. The notation RAS(2,1) means that two a' and one a" orbital were moved from the doubly occupied space into the active space. In the case of AUX(2,1), two a' and one a" virtual orbitals were moved into the active space. The CAS(8,6) remained unchanged. Only single excitations were allowed from the RAS and into the AUX space. This configuration space was used as a wave function in the MCSCF calculation and as a reference wave function in the case of the MR-CISD calculation. The corresponding calculation is named MRCI(extref).

The geometries were described by natural internal coordinates⁴⁶ and optimized using the direct inversion in the interactive subspace (GDIIS) method.⁴⁷ Geometry optimizations and determination of the minima on the crossing seam (MXS), i.e., the conical intersections respectively employed analytic MR-CISD gradients^{48,49} and analytic MR-CISD nonadiabatic coupling vectors.^{50–52} The exception was the S_2/S_1 and S_5/S_4 conical intersections, which were converged only at the CAS (8,6) level; in these cases, a single-point MR-CISD with the same type of wave function used for the other geometries were done. The potential energy curves were obtained with full optimization of all coordinates except the selected coordinate for constructing the potential curve.

The COLUMBUS suite of programs was used for all calculations.⁵³⁻⁵⁵ The calculations were in part performed using the parallel version of COLUMBUS.^{56,57} The atomic orbital (AO) and the derivative integrals were calculated using the DALTON program.⁵⁸

3. RESULTS AND DISCUSSION

3.1. Vertical Transitions. The DFT/B3LYP//6-311G +(2df,2p) planar ground state geometry obtained in our previous work²⁸ was employed for the vertical calculations. Transition energies and oscillator strengths from the S₀ ground state to the S₁–S₅ valence excited states were computed. The results are collected in Table 1 together with our previous DFT/MRCI results²⁸ and available experimental data. The SA-6-CASSCF(8,6) molecular orbitals are depicted in Figure 1. The results of the two additional vertical calculations are shown in Table 2.

Table 2. Computed MCSCF, MR-CISD, and MR-CISD+Q Vertical Excitation Energies ΔE (eV) Using the MRCI(extbas) and MRCI(extref) Approaches

			MRCI			MRCI
state	MCSCF (extbas)	MRCI (extbas) ^a	$^{+\mathrm{Q}}_{(\mathrm{extbas})^a}$	MCSCF (extref)	MRCI (extref) ^b	+Q (extref) ^b
$\overset{S_1-}{1^1A''}$	3.45	3.86	4.00	3.64	3.94	4.07
$S_{2^{-}} 2^{1}A''$	4.03	4.39	4.47	4.53	4.48	4.40
S ₃ - 2 ¹ A'	6.07	6.30	6.10	6.34	6.09	5.74
S ₄ - 3 ¹ A"	6.98	7.62	7.28	7.13	7.39	7.38
S ₅ - 3 ¹ A'	7.12	7.23	6.15	8.18	7.34	5.83
^a 6-311+	G(2d1f)/	6-31C**/	CAS(8)	5)-MR-CIS	D ^b 6-31	++C**//

"6-311+G(2d1f)/ 6-31G**// CAS(8,6)-MR-CISD. "6-31++G**// RAS(2,1)CAS(8,6)AUX(2,1)-MR-CISD.

The measured weak band is assigned to the two $n-\pi^*$ transitions involving states S_1 and S_2 . The MR-CISD transition energies computed in this work for the S_1 and S_2 transitions are (MR-CISD+Q values between parentheses): 3.69 eV (4.03 eV) and 4.14 eV (4.48 eV). These two dark transitions are respectively excitations from the n_- (16a') and n_+ (15a') orbitals localized on the NO₂ moiety to the same π^* (4a") final orbital. There is another dark transition (S_4) that also originates from the n_+ (15a') orbital, but in this case is an excitation to the higher π^* (Sa") orbital.

