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A multireference perturbation study of the NN stretching frequency of trans-azobenzene in nπ∗ excitation and an implication for the photoisomerization mechanism

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A multireference second-order perturbation theory is applied to calculate equilibrium structures and vibrational frequencies of trans-azobenzene in the ground and nπ∗ excited states, as well as the reaction pathways for rotation and inversion mechanism in the nπ∗ excited state. It is found that the NN stretching frequency exhibits a slight increase at the minimum energy structure in the nπ∗ state, which is explained by the mixing of the NN stretching mode with the CN symmetric stretching mode. We also calculate the NN stretching frequency at several selected structures along the rotation and inversion pathways in the nπ∗ state, and show that the frequency decreases gradually along the rotation pathway while it increases by ca. 300 cm⁻¹ along the inversion pathway. The frequencies and energy variations along the respective pathways indicate that the rotation pathway is more consistent with the experimental observation of the NN stretching frequency in nπ∗ excitation. © 2013 American Institute of Physics. [http://dx.doi.org/10.1063/1.4790611]

I. INTRODUCTION

The photoisomerization of azobenzene has attracted a great deal of interests in both science and industry since the photoisomerization can be used as light-driven nanoscaled devices such as optical switches, and a great number of experimental studies have been reported. An earlier study on the absorption spectra indicated that the first and second excited states of both cis- and trans-azobenzene were attributed to ππ∗ and ππ∗ excitations, respectively. Quantum yields of the isomerizations for cis- and trans-azobenzene were reported to be different between nπ∗ and ππ∗ excitations. Rau and Luddeke proposed that the photoisomerization of trans-azobenzene should follow different pathways in nπ∗ and ππ∗ excitations, through a comparison of the simple azobenzene and the sterically hindered azobenzenes. It was reported that the nπ∗ excitation invokes the inversion of the NNC bond angle within a planar geometry, while the ππ∗ excitation excitation invokes the rotation of two phenyl rings around the NN bond (Figure 1). Ultrafast UV-visible absorption spectroscopy and time resolved fluorescence experiments on trans-azobenzene suggested that decay in nπ∗ excitation can be described by three processes: (1) relaxation from the Franck-Condon region to the minimum in the S₁(nπ∗) state, (2) motion toward S₁/S₀ conical intersections, and (3) vibrational cooling in the S₀ state through an interaction with solvent molecules.

There have been several transient absorption experimental studies for the ππ∗ excitation of trans- and cis-azobenzene. Tahara and co-workers carried out the time-resolved Raman, fluorescence, and absorption measurements with the ππ∗ excitation of trans-azobenzene in solution. They clarified that isomerization proceeds in the ππ∗ state after electronic relaxation, and proposed that the isomerization pathway is the same as that of direct nπ∗ excitation. They also found that the NN stretching frequency shows only a slight decrease in the S₁(nπ∗) state.

Monti et al. showed that in the S₁(nπ∗) state there is a large activation barrier along the rotation pathway while there is no barrier along the inversion pathway at the singly-excitation configuration interaction (CIS) level, indicating that the photoisomerization of trans- and cis-azobenzene is likely to proceed via the inversion pathway in nπ∗ excitation. Monti et al. also suggested that, in ππ∗ excitation, the preferred pathway is a rotational route, which leads to either the photoisomerization via S₁(nπ∗) or no isomerization with direct transition to S₀.

In 1999, complete active space self-consistent field (CASSCF) method was employed to determine the rotation and inversion pathways between trans- and cis-azobenzene in the respective S₁(nπ∗) and S₂(ππ∗) states, in which the adiabatic potential energy was calculated with a multireference perturbation theory along the CASSCF reaction pathways. It was concluded that the inversion is probably the preferred pathway for nπ∗ excitation while the rotation is preferred for ππ∗ excitation.

