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Ab initio molecular dynamics (AIMD) simulations for the excited-state hydrogen transfer (ESHT) reaction of 7-azaindole (7AI−(H₂O)ₙ; n = 1, 2) clusters in the gas phase and in water are presented. The effective fragment potential (EFP) is employed to model the surrounding water molecules. The AIMD simulations for 7AI−H₂O and 7AI−(H₂O)₂ clusters show an asynchronous hydrogen transfer at t ∼ 50 fs after the photoexcitation. While the ESHT mechanism for 7AI−H₂O in water does not change appreciably compared with that in the gas phase, the AIMD simulations on 7AI−(H₂O)₂ in water solution exhibit two different mechanisms. Since the tautomer form is lower in energy compared to the normal form in the S₁ state, 7AI and (H₂O)ₙ fragments separate from each other after the ESHT. With the use of the results of the AIMD trajectories, the minimum energy conical intersection point in the tautomer region has also been located.

I. Introduction

Excited-state hydrogen transfer (ESHT) is a fundamental reaction which plays an important role in a variety of biological processes.1–3 Typically, this type of transfer occurs through a hydrogen-bonded network. 7-Azaindole (7AI) is a molecule in which this process is important. Tautomerization of 7AI, accompanying the ESHT from the five-membered ring to the six-membered ring, has been studied with much attention both experimentally4–21 and theoretically,22–30 since 7AI can be regarded as a simple model for DNA base pairs. Very recently, Sakota and co-workers measured the dispersed fluorescence (DF) spectra and resonance-enhanced multiphoton ionization (REMPI) spectra for the 7AI−(H₂O)₂ (n = 2, 3) cluster at low temperature in the gas phase and found that the tautomerization of 7AI−(H₂O)₂ (n = 2, 3) occurs in the S₁ state,21 although it was not observed at first based on the measurement of the fluorescence excitation (FE) and DF spectra.16 It was also reported18,19 by the same group that the excited-state triple-H-transfer occurs in the 7AI−(MeOH)₂ cluster, based on the measurement of DF and REMPI spectra, but the tautomerization does not occur in 7AI−MeOH and 7AI−(MeOH)₃ clusters. They pointed out that ESHT in 7AI−(MeOH)₂ proceeds via a tunneling mechanism in their experimental conditions, and the excitation of the low-energy intermolecular stretching mode accelerates ESHT in 7AI−(MeOH)₂.

An early theoretical effort on the ESHT reaction of 7AI−H₂O was performed by Chaban and Gordon,22,24 in which the intrinsic reaction coordinate was calculated for the tautomerization in an isolated 7AI molecule and the 7AI−H₂O complex in the singlet ground (S₀) and first excited (S₁) states at the CASSCF (complete active space self-consistent field) level of theory, with improved energetics obtained using multireference second-order perturbation theory (MCQDPT2).31 It was shown that the normal form of 7AI is more stable than the tautomer in the S₀ state, whereas the relative energies are reversed in the S₁ state. The activation energy for tautomerization in 7AI is significantly reduced by the complexation with water, because then the H transfer occurs via the 7AI−H₂O hydrogen-bonded network. Casadesús et al.27 examined the reaction profile for the tautomerization of 7AI−(H₂O)ₙ (n = 0–4) in the S₁ state using single-excitation configuration interaction (CIS) geometry optimization, followed by time-dependent density functional theory (TDDFT) energy calculations, and discussed the dependence of the activation barriers on the number of attached water molecules. They found that the ESHT mechanism in 7AI−(H₂O)₂ is the most efficient one among 7AI−(H₂O)ₙ (n = 0–4) clusters.