The S₅ state corresponds to the measured band maximum and the S₃ corresponds to the band shoulder; both transitions are of $\pi - \pi^*$ type. The S₃ state has MR-CISD transition energies (MR-CISD+Q between parentheses) of 6.33 eV (6.05 eV) and oscillator strength f = 0.034 (MR-CISD) and the S₅ has 7.58 eV (6.00 eV) and f = 0.375. The S₃ state is a transition from a π (2a") orbital localized on the NO₂ group to a π^* (4a") orbital while the S₅ state, the brightest of the computed set, is an excitation from the π (3a") orbital to the π^* (4a") having contributions from electron density distributed over the CC bond as well. As it is typical of energetic molecules, the bright $\pi - \pi^*$ transition involve orbitals mostly localized on the NO₂ moiety.

The transition energy of the S₅ bright $\pi - \pi^*$ transition computed at the MR-CISD+Q level agrees quite well with the experimental band maximum. The corresponding MR-CISD value is too high in energy. The weaker $\pi - \pi^*$ transition (S₃) is too high at both the MR-CISD and MR-CISD +Q levels.

In contrast to our previous results,²⁸ there is an inversion of the MR-CISD energy ordering of the $n-\pi^*$ (S₄) and $\pi-\pi^*$ (S₅) transitions in comparison with the DFT/MRCI calculations (Table 1). However, the vertical energy ordering becomes compatible with DFT/MRCI after inclusion of the Davidson (MR-CISD-Q) correction.

An overall comparison of the MCSCF transition energies with all the available MR-CISD values (Table 1 and Table 2) shows relatively little changes except for the MCSCF(extref) results of the S_5-3^1A' state. This shows that our standard CAS(8,6) reference space is in principle flexible enough to represent the major features of all states investigated except the one mentioned. In this case, the MCSCF(extref) enhances the weight of $\pi - \pi^*$ excitations on the cost of artificial $n^2 - \pi^2$ excitations; thus, the character of the S5 state is considerable improved in spite of the higher MCSCF excitation energy. The MR-CISD approach actually increases in most cases the excited state energies relative to the ground state; the most prominent exception is the S₅ state for which the MRCI(extref) not only enhances the weight of the excitation from the π (3a") to the π^* (4a") orbital, but also stabilizes this state relative to the others. Significant changes are observed by applying the Davidson correction which decreases the excitation energies of the two $\pi - \pi^*$ states 2¹A' and 3¹A' considerably, especially of the latter state. The energetic location of both states is especially improved by extension of the active space to the extref case, but the $2^{1}A'$ state is still about 0.6 eV above the experimental value. Further extension of the active space beyond the present RAS/CAS/AUX scheme of 14 electrons in 12 active orbitals would be necessary in order to increase especially the stability of the 2¹A' state, but such an approach will render the calculations very costly. Since our focus was not so much concentrated on the vertical excitations but on the exploration of the photodynamical deactivation picture, we did not follow this path further.

Overall, the MR-CISD and MR-CISD+Q methods provide a good starting point for the following discussion of reaction paths on excited-state energy surfaces and the determination of conical intersections. These investigations will lead into regions of the energy surfaces far away from the Franck–Condon region, and it is this aspect where the multireference method will be most useful. For this purpose, we have chosen the standard MRCI/CAS(8,6) approach which gives a good balance between computational efficiency and flexibility of the wave function.

3.2. Potential Curves. Following the experience with the photodeactivation mechanisms found for ethylene³¹ and fluoroethylene,³⁶ the MR-CISD potential energy curves of the states S_0-S_5 were calculated for two rigid torsions: of the NO₂ group around the C–N bond (Figure 2) and of the CH₂ group





Figure 2. Rigid MR-CISD potential energy curves for NO_2 rotation around the CN bond starting from the DFT/B3LYP//6-311G +(2df,2p) ground state geometry.

around the C=C bond (Figure 3); the latter torsion also occurs in ethylene.³¹ The MR-CISD CH_2 out-of-plane bending



Figure 3. Rigid MR-CISD potential energy curves for CH_2 rotation around the CC bond starting from the DFT/B3LYP//6-311G +(2df,2p) ground state geometry.

curves (Figure 4) were also computed. In all cases, the calculations started from the DFT/B3LYP//6-311G+(2df,2p) ground state geometry.²⁸