Ishikawa and co-workers examined two-dimensional potential energy surfaces (PES) of the S₁(nπ∗) state as a function of the CNNC torsion and NNC bond angles at the multi-reference single and double excitation configuration...
interaction (MR-SDCI) level, and found an S1/S0 conical intersection along the rotation pathway, indicating that the rotation pathway is preferred in the S1(nπ∗) state. After Ishikawa’s report,24 all theoretical studies at the CASSCF, multireference complete-active-space second-order perturbation theory (CASSPT2), second-order approximated coupled-cluster model with the resolution-of-the-identity approximation (RI-CC2), and time-dependent density functional theory (TDDFT) levels indicate that the rotation is the preferred pathway for nπ∗ excitation in the gas phase.27,28,30,31,42,58

On-the-fly dynamics simulations were also performed for the photoisomerization of azobenzene on the basis of semiempirical molecular orbital calculations with the surface hopping method29,33,55,61 and with the multiple spawning method.32 Recently, ab initio molecular dynamics (AIMD) simulations at the CASSCF level26,47,54,57 and Car-Parrinello molecular dynamics simulations34,45,49,50 were also performed for the photoisomerization of azobenzene in nπ∗ excitation. In our surface hopping AIMD simulation67 at the state-averaged CASSCF (SA-CASSCF) level, it was shown that cis to trans isomerization in nπ∗ excitation occurs via two-step rotation mechanism, accompanying rotations of the central NN part and two phenyl rings, and this process can be classified into two types with respect to the orientation of the rotation, namely, clockwise and counterclockwise rotation pathways; the calculated quantum yields and lifetime in the excited states are in very good agreement with the corresponding experimental results.47 The similar reaction mechanism was also reported in the dynamics simulations by Doltsinis et al.45,49,50 and Thiel et al.55

As described above, the transient Raman spectra indicate that the NN stretching frequency of trans-azobenzene is almost unchanged (decreases by only 12 cm−1) in the S1(nπ∗) state.12 There are several theoretical reports on vibrational frequencies for trans-azobenzene in the ground state at the MP2, density functional theory (DFT), and CASSCF levels,20,21,23,25,30 while, to our knowledge, there is only one report on frequencies of trans-azobenzene in the S1(nπ∗) state, which employed the CASSCF method.30 However, the CASSCF method is sometimes insufficient for the quantitative discussion of geometries and frequencies because dynamical correlation effects are not taken into account.52,63

The goal of the current article has two-folds. One is to clarify the origin of small frequency shift in the NN stretching in the nπ∗ state, and another is to show likely mechanism for trans-cis photoisomerization.

In order to gain insight to the photoisomerization mechanism of trans-azobenzene with reference to the Raman spectra, we employ the CASPT2 method to determine stationary points and vibrational frequencies for trans-azobenzene in both S0 and S1(nπ∗) states. This is the first attempt to discuss vibrational frequencies for the excited state of azobenzene by ab initio multireference theory.

II. COMPUTATIONAL DETAILS

Geometry optimizations and normal mode analyses were performed for trans-azobenzene in both S0 and S1 states under the C2h symmetry restriction. In the S0 state, DFT (B3LYP and CAM-B3LYP), Møller-Plesset second-order perturbation theory (MP2), quadratic configuration interaction with singles and doubles including triples perturbationally (QCISD(T)), SA-CASSCF, and CASPT2 methods were employed, while in the S1(nπ∗) state TDDFT (B3LYP and CAM-B3LYP), SA-CASSCF, and CASPT2 methods were used. Normal mode analyses were also performed for an isomer of trans-azobenzene at optimized structures in both S0 and S1 states by substituting 14N with 15N to compare with the Raman spectra.12 To examine the anharmonic effect, the direct vibrational self-consistent field (VSCF) and the following correlation-corrected VSCF (cc-VSCF) methods66,65 were applied to both S0 and S1 states with quartic force field option66 at the DFT(B3LYP)/6-31G* level.

