Effect of an S_1/S_0 Conical Intersection on the Chemistry of Nitramide in Its Ground State. A Comparative CASPT2 Study of the Nitro-Nitrite Isomerization Reactions in Nitramide and Nitromethane

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The potential energy surfaces for the dissociation of nitramide $(NH_2NO_2 \rightarrow NH_2 + NO_2)$ and nitromethane $(CH_3NO_2 \rightarrow CH_3 + NO_2)$ and the nitro-nitrite rearrangement of these nitrocompounds $(RNO_2 \rightarrow RONO)$ as well as the dissociations of the nitrite isomers $(RONO \rightarrow RO + NO)$ have been studied with the second-order multiconfigurational perturbation theory (CASPT2) by computation of numerical energy gradients for stationary points. It is found that multiconfigurational methods [CASPT2] and complete active space SCF(CAS-SCF) predict that the isomerization of nitramide to NH_2ONO occurs in a two-step mechanism: (i) $NH_2NO_2 \rightarrow NH_2 + NO_2$ and (ii) $NH_2 + NO_2 \rightarrow NH_2ONO$, the second step involving surmounting an activation barrier. Contrastingly, Hartree-Fock based approaches give isomerization as a one-step reaction. Additionally, both mono- and multiconfigurational methods predict that nitro-nitrite rearrangement of CH_3NO_2 is a one-step process. The difference in the reaction mechanisms of these two isoelectronic molecules arises from the presence of an S_1/S_0 conical intersection in nitramide which is absent in nitromethane.

Introduction

It is firmly established that conical intersections are key features in the photochemistry of polyatomic molecules. ^{1,2} One of their most important implications is that internal conversion from upper to lower excited states occurs efficiently through or near these crossing points. On the other hand, it is commonly asserted that upward transitions from the lower surface to the upper surface should be less likely because there exists a peak in the lower surface which tends to guide trajectories away from the intersection. ^{3–5} Therefore, the role of surface crossings for the chemistry of molecules in the ground state should be less important. Otherwise, there are several cases in which conical intersections are relevant in the chemistry of molecules even in the ground state. These cases correspond to the so-called diabatic trapping effect, which implies nonadiabatic recrossing of the potential energy surfaces. ^{5–11}

The present work reports how the existence of an S_1/S_0 conical intersection determines the chemistry of nitramide (NH₂NO₂) in its ground state. The observed effect consists of the distortion of the ground-state potential energy surface by the conical intersection. Such a topological alteration of the surface extends to a wide region and leads to a dramatic change in the expected reaction mechanism. It must be noticed that a diabatic trap is not invoked at all.

To enlighten this effect, we must compare the reaction mechanisms for nitro-nitrite isomerizations of nitromethane (CH₃NO₂ \rightarrow CH₃ONO) and nitramide (NH₂NO₂ \rightarrow NH₂ONO). Great attention has been paid to these reactions in computational studies ^{12–20} since the pioneer work of Dewar et al. ¹² on the rearrangement of nitromethane to methylnitrite (CH₃ONO). The infrared multiphoton dissociation experiments of Wodtke, Hintsa, and Lee²¹ indicate that isomerization of nitromethane (eq 1) is a one-step reaction competing with C–N bond

dissociation (eq 2). A transition state exists in the isomerization reaction, while dissociation does not exhibit any exit energy barrier.

$$CH_3NO_2 \rightarrow CH_3ONO$$
 (1)

$$CH_3NO_2 \rightarrow CH_3 + NO_2$$
 (2)

The accepted reaction mechanisms for nitramides give the same pictures. That is

$$NH_2NO_2 \rightarrow NH_2ONO$$
 (3)

$$NH_2NO_2 \rightarrow NH_2 + NO_2$$
 (4)