We have seen above that the $\pi - \pi^* S_3$ state $(2^1A')$ is a vertical transition characterized by molecular orbitals mostly localized in the NO₂ moiety. Upon NO₂ torsion, the S₃ excited state is destabilized by about 0.5 eV (Figure 2). The other states are destabilized also, albeit to a lesser degree. A partial exception to this behavior is the S₄ state that, after initially stabilizing, destabilizes also following the behavior of the other states. There is an avoided crossing between S₃ and S₄ at around 45° where the character of the two states is exchanged. S₄ now acquires $\pi - \pi^*$ character and S₃ becomes $n_+ - \pi^*$. For greater torsion angles, the S₄ and S₅ states become almost degenerate. The small gap in the curves between the pair of states S₃-S₄ and S₄-S₅ indicates the possible presence of conical intersections and crossing seams involving torsion of



Figure 4. MR-CISD CH_2 out-of-plane bending starting at the DFT/ B3LYP//6-311G+(2df,2p) ground state geometry.

the NO_2 moiety. In fact, these conical intersections were found (see below).

For the CH₂ torsion curves (Figure 3), the picture has more features. The S₃ state in nitroethylene, a $\pi - \pi^*$ vertical excitation, is stabilized by about 0.5 eV for increasing CH₂ torsion angles. The S₂ and S₃ states reveal an avoided crossing and exchange their character at an angle of about 56°. From this angle onward, the dominant character of S₃ is n₊ $-\pi^*$ while the S₂ state becomes $\pi - \pi^*$. Both states become almost degenerate at 70° with an energy difference of 0.3 eV, thus indicating a crossing.

Similar to ethylene, the ground state S_0 (1¹A') in nitroethylene destabilizes strongly upon CH₂ torsion (Figure 3) by almost 3 eV. This situation leads to a reduction of the gap to the first excited state (S₁) by about 2 eV, a 50% decrease. This contrasts with the very small destabilization of the ground state upon NO₂ torsion (Figure 2). For a CH₂ torsion angle of ~70°, the character of the S₀ state, π^2 at 0°, now has a $\pi - \pi^*$ configuration that contributes with a weight of about 20%. For CH₂ torsion angles of 90°, the S₀ state is dominated by a $\pi - \pi^*$ configuration with weight of 57%, while for S₁ another $\pi - \pi^*$ configuration (weight = 12%) mixes with the dominant n₋- π^* character (weight = 36%). This character is essentially maintained in the MXS (S₁/S₀) found (see Table 3).

For ethylene, it has been shown that pyramidalization of the CH₂ group (combined with partial hydrogen migration) is important for reaching the conical intersection between a $\pi - \pi^*$

Table 3. MR-CISD Character of the Conical Intersections with Corresponding Coefficients

MXS	character
S_{1}/S_{0}	$\begin{array}{l} (0.72)\mathbf{n_{+}}{-\pi^{*}} + (0.47)\pi{-\pi^{*}}/(0.85)\mathbf{n_{-}}{-\pi} + (0.17)\mathbf{n_{-}}^{2}{-\pi^{*}}{\pi^{*}} + (0.16)\\ (\pi^{*})^{2\alpha} \end{array}$
S_2/S_1	$(0.79)n_{+}-\pi^{*}+(0.27)\pi-\pi^{*}/(0.83)n_{-}-\pi^{*}+(0.29)(\pi^{*})^{2}$
S ₃ /S ₂	$\begin{array}{l} (0.38) \mathrm{n_{-}}\pi^{*} + (0.30) \ \pi^{2} - (\pi^{*})^{2} + (0.29) \ \pi - \pi^{*} / \ (0.46) \ \mathrm{n_{-}}\pi^{*} + \\ (0.28) \ \mathrm{n_{+}}\pi^{*} \end{array}$
S_{4}/S_{3}	$n_{\pi}\pi^{*}\pi^{*}/(0.64)\pi^{-}\pi^{*} + (0.29) n_{\pi}\pi^{*}$
S_{5}/S_{4}	$ (0.54)(n_{-})^{2} - (\pi^{*})^{2} + (0.42) \pi^{2} - (\pi^{*})^{2} / (0.47)n_{+} - \pi^{*} + (0.35) n_{-} \pi - \pi^{*} \pi^{*c} $

^{*a*}S₀: double excitation from n₋ to the two distinct virtual π^* orbitals. ^{*b*}S₄: double excitation from n₋ and π to the two distinct virtual π^* orbitals. ^{*c*}S₅: double excitation from n₋ to a π^* . state and the S_0 .^{29,31,59–61} Therefore, we also investigated the effect of the CH₂ out-of-plane bending (i.e., pyramidalization) coordinate through the potential energy curves of nitro-ethylene.