In SA-CASSCF calculations, the S0 and S1(nπ∗) states are averaged with an equal weight, and the active space is chosen as 14 electrons in 12 orbitals (14,12), which includes two n orbitals and π/π∗ orbitals of nitrogens, and four sets of π/π∗ orbitals of phenyl rings. In CASPT2 calculations, the reference SA-CASSCF wavefunction is determined so that the active space includes two n orbitals and π/π∗ orbitals of nitrogens ((6,4) active space); the lowest 17 orbitals are treated as frozen orbitals in the perturbation calculations for dynamic electron correlation. The active space of the reference CASSCF wavefunction for CASPT2 calculations was determined carefully to get the reliable results with available computational resources, through preliminary calculations; we verified that the equilibrium geometry in the ground state determined at the CASPT2 level with the CASSCF(6,4) reference is very similar to the equilibrium geometry determined at the CASPT2 level with the CASSCF(10,8) reference.67

We also investigated the trans-cis isomerization via the rotation pathway as a function of a CNCN torsion angle dCNCN and the inversion pathway as a function of an NNC bond angle dNNC on the S1(nπ∗) PES. Optimization of the geometrical parameters was carried out by fixing dCNCN to be 180◦ for the inversion pathway. Then, we calculated the NN stretching frequency along the respective pathways by applying a projection technique68 at the CASPT2 level.

In the present calculations, Sapporo-DZP basis set69 were employed except for VSFC/cc-VSCF calculations. DFT calculations were performed by the GAMESS program package,70 while MP2, CASPT2, and QCISD(T) calculations were performed by the MOLPRO2010 program package.71

III. RESULTS AND DISCUSSION

Table I shows calculated and experimental values for geometrical parameters and the NN stretching frequency at the

FIG. 1. A scheme of rotation and inversion mechanism for cis-trans isomerization of azobenzene.
equilibrium structure in the ground state of trans-azo benzene (referred to as \((S_0)_{\text{min}}\)). The calculated geometrical parameters show good agreements with the experimental values so that the maximum differences are \(\Delta r_{\text{NN}} \sim 0.032\ \text{Å}, \Delta r_{\text{CN}} \sim 0.012\ \text{Å}, \) and \(\Delta \theta_{\text{NNC}} \sim 0.9^\circ\). The NN stretching frequency \(v_{\text{NN}}\) is overestimated by ca. 80 cm\(^{-1}\) relative to the fundamental frequency at the CASPT2 and QCISD(T) level, while it is underestimated by 23 cm\(^{-1}\) at the MP2 level. The cc-VSCF calculations at the DFT(B3LYP)/6–31G\(^*\) level show that the harmonic and fundamental frequencies are 1561 and 1531 cm\(^{-1}\), respectively, in the S\(_0\) state, and thus the anharmonicity reduces the frequency by 30 cm\(^{-1}\). By using this value in the other computational levels, a fundamental frequency for the NN stretching mode is estimated to be 1490 and 1492 cm\(^{-1}\) at the CASPT2 and QCISD(T) levels, respectively, which is comparable to the experimental value, 1440 cm\(^{-1}\). There are several theoretical reports on equilibrium geometries and vibrational frequencies for trans-azo benzene in the ground state at the MP2, DFT, and CASCCSF levels.\(^{20,21,23,25,30}\) In an early study by Armstrong et al.,\(^{20}\) \(r_{\text{NN}}\) and \(v_{\text{NN}}\) were calculated to be 1.310 Å (\(\Delta r_{\text{NN}} \sim 0.063\ \text{Å}\)) and 1296 cm\(^{-1}\), respectively, at the MP2/6–31G level, while by replacing the basis set of nitrogen atoms with 6–31+G(d), \(r_{\text{NN}}\) and \(v_{\text{NN}}\) were calculated to be 1.275 Å (\(\Delta r_{\text{NN}} \sim 0.028\ \text{Å}\)) and 1450 cm\(^{-1}\), respectively, so the addition of diffuse functions on nitrogen atoms improves the NN bond strength extensively.