In this study we demonstrate that the nitro-nitrite rearrangement of nitramide occurs in a two-step mechanism (5) that implies dissociation into $NH_2 + NO_2$ and subsequent recombination of the radicals after surmounting an energy barrier. In contrast, nitro-nitrite isomerization of nitromethane takes place in only a one-step reaction. The reason for this different mechanistic behavior of nitramide with respect to that of nitromethane arises from the presence of an S_1/S_0 conical intersection in nitramide which is absent in nitromethane.

$$NH_2NO_2 \rightarrow NH_2 + NO_2 \rightarrow NH_2ONO$$
 (5)

Computational Details

Generally contracted basis sets of atomic natural orbital (ANO) type obtained from C,N,O(14s9p4d3f)/H(8s4p3d) primitive sets,²² the so-called ANO-L basis sets, with the C,N,O-[4s3p2d1f]/H[3s2p1d] contraction schemes, were used in all of the geometry optimizations of the relevant species involved in the reactions of nitromethane and nitramide. These optimizations were performed following two strategies: (i) optimization at the complete active space self-consistent field (CAS-SCF)²³

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level of theory by computation of analytical energy gradients and (ii) optimizations with the second-order multiconfigurational perturbation theory (CASPT2)²⁴ by computation of numerical energy gradients. Both methods were applied as implemented in the MOLCAS 6.2 program.²⁵ In the CASPT2 calculation, the 1s electrons of the carbon, oxygen, and nitrogen atoms, as determined in the SCF calculations, were kept frozen. On the other hand, the localization of the crossing points was done with the algorithm implemented in the GAUSSIAN program²⁶ and with the cc-pVDZ basis sets.²⁷ The gradient difference and nonadiabatic coupling vectors are computed by using state average orbitals in the manner suggested by Yarkony.²⁸

The stationary points (minima and saddle points) were characterized by their CASSCF analytic harmonic vibrational frequencies computed by diagonalizing the mass-weighted Cartesian force constant matrix.

The selection of the active space plays a critical role in the CAS–SCF studies of nitromethane and nitramide. If we want to avoid undesirable effects such as symmetry breaking when the systems move on the potential energy surfaces, the minimum active space must comprise 14 electrons distributed in 11 orbitals. These orbitals correspond to $2s_{\rm N}$, two NO bonding orbitals σ NO, one delocalized ONO π -bonding orbital π NO, one σ nonbonding orbital n σ O, one π nonbonding orbital n π O, two NO antibonding orbitals $\sigma^*_{\rm NO}$, one delocalized ONO π -antibonding orbital $\pi^*_{\rm NO}$, plus CN bonding and antibonding orbitals σ CN and $\sigma^*_{\rm CN}$ on CH3NO2 or NN bonding and antibonding orbitals σ NN and $\sigma^*_{\rm NN}$ orbitals on NH2NO2, respectively.

Results and Discussion

This section is organized as follows. Prior to discussion of the reaction mechanisms, all of the critical points found in this study are illustrated in Figure 1, which have been plotted with the MacMolplt program. Second, it is shown that the dissociation reactions of the four isoelectronic molecules dealt with in this work (CH₃NO₂, CH₃ONO, NH₂NO₂, and NH₂ONO) occur without any exit energy barrier. Third, the nitro-nitrite isomerization reactions of nitromethane (CH₃NO₂ \rightarrow CH₃ONO) and nitramide (NH₂NO₂ \rightarrow NH₂ONO) are discussed, respectively. Comparison of both reaction mechanisms will allow us to obtain the main conclusion of this work.

Conformers on the Potential Energy Surfaces of the Nitro and Nitrite Compounds. Nitromethane has two C_s conformers: (i) staggered (M1, Figure 1a) and (ii) eclipsed. They are almost isoenergetic but with the staggered structure slightly favored. Methylnitrite has four conformational isomers with C_s symmetry depending on whether the terminal NO and in-plane hydrogen are placed cis or trans with respect to the central NO bond: (i) trans—trans (M2, Figure 1b), (ii) trans—cis, (iii) cis—trans, and (iv) cis—cis. The cis—trans conformer is energetically the most stable one (Table 1). The cis—cis isomer is a saddle point on the potential energy surface at the CAS—SCF level.