The MR-CISD CH₂ out-of-plane bending curves of nitroethylene starting from the ground state geometry are shown in Figure 4. Overall, the six states including the ground state steadily destabilize by about 1 eV. The S₄ and S₅ exchange $n_+-\pi^*$ and $\pi-\pi^*$ character via an avoided crossing for a CH₂ out-of-plane angle of 55°. The energies of the pair of S₄/S₅ curves slightly decreases with increasing CH₂ out-of-plane angle, and the energy splitting remains very small. Thus, for nitroethylene there is no significant effect of the CH₂ out-ofplane motion on the expected conical intersection structures.

3.3. Ground State and Structure of the Conical Intersections. The SA-6-MCSCF/MR-CISD optimized geometries of the ground state and of the minima on the crossing seam (MXS) are shown in Figures 5–8. The optimized ground state distances and angles computed in this work at MR-CISD level are displayed in Figure 5a. They agree very well with the ones obtained previously by means of the DFT/B3LYP approach.²⁸

The main feature of the MXS (S_1/S_0) (Figure 5b) as compared with the MR-CISD ground state is the position of the CH₂ group, twisted by 97°: it is now essentially perpendicular to the original molecular plane. There is a CC bond elongation of 0.14 Å and also an elongation of the NO bonds: the larger NO elongation is 0.14 Å, and the smaller one



Figure 5. Optimized geometries of nitroethylene. (a) MR-CISD ground state. Computed conical intersections (MXS): (b) MR-CISD MXS (S_1/S_0), and (c) CASSCF MXS (S_2/S_1). Angles are in degrees and distances are in Å. The dihedral angles are shown according to the numbering given in the inset at panel a. Insets of panels b and c are front views of the corresponding structures.

is 0.02 Å. In contrast, the C–N bond shortens by -0.16 Å. Although the torsional structure of the MXS (S₁/S₀) could be deduced from the large CH₂ torsion angle of the corresponding potential curves, Figure 3, the just indicated additional bond length variations are crucial to reach the conical intersection. As already mentioned above, out-of-plane bending of the CH₂ group is not important in nitroethylene.

The MXS (S_2/S_1) (Figure 5c) is overall similar to the MXS (S_1/S_0) (Figure 5b); the torsion of the CH₂ group by about 90° is also the prominent feature. There are differences in the elongation of the NO and CN bonds of the MXS (S_2/S_1) as compared to the MXS (S_1/S_0) as well as differences in the angular values despite the two insets looking very similar.

The MXS (S_3/S_2) (Figure 6a) has distinct features. There is a torsion of the NO₂ group and of the C₂H₈ bond (for atomic



Figure 6. Computed conical intersections (MXS): (a) MR-CISD MXS (S_3/S_2) , (b) MR-CISD MXS (S_4/S_3) , and (c) CASSCF MXS (S_5/S_4) . Angles are in degrees and distances are in Å. The dihedral angles are shown according to the numbering given in the inset in Figure 5, panel a. Insets are front views of the corresponding structures.