For the \(n\pi^*\) excited state, geometry optimization was performed under the C\(_{2h}\) symmetry constraint (the obtained structure is referred to as \((S_1)_{\text{C2hmin}}\)). Normal mode analyses indicate that \((S_1)_{\text{C2hmin}}\) has one imaginary frequency mode, which corresponds to a torsional motion of the central -CNNC- (A\(_a\) symmetry) at DFT(CAM-B3LYP) (20i cm\(^{-1}\)) and CASPT2 (49i cm\(^{-1}\)) levels, while it has no imaginary frequency mode at the DFT(B3LYP) and SA-CASSCF levels. To examine instability of the planar geometry of trans-azo benzene in the \((S_1(n\pi^*))\) state further, we performed additional DFT (B3LYP and CAM-B3LYP) and CASPT2 calculations using Sapporo-DZP plus diffuse functions, and found that the inclusion of diffuse functions does not change existence of the imaginary frequency mode. This result indicates that the planar structure of \((S_1)_{\text{C2hmin}}\) is unstable with respect to the CNNC torsional motion, although an absolute value of these imaginary frequencies is not so large.

We will come back to instability of the planar geometry later.

Table II shows geometrical parameters and the NN stretching frequency at \((S_1)_{\text{C2hmin}}\), as well as \(n\pi^*\) vertical and adiabatic excitation energies within the C\(_{2h}\) coordinate space. The calculated vertical excitation energy is in good agreement with the experimental value.\(^{13}\) The comparison of geometrical parameters between S\(_0\) and \((S_1(n\pi^*))\) indicates that, by \(n\pi^*\) excitation, the NN bond length is almost unchanged while the NNC bond length increases by ca. 14° and the CN bond length decreases by ca. 0.06 Å. These geometrical changes can be explained by nature of molecular orbitals related to \(n\pi^*\) excitation. Figure 2 shows natural orbitals relevant to \(n\pi^*\) excitation, i.e., the lone pair n orbital of N atoms and the NN-π orbital shows anti-bonding nature for the NN bond, while the \(\pi^*\) orbital shows anti-bonding nature for the NN bond and bonding nature for two CN bonds; thus, due to \(n\pi^*\) excitation, CN bonds should be strengthened while NN bond is not so affected. An increase in the NNC bond angle can be understood by noting the reduction of a repulsive force between bonding electron pairs (NN and CN bonds) and the lone pair of N atoms, which follows a valence shell electron pair repulsion (VSEPR) rule. A similar finding in geometrical changes of trans-azo benzene due to \(n\pi^*\) excitation was also reported very recently at the semiempirical OM2/MRCI level.\(^{61}\)

Experimentally, vibrational frequency was also measured for the isotopomer of trans-azo benzene (where \(^1\)N is

![FIG. 2. Natural orbitals corresponding to (a) the lone pair orbital of nitrogen and (b) the NN anti-bonding \(\pi^*\) orbital of trans-azo benzene related to \(n\pi^*\) excitation determined by the SA-CASSCF(6,4) calculation at \((S_0)_{\text{min}}\).](image)
substituted by $^{15}$N), and the isotopic shift in the NN stretching frequency was reported to be $-27$ and $-29$ cm$^{-1}$ in the S$_0$ and S$_1$ states, respectively. The corresponding isotopic shifts are calculated to be $-31$ and $-37$ cm$^{-1}$ in the S$_0$ and S$_1$ states, respectively, at the DFT(B3LYP)/DZP level, which are in good agreement with the experimental values. This agreement indicates that the experimentally observed frequency in the Raman spectra$^{12}$ is definitively ascribed to the NN stretching mode. As shown in Tables I and II, due to $n\pi^*$ excitation, the experimental NN stretching frequency decreases by 12 cm$^{-1}$, while the calculated frequency increases by 116 cm$^{-1}$ (TDDFT(B3LYP)), 90 cm$^{-1}$ (TDDFT(CAM-B3LYP)), 76 cm$^{-1}$ (CASSCF), and 61 cm$^{-1}$ (CASPT2). The anharmonicity effect in the NN stretching frequency is estimated to be 29 cm$^{-1}$ (1679 → 1650 cm$^{-1}$) in the S$_1$ state as indicated by the cc-VSCF calculation at the TDDFT(B3LYP)/6-31G$^\text{d}$ level. The anharmonicity effect is similar to that in the ground state, and thus, change of the NN stretching fundamental frequency via S$_0$ → S$_1$ excitation seems to be similar to the change of the harmonic one.