Nitramide has only one C_s conformer (M3, Figure 1d). As methylnitrite, there are four isomers of NH₂ONO depending on whether the terminal NO and hydrogens are placed cis or trans with respect to the central NO bond. The cis—cis isomer is the most stable one (Table 2). In Figure 1e we show the trans—trans conformer.

NO₂ and **NO** Eliminations. We have studied dissociations of the nitro and nitrite compounds by calculating the energy profiles of the reactions along linear interpolations as in previous works. ^{29,31} That is, an interpolation vector, $\Delta \mathbf{R}$, which connects ground states of stable molecules and their respective dissocia-

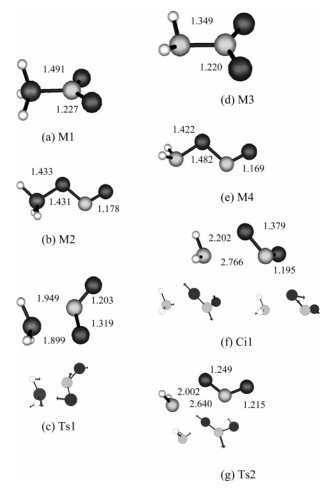


Figure 1. CASPT2 geometries of the relevant species for the decomposition and isomerization of both nitromethane and nitramide: (a) staggered nitromethane (M1); (b) trans—trans methyl nitrate (M2); (c) transition state for M1 \rightarrow M2 isomerization (Ts1), the arrows in the smaller figures correspond to the transition vector; (d) nitramide (M3); (e) trans—trans NH₂ONO (M4); (f) S1/S0 conical interaction (Ci1), the arrows in the smaller figures correspond to nonadiabatic coupling (left) and gradient difference (right) vectors; (g) transition state for M3 \rightarrow M4 isomerization (Ts2), the arrows in the smaller figures correspond to the transition vector.

tion products, is obtained by calculating the difference between internal coordinates of reactant and dissociation products, $\Delta R = R_i - R_j$, where R_i and R_j represent the internal coordinate vectors of the initial and dissociation products, respectively. Units of the interpolation vector are measured in Å for internuclear distances and degrees for valence and dihedral angles.

Figure 2 displays the energy profiles for dissociation of CH_3 - NO_2 and CH_3ONO into $CH_3 + NO_2$ and $CH_3O + NO$, respectively. The S_1 state for dissociation of CH_3ONO is also plotted in the same figure, which shows that S_1 is dissociative.

Figure 3 shows the energy profiles for dissociation of NH_2 - NO_2 and NH_2ONO into $NH_2 + NO_2$ and $NH_2O + NO$, respectively. As occurs in CH_3ONO , the S_1 state of NH_2ONO results in being dissociative.

The two-dimensional plots given in Figures 2 and 3 are not true reaction paths but cuts of the multidimensional potential energy surfaces along an arbitrary direction. However, provided that no barrier is found along these dissociation interpolations, it should not exist along the true energy paths. In other words, there will not be saddle points related to transition states on the potential energy surfaces.

TABLE 1: Energetic of the Critical Points on the S₀ Potential Energy Surfaces of CH₃NO₂^a