numbering, see the inset of Figure 5a) in the opposite sense leading to an almost perpendicular orientation of both relative to the ground state molecular plane (see inset of Figure 6a). There is an opening of the ONO bond and a visible pyramidalization of the NO₂ group. The CN distance in the MXS (S_3/S_2) structure decreases by 0.03 Å as compared to the ground state. The three CH distances decrease slightly by 0.04 Å, while the larger NO distance increases by 0.16 Å and the smaller one increases by 0.02 Å. The CN bond shortens by 0.04 Å. There are considerable variations of the valence angles, the most pronounced one being the 9° increase in the NO₂ opening angle and the CH₂ opening angle decreasing by 2.6°. Although the shape of the MXS (S_4/S_3) (Figure 6b) is overall similar to the MXS (S_1/S_o) (Figure 5b) and MXS (S_2/S_1) (Figure 5c) conical intersections, there are important differences. In the following, we continue to compare distances with the ground state values. The CC bond of the MXS (S_4/S_3) (Figure 6b) elongates by 0.13 Å and the CN bond by 0.3 Å. The N₃O₅ bond stretches by 0.07 Å, while the other NO bond barely changes. In comparison with the other conical intersections, the MXS (S_4/S_3) stretches almost all bonds, and the CH₂ group twisted by about 80° as compared to the ground state structure. The NO₂ group is also twisted almost by 90° and displays some pyramidalization.

In the MXS (S_5/S_4) (Figure 6c), the structure is almost planar; the CH₂ group is twisted by a small angle as compared to the other conical intersections, but overall the geometry is more planar in comparison with the other MXS's. The NO₂ angle increases by 5.5° and all the bonds elongate, excepting the CN and CH distances.

Table 3 collects the character of the conical intersections found. It can be seen that the character of the MXS's could be qualitatively inferred from the computed potential energy curves. In particular, the overall similarities of the MXS (S_1/S_0) and MXS (S_2/S_1) geometries discussed above have correspondence with the respective character. The expected multiconfiguration character of the states in most intersections is clear, and $\pi - \pi^*$ type transitions are relevant in the five intersections. Moreover, there is an important role in the character of all conical intersections played by the lone pair (n) orbitals localized on the NO₂ moiety.

3.4. Photodynamical Mechanism. The g- and h-vectors characterizing the conical intersection^{50,52,62,63} are shown in Figures 7 and 8 for the five computed MXS structures. The g-vector is defined by the energy gradient difference between the two states, and the h-vector is the nonadiabatic coupling vector. A molecular deformation along these vectors determines the branching space, lifts the degeneracy of the states, and describes possible reaction coordinates.

For the MXS (S_1/S_0) , the g-vector (Figure 7a) is dominated by the CN stretching and NO₂ wagging. The h-vector represents a CH₂ torsion around the CC bond. The CC torsion leading to a reduction of the S_1/S_0 gap also occurs in ethylene and fluorethylene. As discussed before for ethylene, the S_1/S_0 gap is reached via CH₂ pyramidalization and partial hydrogen migration³¹ while in fluorethylene CC torsion and stretching reduce the S_1/S_0 gap.³⁶ Therefore, in this case nitroethylene resembles more the fluorethylene case because for both molecules, in contrast to ethylene, the CH₂ pyramidalization does not play a role.

Although the geometry of the MXS (S_2/S_1) is similar to the MXS (S_1/S_0) , their g- and h- vectors (Figure 7b) display distinct features. The most important ones are the bending deformation of the NO₂ group and the absence of CH₂ motion. Therefore, only NO₂ deformation is relevant to reach the S_2/S_1 intersection.

The MXS (S_3/S_2) g- and h- vectors (Figure 7c) display pronounced stretching modes of the NO₂ bonds, combined with an angular deformation of the NO₂ group and a stretching of the CC bond. The h-vector also indicates a CH₂ torsion and an angular deformation of the CH₂ bond. The g- and h- vectors of the MXS (S_4/S_3) (Figure 8a) are dominated by an out-ofplane deformation of the NO₂ group with a contribution of a CC stretching. The MXS (S_5/S_4) (Figure 8b) g- and h-vectors

The Journal of Physical Chemistry A



Figure 7. Computed g- and h-vectors of the conical intersections found. (a) MR-CISD MXS (S_1/S_0) , (b) CASSCF MXS (S_2/S_1) , and (c) MR-CISD MXS (S_3/S_2) . The arrows are not to scale between different pictures.