To our knowledge, there is one theoretical report on vibrational frequencies of trans-azobenzene in the S$_1(n\pi^*)$ state, which state-specific CASSCF calculations were performed to determine minimum energy structures and harmonic frequencies for the ground and S$_1(n\pi^*)$ states under the C$_2h$ symmetry restriction, using the analytical gradient and analytical Hessian matrix, implemented in the MOLCAS-5.2 program. The calculated frequencies, multiplied by scaling factor 0.91 to take into account effects of dynamic correlation and anharmonicity, were reported as 1440 cm$^{-1}$ (S$_0$) and 1438 cm$^{-1}$ (S$_1$), which were in good agreement with the experimental values. Since their results are totally different from our calculated values at the CASSCF/DZP level, we also performed geometry optimization and normal mode analyses by the same method, namely, the state-specific CASSCF with the (14,12) active space and the atomic natural orbital (ANO) basis sets, using the MOLCAS-6.4 program. The calculated geometries and frequencies are almost identical to those in Ref. 30, and we have noticed, however, that the assignment for the NN stretching frequency at the CASSCF level in Ref. 30 is erroneous. In our CASSCF calculations, the frequencies for the NN stretching mode are calculated to be 1701 cm$^{-1}$ (S$_0$) and 1771 cm$^{-1}$ (S$_1$), which are reduced to 1548 cm$^{-1}$ and 1612 cm$^{-1}$, respectively, by applying the scaling factor of 0.91. This result is consistent with our CASSCF calculations with Sapporo-DZP basis sets, i.e., 1694 cm$^{-1}$ (S$_0$) and 1770 cm$^{-1}$ (S$_1$).

The increase in the NN stretching frequency due to n\pi excitation can be understood by considering the mixing of normal modes. As described above, n\pi excitation strengthens both CN bonds, leading to an increase of force constants in the CN symmetric and anti-symmetric stretching modes. Then, an interaction between the CN symmetric stretching mode and the NN stretching mode of total symmetry representation results in the mixing of these two normal modes. Figure 3 shows the mechanism of the mixing of the NN stretching mode and the CN symmetric stretching mode due to n\pi excitation. As shown in the figure, the NN stretching mode in the ground state shows almost a pure NN stretching motion while in the S$_1(n\pi^*)$ state, the corresponding mode contains a small contribution from a CN symmetric stretching motion; similarly, the original CN symmetric stretching mode contains a small contribution from NN stretching motion with anti-phase. As the result of this mixing, the frequency of the NN stretching mode increases from 1520 to 1581 cm$^{-1}$, while the frequency of the CN symmetric stretching mode decreases from 1235 to 1207 cm$^{-1}$.

In order to examine the effect of CN symmetric stretching mode on the NN stretching frequency, we also performed geometry optimization and normal mode analyses for trans-diazene, N$_2$H$_2$, in the S$_0$ and S$_1(n\pi^*)$ states under the restriction of C$_2h$ symmetry, by the CASPT2 method with Sapporo-DZP basis sets. The calculations show that, in N$_2$H$_2$, the $\pi^*$ anti-bonding orbital is completely localized on the NN bond while the n orbital looks very similar to the one for trans-azobenzene in Fig. 2(a). Table III summarizes the calculated results. According to the present calculations, $\gamma_{NN}$, $\gamma_{NH}$, and $\gamma_{NNH}$ change as 1.265 → 1.290 Å, 1.045 → 1.026 Å, and 105.0 → 118.9$^\circ$, respectively, and the NN stretching frequency changes as 1541 → 1478 cm$^{-1}$, due to n\pi excitation, indicating that NN is weakened in the S$_1(n\pi^*)$ state. This result is consistent with chemical intuition, and therefore the origin of the increase in the NN stretching frequency of trans-azobenzene in the S$_1(n\pi^*)$ state can be ascribed to the mixing of the NN stretching mode and the CN symmetric stretching mode.

| S$_0$  | 1.265 | 1.045 | 105.0 | 1541 | 1308 | 3.38 | ... | 3.00 |
| S$_1$ (C$_2$h)$_{min}$ | 1.290 | 1.026 | 118.9 | 1478 | 621 i | ... | 3.00 |