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description	CASPT2 ^a	$ZPE^{b,c}$	$TC^{c,d}$	ΔH^e	exp.h
s-CH ₃ NO ₂	-244.620 77	29.2	2.3	0.0	
		$(30.2)^g$	(2.2)	(0.0)	
e-CH ₃ NO ₂	-244.62079	29.2	2.8	0.6	
t-t-CH ₃ ONO, m ^j	-244.61224	28.1	3.2	5.2	
t-c-CH ₃ ONO, m ^j	-244.61265	28.1	2.6	4.3	
c-t-CH ₃ ONO, m ^j	-244.61449	28.5	2.9	3.9	1.36
c- c -CH ₃ ONO, sd ^{k}	-244.61154	28.2	2.5	5.1	
Ts1	-244.50964	26.6	3.0	67.8	
$CH_3 + NO_2^f$	-244.51966	22.0	4.0	58.6	60.89^{i}
		(23.5)	(3.7)	(58.8)	
$CH_3O + NO^f$	-244.54208	23.0	3.7	43.9	41.87^{i}

^a CASPT2/ANO-L energy in hartrees. ^b Zero point energy in kcal/ mol. ^c Vibrational frequencies scaled by 0.90. ^d Thermal correction (298.15 K) in kcal/mol. e Relative enthalpies in kcal/mol with respect to the ground state of s-CH₃NO₂. ^f Distance between fragments 10 Å. g In parentheses values obtained including experimental frequencies. ^h From: In CRC Handbook of Chemistry and Physics, 67th ed. 1986-1987; Weast, R. C., Ed.; CRC Press: Boca Raton, FL, 1986. ⁱ Bond dissociation energy calculated as $D^{\circ} = \sum \Delta_{H}^{\circ}(\text{product}) - \Delta_{H}^{\circ}(\text{reactant})$. ^j m: minimum. ^k sd: saddle point.

TABLE 2: Energetic of the Critical Points on the S_0 Potential Energy Surfaces of NH₂NO₂^a

description	$CASPT2^a$	$ZPE^{b,c}$	$TC^{c,d}$	ΔH^e
NH ₂ NO ₂	-260.64270	23.2	2.5	0.0
t-t-NH ₂ ONO, m ⁱ	-260.60458	21.6	3.0	22.9
t-c-NH ₂ ONO, m ⁱ	-260.60356	21.6	3.0	23.5
c-t-NH ₂ ONO, m ⁱ	-260.60558	21.5	3.0	22.2
c-c-NH ₂ ONO, m ⁱ	-260.60783	21.8	2.8	20.9
Ts2	-260.54142	17.6	3.5	58.9
$Ci1 S_1/S_0$	$-260.515\ 10^{f}$	-	-	80.5
$NH_2 + NO_2^g$	$-260.557\ 13$	15.3	3.6	47.6
$NH_2O + NO^g$	-260.58148^{h}	17.5	3.6	36.2

^a CASPT2/ANO-L energy in hartrees. ^b Zero point energy in kcal/ mol. ^c Vibrational frequencies scaled by 0.90. ^d Thermal correction (298.15 K) in kcal/mol. e Relative enthalpies in kcal/mol with respect to the ground state of s-NH₂NO₂ f Multistate-CASPT2 energy referred to -260.643 42 hartrees [reference wave function CAS(14,12), 2 roots]. g Distance between fragments 10 Å. h Multistate-CASPT2 energy referred to -260.645 40 hartrees [reference wave function CAS(16,12), 4 roots]. i m: minimum.

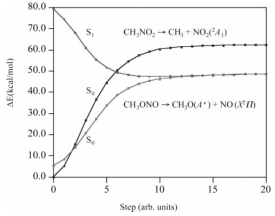


Figure 2. Singlet CASPT2/ANO-L dissociation curves (C_s symmetry) of nitromethane (S₀) and trans-trans methyl nitrate (S₀ and S₁). The geometry of the initial points corresponds to the minima of nitromethane and methyl nitrite, respectively.