Figure 8. Computed g- and h-vectors of the conical intersections found. (a) MR-CISD MXS (S_4/S_3) and (b) CASSCF MXS (S_5/S_4) . The arrows are not to scale between different pictures.

also display a deformation of the NO_2 group and CC stretching, similarly to the other conical intersections.

The existence of a series of conical intersections in nitroethylene suggests nonradiative deactivation processes via internal conversion (IC). In Figure 9, an overview of possible deactivation paths based on the energies and geometries of the conical intersections is presented. It can be seen that the S_5/S_4 ,



Figure 9. Proposed photochemical nonradiative deactivation process. The path leading to the possible release of NO_2 is also indicated. The main structural modifications of the conical intersections with respect to the ground state geometry are displayed.

 S_4/S_3 and the S_3/S_2 conical intersections are quite close in energy. Based on this scheme and along with the computed potential energy curves, the following nitroethylene deactivation photodynamic process can be deduced.

Bearing in mind that because nitroethylene is a multistate problem, the dynamics need not proceed necessarily through minimum-energy conical intersections, but may take shortcuts through near-degeneracies of higher energy (e.g., as observed in the ethylene dynamics³⁴), we suggest the following photodynamic mechanism. The photodynamics starts from the Franck-Condon region by populating the bright S₅ state. The torsion around the CC bond (Figure 3) does not lead to a pronounced stabilization as is found in ethylene³¹ or fluoroethylene.³⁶ Therefore, this torsion alone does not lead to a region of intersection with the S₄ state. The NO₂ torsion (Figure 2), however, points to an S_5/S_4 intersection at a relatively small torsion angle. The internal conversion to the S₄ state occurs mostly by CC stretching and NO₂ torsion (MXS (S_5/S_4) - see Figure 6c and Figure 8b), with contribution of CH₂ twisting. The CC bond lengthens from the ground state value of 1.33 Å to 1.47 Å and the CN bond shortens from 1.46 to1.37 Å. The energy of this MXS is stabilized by 0.4 eV as compared to the vertical excitation. To continue to the S_4/S_3 conical intersection (Figure 6b) requires significant torsions around the CH₂ and NO₂ groups (Figure 8a). In the MXS (S_4 / S₃) structure the C₂H₈ bond is now almost perpendicular to the planes of the CH₂ and NO₂ groups, with the CC and CN bond lengths decreasing and increasing, respectively, as compared to the MXS (S_5/S_4) . The computed potential curves confirm this picture: upon NO₂ (Figure 2) and CH₂ (Figure 3) torsion, the S_4 state slightly stabilizes. The MXS (S_4/S_3) is only 0.14 eV higher in energy than the MXS (S_5/S_4) and certainly accessible energetically within the deactivation dynamics. The S₃ state is also a bright state. Thus, it can be also populated directly through an excitation from the ground state even though with a smaller probability as compared to the S₅ state. The CH₂ torsion stabilizes the S3 state and leads to an intersection with S2. On the other hand, NO2 torsion destabilizes S3. These features are reflected in the structure of the MXS (S_3/S_2) (Figure 6a) where a rotation of the NO_2 group with respect to the MXS (S_4/S_3) is observed. Moreover, the CC bond is stretched in comparison with the latter MXS. The MXS (S_3/S_2) undergoes a significant energetic stabilization by 0.8 eV as compared to the MXS (S_4/S_3) .

The energies of the S_1 and S_2 states upon CH_2 torsion are energetically close in the whole range (Figure 3), particularly at the 90 °CH₂ torsion angle, a torsional region that is also assumed by the preceding MXS (S_3/S_2). In fact, the MXS (S_2/S_1) is characterized by a CH₂ torsion of ~90° (Figure 5c), though the NO₂ bending (see Figure 7c), combined with a relevant shortening of the CC bond and elongation of the CN and NO bonds, led to the S_1/S_0 intersection (Figure 7a).

The MXS (S_1/S_0) is characterized by a CH₂ torsion of ~90° similarly to the MXS (S_2/S_1) . As was already noted above, this intersection is not only achieved by this torsion but also by non-negligible changes of the CC, CN, and NO bond distances with respect to the ground state values suggestive of hot ground state dynamics after transition to S_0 . The energy of the MXS (S_1/S_0) is stabilized by ~0.4 eV in relation to the MXS (S_2/S_1) .