FIG. 3. A scheme of the mixing between the NN stretching mode and the CN symmetric stretching mode due to n\pi excitation.
As mentioned above, CASPT2 calculations show that the C$_{2h}$ minimum of trans-azobenzene in the S$_1$($\pi\pi^*$) state has one imaginary frequency mode of A$_g$ symmetry, which is directly related to a reaction coordinate of the rotation pathway, i.e., a torsion angle $d$CNNC. To examine the feature of the reaction pathways from (S$_1$-C$_{2h}$)$_{min}$, we calculated the rotation pathway by choosing $d$CNNC as a reaction coordinate and optimizing the other geometrical parameters by the CASPT2 method. Along the rotation pathway, vibrational frequencies were also calculated at the CASPT2 level with a projection technique, which gives a set of normal modes orthogonal to those natural orbitals in the active space determined at $d$CNNC as a reaction coordinate and optimizing the other geometrical parameters by the CASPT2 method. Figure 4 shows variations of (a) S$_0$ and S$_1$ energies, (b) important geometrical parameters, and (c) the NN stretching frequency along the rotation pathway. Cartesian coordinates of geometries along the rotation pathway, as well as a variation of S$_1$ energies plus zero-point vibrational energies of transverse modes, are given in the supplementary material. As shown in Fig. 4(a), the S$_1$ energy shows a gradual decrease until reaching a conical intersection of (S$_1$/S$_0$)$_{CI}$ around $d$CNNC $\sim$ 95°. This result is consistent with a downhill feature of the rotation pathway derived from the semiempirical OM2/MRCI surface hopping simulations recently reported by Thiel et al. It is very interesting to note that both $r_{NN}$ and $r_{CN}$ are almost unchanged along the rotation pathway as shown in Fig. 4(b). This result indicates that the NN bond order does not change while proceeding along the rotation pathway. Figure 4(c) shows that the NN stretching frequency shows only a small decrease ($\sim$ 50 cm$^{-1}$) as $d$CNNC changes from 180° to 100°; this small decrease is consistent with an almost constant bond length of the central NN bond.

From chemical intuition, the rotational motion about the central NN bond should weaken the NN bond because $\pi$-bonding orbital possibly breaks down. The calculated NN bond lengths and NN stretching frequencies, however, indicate that the NN bond is not weakened along the rotation pathway. To get insights to this mechanism, we examined variations of natural orbitals in the active space of SA-CASSCF(6,4) wavefunction. Figures 5(a)–5(c) show those natural orbitals in the active space determined at $d$CNNC $= 180°$, 140°, and 100°, respectively. As shown here, each orbital almost keeps its shape (i.e., n, $\pi$, n, $\pi^*$) while the molecule proceeds along the rotation pathway. The occupation numbers in these orbitals are also constant, (n)$^2$($\pi$)$^2$($n^*$)$^1$ in the main configuration for different values of $d$CNNC. No variation in the orbital shapes during the course of rotation is against chemical intuition, and this is the origin of the unchanged bond order of the central NN bond during the rotational motion.

We also calculated the inversion pathway by choosing a bond angle $\alpha$NNC as a reaction coordinate where two carbon atoms of CNNC part are distinguished by C$_1$ and C$_2$, and optimizing other geometrical parameters by the CASPT2 method. Vibrational frequencies were also calculated along the inversion pathway at the same computational level. Figure 6 shows variations of (a) S$_0$ and S$_1$ energies, (b) important geometrical parameters, and (c) the NN stretching frequency along the inversion pathway; natural orbitals in the active space determined at $\alpha$NNC $= 180°$ are also shown in Figure 5(d). Cartesian coordinates of geometries along the inversion pathway, as well as a variation of S$_1$ energies plus zero-point vibrational energies of transverse modes, are given in the supplementary material.

FIG. 4. Variations of (a) S$_0$ and S$_1$ energies, (b) important geometrical parameters, and (c) the NN stretching frequency along the rotation pathway.