NO₂-ONO Rearrangement of CH₃NO₂ and NH₂NO₂. Isomerization of nitromethane (M1, Figure 1a) to methylnitrite (M2, Figure 1b) occurs in a one-step reaction after passing through a transition state (Ts1, Figure 1c) lying 68 kcal/mol above the S₀ minimum of nitromethane. The intrinsic reaction

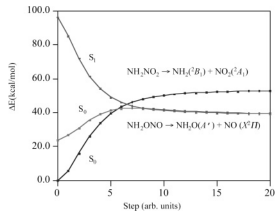


Figure 3. Singlet CASPT2/ANO-L dissociation curves (C_s symmetry) of nitramide (S₀) and trans-trans NH₂ONO (S₀ and S₁). The geometry of the initial points corresponds to the minima of nitramide and transtrans NH2ONO, respectively.

coordinate (IRC) starting at Ts1 leads to nitromethane and methylnitrite, respectively (Figure S1). The geometry and energertic of this CASPT2 transition state is in agreement with the structures reported by Nguyen et al. 16 using the CCSD(T) and CAS-SCF methods.

Nitro-nitrite rearrangement in nitramide deserves much attention. We were able to find the transition state for isomerization of nitramide (M3, Figure 1d) to NH2ONO (M4, Figure 1e) by applying different approaches: (i) restricted Hartree-Fock (RHF), (ii) restricted density functional theory (DFT), and (iii) restricted Møller-Plesset (MP2). The geometrical parameters of these transition states are collected in Table S1. As an example, Figure S2 shows the energy profile of the IRC starting at the saddle point found with the DFT/B3-LYP method. This IRC clearly demonstrates that this structure is the transition state for the nitro-nitrite rearrangement of nitramide at this level of theory.

Surprisingly, despite an exhaustive search, it was impossible at all to find the nitro-nitrite transition state of nitramide at the CAS-SCF and CASPT2 levels of theory. We expected to find such a transition state close to the geometry predicted by DFT methods. However, what we found in the neighboring of this region was an S₁/S₀ conical intersection (Ci1, Figure 1f). Minimum energy paths starting at this conical intersection and following both directions of the gradient difference and nonadiabatic coupling vectors always led the system to NH₂ONO or dissociation products (NH₂ + NO₂), alternately.

To clarify the discrepancy found between multiconfigurational and Hartree-Fock based methods, the reduced three-dimensional potential energy surfaces for the isomerization reaction of nitramide have been built at the state averaged CAS-SCF level of theory including both the S_0 and S_1 states (Figure 4a) and applying the N,O[3s2p1d]/H[2s1p] contraction schemes to the ANO-L basis sets. The points which generate these surfaces span the geometries of four critical molecular arrangements: (i) nitramide (M3), (ii) S₁/S₀ conical intersection (Ci1), (iii) NH₂-ONO (M4), and (iv) dissociation products ($NH_2 + NO_2$). The most striking feature of this figure is the S₁/S₀ conical intersection. Additionally, a reaction path leading to the dissociation of nitramide into $NH_2 + NO_2$ on the S_0 surface is observed, but no transition state appears for the direct isomerization. The reason for this behavior can be attributed to the presence of such a conical intersection, which repels down the S₀ surface destroying the transition state for isomerization. On the other hand, the S₀ surface exhibits a saddle point near the conical intersection, which is located approximately along the

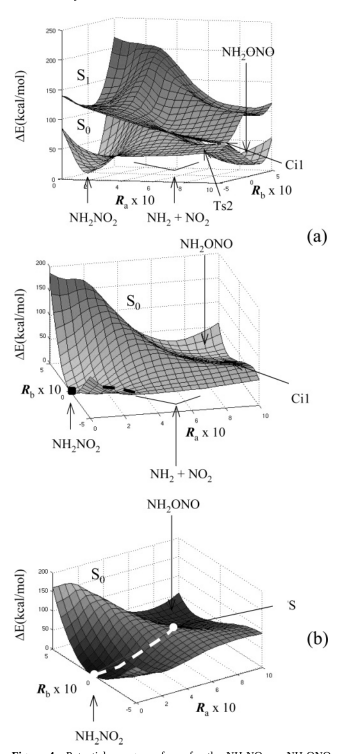


Figure 4. Potential energy surfaces for the NH₂NO₂ \rightarrow NH₂ONO reaction (a) (upper) state average CAS-SCF/ANO-L (C,N,O[3s2p1d]/H[2s1p]) surfaces of the S₁ and S₀ states showing the S₁/S₀ conical intersection and (lower) S₀ surface from a different perspective and (b) S₀ B3-LYP/aug-cc-pvdz surface showing a saddle point between NH₂NO₂ and NH₂ONO.