In addition to the experimental observation in the gas phase, the tautomerization in 7AI was also observed in the condensed phase in alcohol or water solution,4–10,11 where the tautomerization exhibits a strong dependence on the water concentration in mixtures of water and aprotic solvents.8 It is possible that ESHT would exhibit different mechanisms for the reactions in the gas phase and in the condensed phase. The spectroscopic experiments in the gas phase have been carried out at very low temperature, and so quantum mechanical tunneling could contribute significantly to the ESHT reaction. On the other hand, the experiments in solution have been carried out at room temperature, and ESHT reactions could occur classically (without tunneling), because there is sufficient energy in the system to overcome the small activation barrier. Fernández-Ramos et al.26 calculated the rate constant for the ESHT reaction of 7AI in solution using the Onsager model. Although the rate constants calculated for 7AI−H₂O and 7AI−(H₂O)₂ embedded in a dielectric continuum did not reproduce the experimental ones in dilute solution, those calculated for the 7AI−(H₂O)₃ complex were shown to reproduce the observed order of magnitude.
The Onsager model has limited accuracy for predicting solvent effects and cannot account for explicit solute—solvent interactions, so a more sophisticated approach would be useful for improving the understanding of the effects of solvation structures on the dynamics. One alternative approach is the effective fragment potential (EFP) method, implemented in the program package GAMESS, which in which each solvent molecule is represented by a discrete model potential. When combined with an electronic structure calculation for the solute, this method can be classified as a quantum mechanics combined with molecular mechanics (QM/MM) approach. The solvent molecules that participate directly in the reaction process should be included in the QM part, while the environmental solvent molecules can be represented by EFPs. In this work, we present ab initio QM/MM molecular dynamics (AIMD) simulations for the excited-state full tautomerization process in 7AI \((\text{H}_2\text{O})_n\) \((n = 1, 2)\) clusters in a water solvent environment and investigate the role of solvent water molecules on the ESHT reaction. The AIMD method is based on a classical trajectory, in which the force acting on each atom is calculated “on-the-fly” by ab initio calculations, and it has been extended to excited-state dynamics very recently, including a nonadiabatic surface-hopping scheme.

II. Computational Details

Geometry optimizations and normal-mode analyses were carried out for 7AI and 7AI \((\text{H}_2\text{O})_n\) \((n = 1, 2)\) in the ground and first excited singlet states by the state-specific CASSCF method with the segmented DZP basis set, using the quantum chemical program package, GAMESS. The CASSCF active space includes 10 electrons in 9 orbitals which comprise all of the 7AI \(\pi\) orbitals and electrons. The transition state geometries for tautomerization in 7AI \(\sim \text{H}_2\text{O}\) and 7AI \(\sim \text{H}(\text{H}_2\text{O})_2\) were determined by the same methods. Following the examination of the energetics of the reactions, AIMD simulations were performed for the ESHT processes in 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\) \((n = 1, 2)\). The AIMD code used here is based on the leapfrog algorithm and utilizes ab initio energy gradients. The initial conditions for the dynamics simulations were the equilibrium structures in the ground state, with the atomic velocities given in the directions of the transition state. The latter consists of two normal modes, shown in Figure 1 (plotted with MacMolPlt). Preliminary AIMD simulations were performed in which the vibrational quantum numbers were successively incremented by one. It was determined that at least a vibrational energy equivalent to three photons is required to invoke H-transfer reactions within a tractable amount of time. Therefore, the kinetic energy given to the respective modes in the production simulations corresponds to three photons. These initial conditions were employed to promote the H transfer efficiently.

To simulate the ESHT processes in water, 100 Hartree–Fock (HF)-based effective fragment potential (EFP/HF) water molecules were distributed around 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\). The EFP/HF method was developed to describe water clusters, either with or without an explicit ab initio solute system. The EFP method accounts for three important interaction terms: (a) Coulomb interactions for solute–solvent and solvent–solvent, (b) solute–solvent and solvent–solvent polarization interactions, and (c) exchange repulsion + charge transfer interactions. In the presence of an ab initio solute, these terms are added as one-electron terms to the ab initio Hamiltonian. The Coulomb interaction term is approximated with a distributed multipolar analysis (DMA) of the solvent molecule, expanded through octopoles. The polarization term is treated by a self-consistent distributed perturbation model with localized molecular orbital (LMO) polarization tensors. In order to obtain the remaining (exchange repulsion + charge transfer) interaction, the first two terms are calculated for many points on the water dimer potential energy surface and subtracted from the total Hartree–Fock interaction potential. This remainder (exchange repulsion + charge transfer) interaction is then fitted to a functional form. Note that the internal geometry of an EFP water is fixed.

The AIMD simulations for an ab initio solute in the excited state surrounded by EFP solvent molecules require analytic energy gradients for both the ab initio and EFP parts. In the current version of GAMESS, such excited-state simulations can only be performed with state-specific CASSCF. The electron density used to determine induced dipoles in EFP is obtained from the orbitals and CI coefficients of the selected CASSCF state. That is, the density in the MO basis is formed from the CI coefficients of the chosen state and then transformed to the AO basis via the MO coefficients; the resulting AO density matrix is then used for the polarization step (this is performed in each CASSCF iteration until self-consistency is attained).