FIG. 5. A set of natural orbitals in the active space of SA-CASSCF(6,4) wavefunction at (a) $d$CNNC $= 180°$ ((S$_1$-C$_{2h}$)$_{min}$), (b) $d$CNNC $= 140°$, and (c) $d$CNNC $= 100°$, along the rotation pathway. The corresponding orbital set at $\alpha$NNC $= 180°$ on the inversion pathway is also given in (d).
material. Along the inversion pathway, the $S_1$ energy shows an increase by ca. 0.8 eV at $\alpha_{\text{NNC}} \sim 180^\circ$, and simultaneously the NN bond length decreases by 0.03 Å, and the NN stretching frequency increases by ca. 300 cm$^{-1}$. The increase of the NN bond order at $\alpha_{\text{NNC}} \sim 180^\circ$ can be understood by noting that one of n orbitals gets $\pi$-bonding character as shown in Fig. 5(d). The increase in the adiabatic energy along the inversion pathway clearly indicates that trans-azobenzene prefers the rotation pathway, compared to the inversion pathway, in $\pi\pi^*$ excitation. This finding can explain, why in dynamics simulations the isomerization occurs through a rotation of the $\sim$N$\rightleftharpoons$N$\leftleftharpoons$ fragment rather than a true torsion along the N-N bond.

Finally, we summarize significant results from the present calculations. First, we should mention that we employ the CASPT2 method in determinations of geometrical structures, vibrational frequencies, and reaction pathways. Several studies employed $ab\ initio$ multireference theory in the evaluations of the energy for geometries determined by the CASSCF method, but to our best of knowledge, there has been no study to determine geometrical structures of stationary points and reaction pathways at the multireference correlated level. The energy variations along the reaction pathways clearly indicate that the rotation pathway is preferred to the inversion pathway. We also calculated the NN stretching frequencies for $(S_0)_{\text{min}}, (S_1-C_2h)_{\text{min}}$, and selected structures along the reaction pathways at the CASPT2 level, and compared those frequencies with the experimental values. A shift of the NN stretching frequency due to $\pi\pi^*$ excitation is relatively small, agreeing to the frequency shows a decrease in the experiment while it shows an increase in the calculations.

**IV. CONCLUDING REMARKS**

In this work, we have examined the NN stretching frequency of trans-azobenzene in both $S_0$ and $S_1(\pi\pi^*)$ states thoroughly by $ab\ initio$ multireference perturbation theory, CASPT2, to gain insights into the photosomerization mechanism of trans-azobenzene. Through the time-resolved Raman measurements in solution, Tahara et al. found that the NN stretching frequency shows only a slight decrease in the $S_1(\pi\pi^*)$ state. Our CASPT2 calculations show that, due to $\pi\pi^*$ excitation, the NN bond length changes little ($\sim$0.004 Å) and the NN stretching frequency increases only by 64 cm$^{-1}$ at the $C_{2h}$ minimum on the $S_1$-PES; the former is explained in terms of molecular orbitals related to $\pi\pi^*$ excitation while the latter is explained by the mixing of the NN stretching normal mode with the CN symmetric stretching normal mode.

It is also shown that the $C_{2h}$ minimum of trans-azobenzene in the $S_1(\pi\pi^*)$ state is unstable with respect to the CNNC torsion angle, although an absolute value of the imaginary frequency of the related mode is not large. We also calculated the rotation and inversion pathways as a function of $\alpha_{\text{CNNC}}$ and $\alpha_{\text{NNC}}$, respectively, at the CASPT2 level, and then calculated the NN stretching frequency along the respective reaction pathways by a projection technique. The calculated NN stretching frequency shows a slight decrease along the rotation pathway while it increases rapidly along the inversion pathway. The almost constant bond order of the central NN bond along the rotation pathway is explained in terms of little changes of orbital shapes. A small variation of the NN stretching frequency along the rotation pathway suggests the possibility that in the experimental Raman spectra, vibrational motion of the molecule under structural transformation along the rotation pathway is observed, and our CASPT2 calculations support the rotation pathway as a preferred one in the $\pi\pi^*$ excitation of trans-azobenzene.

Finally, it should be noted that our calculations support the rotation mechanism in the gas phase, and there remains a possibility of other reaction mechanism in solution, depending on the type of solvent molecules.

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