line defined by NH₂ONO and dissociation products. Thus, we searched for and localized a transition state in the vicinity of this region (Ts2, Figure 1g). The IRC starting at this stationary point demonstrates that it is the transition state for the bimolecular reaction NH₂ + NO₂ \rightarrow NH₂ONO (Figure S3).

It is worthy to note that surfaces displayed in Figure 4 are built from a set of perfectly ordered points. The two orthogonal directions are obtained as follows:

(1) Two generator vectors are defined in internal coordinates as

$$\mathbf{R}_1 = \mathbf{R}(\mathrm{NH}_2\mathrm{NO}_2) - \mathbf{R}(\mathrm{NH}_2\mathrm{ONO}) \tag{6a}$$

$$\mathbf{R}_2 = \mathbf{R}(\mathrm{NH}_2\mathrm{NO}_2) - \mathbf{R}(\mathrm{Ci1}) \tag{6b}$$

where R(I) denotes the internal coordinate vector of a given species.

(2) Vectors \mathbf{R}_1 and \mathbf{R}_2 are normalized

$$\boldsymbol{r}_1 = \frac{\boldsymbol{R}_1}{R_1} \tag{7}$$

$$r_2 = \frac{R_2}{R_2} \tag{8}$$

(3) Sum and subtraction of r_1 and r_2 give a set of two orthonormal vectors

$$\boldsymbol{r}_a = \boldsymbol{r}_1 + \boldsymbol{r}_2 \tag{9}$$

$$\boldsymbol{r}_b = \boldsymbol{r}_1 - \boldsymbol{r}_2 \tag{10}$$

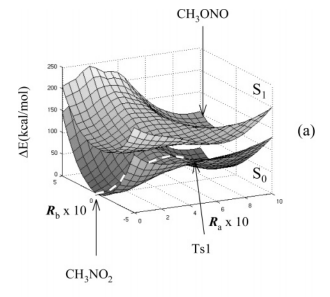
(4) Vectors \mathbf{r}_a and \mathbf{r}_b are scaled by the length of the largest vector between \mathbf{R}_1 and \mathbf{R}_2 , which is \mathbf{R}_2 in this case

$$\mathbf{R}_a = R_2 \cdot \mathbf{r}_a \tag{11}$$

$$\mathbf{R}_b = R_2 \cdot \mathbf{r}_b \tag{12}$$

The B3-LYP/aug-cc-pVDZ potential energy surface of the nitro-nitrite isomerization of nitramide (Figure 4b) has been built with the same geometrical points used in Figure 4a. Although the geometry of the transition state has not been specifically included, it is observed for the reaction path for direct isomerization passing through a saddle point in such a figure. This fact strengthens our confidence in that the reduced potential energy surfaces are representative of the reaction mechanisms. That is, if the transition state for isomerization does exist at the CAS-SCF level, we had been indeed able to find it. There are several reasons to think that the picture given for the isomerization process by the methods based on monoconfigurational wave functions are nothing but artifacts. First, the weight of the Hartree-Fock configuration on the multiconfigurational wave function amounts to \sim 50% around the region of the conical intersection, just where restricted approaches localize the transition state. Second, these methods give an incorrect dissociation limit which topologically results on an overestimation of the energy of the lower right corner of Figure 4b. This fact, in connection with the two minima corresponding to NH₂-NO₂ and NH₂ONO, creates an artificial saddle point. Third, monoconfigurational methods are blind to the S_1 state.