The RHF/STO-3G-AIMD simulations for 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\)–EFP were performed in the ground state for 10 ps at a constant temperature of 300 K with a time step of 1 fs. In this process the geometry of the solute 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\) was fixed. Twenty different configurations and velocities of EFPs were chosen from the above AIMD trajectories and used as the initial conditions for the AIMD simulations in the excited state. The AIMD simulations for phototautomization in 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\) and 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\)–EFP \((n = 1, 2)\) were performed at the CASSCF level of theory with the segmented DZP basis set, and the solvent effects on the ESHT processes were investigated from the dynamical point of view. The time step was taken as 0.5 fs throughout the simulation.

The minimum energy conical intersection point for the S0 and S1 states was also located with the state-averaged CASSCF (SA-CASSCF) method in the Molpro program. The role of this conical intersection in the ESHT process is discussed, based on the results of the AIMD simulations. To estimate more accurate energetics for the reactions, CASPT2 calculations were also carried out at the S0 and S1 states.

III. Results and Discussion

III.A. Energetics and Dynamics of Tautomerization in the 7AI \(\sim \text{H}(\text{H}_2\text{O})_n\) Cluster. Figure 2 shows the calculated equilibrium geometries for the normal and tautomer forms of 7AI–H2O and
the transition state (TS) geometries for tautomerization in both the $S_0$ and $S_1$ states, where the CASSCF interatomic distances and relative energies are given. In the transition states, double-H-transfer occurs in a concerted manner in both the $S_0$ and $S_1$ states. Each transition state has only one imaginary frequency. When the $S_0$ and $S_1$ equilibrium structures are compared, it is observed that most CC and CN bond distances increase upon photoexcitation. This feature is related to the character of the CASSCF wave function in the $S_1$ state, in which several electronic configurations with excitations from $\pi$ to $\pi^*$ orbitals contribute to the $S_1$ wave function. As a result, vibrational excitation of the 7AI ring-skeleton may be induced by the photoexcitation. The normal form is lower in energy by ca. 13 kcal/mol than the tautomer in the $S_0$ state, whereas the tautomer is lower in energy by ca. 32 kcal/mol than the normal form in the $S_1$ state. The activation barriers relative to the normal isomer are 39.7 and 16.9 kcal/mol, respectively, in the $S_0$ and $S_1$ states (the vibrational zero-point corrected barriers are 36.0 and 13.5 kcal/mol), so the H-transfer reaction occurs more easily in the $S_1$ state. The geometries and relative energies are in good agreement with those reported by Chaban and Gordon; the differences in the interatomic distances and relative energies are $\sim 0.025 \text{ Å}$ and 1.3 kcal/mol, respectively, except for the TS geometry in the $S_0$ state. The difference in this TS geometry is caused by the use of different basis sets. The energy profiles for the tautomerization in 7AI$^-$($H_2O)_2$ are almost the same as those in 7AI$^-$(H$_2$O)$_n$. As will be discussed later, it is found that inclusion of dynamic electron correlation by CASPT2 lowers the activation energy in both the $S_0$ and $S_1$ states (see Table 1).

The AIMD trajectory simulations were started on the $S_1$ excited state from the ground-state equilibrium structure for the normal form of 7AI$^-$(H$_2$O)$_n$. The initial velocities are given in the directions of the H transfer consisting of the two normal modes shown in Figure 1, whose frequencies are 3828 and 4013 cm$^{-1}$ for 7AI$^-$(H$_2$O) and 3729 and 3920 cm$^{-1}$ for 7AI$^-$(H$_2$O)$_2$, respectively. The kinetic energy is given to the respective modes with three quanta, and this amounts to 67.3 and 65.6 kcal/mol for 7AI$^-$(H$_2$O) and 7AI$^-$(H$_2$O)$_2$, respectively. Figure 3 displays the time evolution of (a) the relative energy between the $S_0$ and $S_1$ states and (b) the interatomic distances related to the H transfer along the AIMD trajectory for 7AI$^-$(H$_2$O). During the initial 40 fs, the molecule stays in the normal-form region of the potential energy surface, with fluctuations in the excitation energy with a period of $\sim 10$ fs (Figure 3a); this is consistent with the approximate period of vibration of the N···H and O···H hydrogen-bond stretching modes as shown in Figure 3b. The excitation energy in Figure 3a decreases rapidly at $\sim 50$ fs, and the hydrogen-bond distances, N···H and O···H, decrease to covalent bond lengths, indicating that ESHT occurs through the hydrogen-bonded network. The N···H and O···H hydrogen-
the NH on the five-membered ring to OH, completing the tautomerization of 7AI. After the H transfer, the molecular system acquires considerable energy due to the stabilization of the tautomer in the S1 state, and the 7AI and H2O fragments separate, breaking the two hydrogen bonds. This dynamical behavior indicates that the ESHT reaction in 7AI–H2O proceeds in a concerted manner via the TS structure shown in Figure 2, but rather asynchronously.