In view of the 3×3 CI analytical model as developed by Bonačić-Koutecký and Michl,^{37,38} the torsion of the CH_2 in nitroethylene from the planar geometry (Figure 3) reduces the S_1/S_0 gap as seen in the corresponding potential curves. A NO₂ torsion has little influence on the S_1/S_0 gap (Figure 2). This reduction according to that theory is due to an increased electronegativity difference of the CC bond upon CH₂ torsion. However, this gap reduction in nitroethylene is not as pronounced as, e.g., in fluorethylene. In the latter,³⁶ a rigid CC torsion reduces the gap to ~ 1 eV. In contrast, in nitroethylene a rigid CC torsion leads to a gap of ~2 eV and in ethylene to ~2.3 eV. However, as said above, a combination of bond changes, namely, a significant lengthening of the CC (0.15 Å) and one NO bond $(N_3O_5 = 0.14 \text{ Å})$, and also a shortening of the CN bond (-0.16 Å), as compared to the ground state geometry, takes place and is responsible for reaching the S_1/S_0 intersection. Moreover, the energy gradient difference g-vector of the MXS S_1/S_0 (Figure 7a) indicates possible NO₂ release through activation of the CN bond stretching. No pyramidalization of the CH₂ or NO₂ groups is observed. In this sense, the situation in nitroethylene can be considered similar to the polar fluoroethylene, with a significantly larger contribution of changes in the bond distances as compared to the ground state.

After a nonadiabatic transition, the vibrational energy acquired in the process by the molecule concentrates in the small number of vibrational modes that define the conical intersection.⁶⁴ Nitro compounds are usually thermally unstable and composed of easily broken covalent bonds such as N–N, O–C, and C–N. Release of the very labile NO₂ group in these molecules is thus rather probable in the hot ground state dynamics. The g-vector in the MXS (S_0/S_1) (Figure 7a), as discussed above, indicates the activation of a CN stretching mode. Therefore, it is expected that upon nonradiative transitions these modes might favor the release of the NO₂ group.

4. CONCLUSIONS

Extensive MR-CISD calculations were performed to study photodeactivation mechanisms of nitroethylene involving the first five valence excited states. Calculations of vertical excitations were carried out first. These calculations presented good agreement with our previous calculations and the experimental spectrum, excepting an inversion of the energy ordering of the last two transitions, the $n-\pi^*$ (S₄) and $\pi-\pi^*$ (S₅).

The potential curves for torsion and pyramidalization of the CH_2 group and the torsion of the NO_2 moieties were computed. Five conical intersections, one between every pair of adjacent states, were found: S_5/S_4 , S_4/S_3 , S_3/S_2 , S_2/S_1 , and S_1/S_0 .

The computed g- and h-vectors describing the shape of the intersection cone point to an important role played by the torsions of the CH_2 and NO_2 groups, with contributions of other modes such as CC and CN stretching. Similarities between the behavior of nitroethylene and previous work on the ethylene³¹ and fluorethylene³⁶ molecules were explored.

From the computed potential energies and the conical intersections found, a photochemical mechanism of nitroethylene deactivation could be proposed starting in the bright S_5 (π - π^*) excited state. After a Franck–Condon excitation to this state, a nonradiative chain of deactivation processes of nitroethylene from this state downward can proceed through conical intersections mostly characterized by twisting the CH₂, with important contributions from deformations of the NO₂ group and CC and CN bond stretches. Furthermore, the modes of the MXS (S_1/S_0) g-vector indicate the possibility of NO₂ release in the ground state, an important process in nitro molecules with explosive properties.

The results of this work have shed light on the photochemistry of nitroethylene, in particular on the importance and role of the conical intersections between adjacent pair of states. Moreover, these results will allow further chemical dynamics investigations on this molecule.

ASSOCIATED CONTENT

Supporting Information

MR-CISD absolute energies and the Cartesian coordinates of the conical intersections. This material is available free of charge via the Internet at http://pubs.acs.org.

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The Journal of Physical Chemistry A

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