The same procedure has been applied to build the potential energy surfaces of the nitro-nitrite rearrangement of nitromethane (Figure 5). In this case, generator vectors (\mathbf{R}_1 and \mathbf{R}_2) are obtained as $\mathbf{R}_1 = \mathbf{R}(\mathrm{CH_3ONO}) - \mathbf{R}(\mathrm{CH_3NO_2})$ and $\mathbf{R}_2 = \mathbf{R}(\mathrm{Ts1}) - \mathbf{R}(\mathrm{CH_3NO_2})$, respectively. Figure 5 shows the S_1 and S_0 potential energy surfaces built at the state average CAS—SCF level applying the C,N,O[3s2p1d]/H[2s1p] contraction schemes to the ANO-L basis sets. It is observed that the saddle point related to the transition state for direct isomerization is actually well defined. Likewise, it is important to notice that S_0 and S_1 states do not touch each other in the whole region spanned by the surfaces.



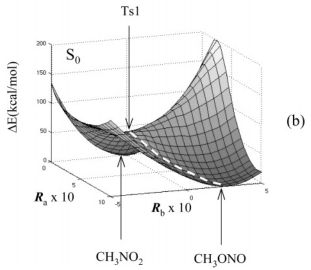


Figure 5. Potential energy surfaces for the $CH_2NO_2 \rightarrow CH_3ONO$ reaction (a) state average CAS-SCF/ANO-L (C,N,O[3s2p1d]/H[2s1p]) surfaces of the S_1 and S_0 states and (b) S_0 surface from a different perspective.

To finish this section, we have checked that dynamic correlation effects do not largely affect the conical intersection localized at the CAS-SCF level of theory. Thus, we have performed single point CASPT2 calculations at the most significant geometries and have found that the energy gap between the S_0 and S_1 surfaces is still small (~ 1 kcal/mol) at the CASPT2 level. These calculations are illustrated in Figure S4 that shows the energy profile connecting NH₂NO₂ minimum with Ci1.

Conclusions

In the present work we have investigated the potential energy surfaces related to nitro-nitrite isomerization of nitramide and nitromethane, respectively. The values predicted by the CASPT2 method for the geometrical parameters of nitromethane and methylnitrite are in excellent agreement with the experimental parameters.32,33 More discrepancies between theoretical and experimental geometries are found in the case of nitramide.³⁴ Tables of the CASPT2 geometrical parameters of all the species considered in this work are given in the Supporting Information (Tables S2-S5). Additionally, we have studied the stability of the R-NO2 and RO-NO bonds for both nitro and nitrite compounds. It is found that the latter one is rather weak. Therefore, whether nitrite isomer is formed in a direct reaction or not, it rapidly decomposes to RO + NO.

The main conclusion of this study is that there is no minimum energy path for rearrangement of nitramide to NH₂ONO. NH₂-ONO is formed after recombination of the NH₂ and NO₂ radicals. Furthermore, such a recombination is not barrierless. It is worthy to notice that it should be dynamically possible to find a direct isomerization path without involving a transition state, as it is suggested in recent articles.^{35–37} The proposed reaction mechanism for nitramide differs from that found in nitromethane, which certainly can isomerize to methylnitrite in a one-step reaction. The reason for these different mechanistic behaviors can be understood by comparing the S_0 and S_1 potential energy surfaces of the NH₂NO₂-NH₂ONO and CH₃- NO_2 -CH₃ONO systems. The presence of an S_1/S_0 conical intersection in the NH₂NO₂-NH₂ONO system, which is absent in CH₃NO₂-CH₃ONO, causes a topological alteration of the potential energy surface destroying the minimum energy path for nitro-nitrite isomerization of NH₂NO₂.

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Supporting Information Available: CASPT2 tables of energetic and geometrical parameters, intrinsic reaction coordinates, and linear interpolations. This material is available free of charge via the Internet at http://pubs.acs.org.

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