It is interesting to note the recent experimental study of Takeuchi and Tahara50 for the double-proton-transfer in the 7AI dimer in solution, in which the 7AI dimer converts to the tautomer configuration in the excited state. They investigated this process by excitation wavelength dependence in steady-state and femtosecond time-resolved fluorescence spectroscopy and concluded that it proceeds in a concerted manner. They noted, however, that this conclusion does not necessarily translate to a synchronous motion of the two protons. A concerted mechanism does not imply strict simultaneity, only that the motions of the two protons are correlated. Ushiyama and Takatsuka51 performed AIMD simulations for the double-H-transfer in formic acid dimer in the ground state. The minimum energy path for this process indicates that the synchronous H transfer is more favorable than the asynchronous one, but AIMD simulations showed that there is a certain time lag (~8 fs) between the two H transfers. This type of motion is an example of a concerted double-H-transfer, which is fundamentally different from the stepwise process.

Figure 4 shows the time evolution of (a) the relative energy of the S0 and S1 states and (b) the interatomic distances related to the H transfer in the AIMD simulation for 7AI–H2O. Solid and dashed lines denote changes in the N–H and O–H distances, respectively.

Changes in the interatomic distances (Figure 4b) indicate that the triple-H-transfer relays occur in the time range of 40 to 60 fs, accompanying the gradual decrease in the excitation energy as shown in Figure 4a. A detailed picture of this process is summarized in the following three steps, i.e., (1) H2 moves from N1 to O3, (2) H3 moves from O3 to O5, and (3) H6 moves from O5 to N7 (the tautomerization completes). In the first step, the sum of the net atomic charges on the H2O moiety is +0.58 and +0.50 according to the Mulliken and ESP population analyses, respectively. The reaction mechanism is a concerted asynchronous process. As illustrated for the 7AI–H2O MD simulation, 7AI and H2O fragments separate after the ESHT reaction.

As is shown in Figure 4c, the adiabatic energies of the S0 and S1 states approach each other in the tautomeric region (indicated by arrows in the figure), suggesting the possibility of a nonradiative decay through a conical intersection of the potential energy surfaces. In order to examine this possibility, an SA-CASSCF geometry optimization was performed to determine the minimum energy conical intersection (MECI) point between the S0 and S1 states for an isolated 7AI molecule, starting from the structure with the smallest relative energy on the 7AI–(H2O)2 AIMD trajectory. Note that the AIMD simulations were carried out using the state-specific CASSCF method. The energetics obtained using the state-specific and state-averaged CASSCF calculations are slightly different from each other. Figure 5a shows two geometrical structures. On the left is the starting structure in the search for the MECI. On the right is the MECI structure, in which the participating H atom attached to the six-membered ring is out of the molecular plane.

**Table 1: Relative Energies (kcal/mol) for Points Related to the H-Transfer Reactions in the S0 and S1 States for 7AI, 7AI–H2O, and 7AI–(H2O)2 Calculated with SA-CASSCF and CASPT2**

<table>
<thead>
<tr>
<th></th>
<th>normal S0-min</th>
<th>S1-min</th>
<th>TS</th>
<th>tautomer S0-min</th>
<th>S1-min</th>
<th>MECI*</th>
</tr>
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<tbody>
<tr>
<td>7AI SA-CASSCF</td>
<td>0.0</td>
<td>4.8</td>
<td>85.8</td>
<td>16.5</td>
<td>33.1</td>
<td></td>
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<tr>
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<td>75.3</td>
<td>88.0</td>
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<tr>
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<td>1.3</td>
<td>61.1</td>
<td>13.0</td>
<td>22.3</td>
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<tr>
<td>7AI CASPT2</td>
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<td>98.0</td>
<td>138.6</td>
<td>85.3</td>
<td>71.9</td>
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<tr>
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<tr>
<td>7AI–H2O CASPT2</td>
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<td>105.8</td>
<td>127.0</td>
<td>92.4</td>
<td>76.3</td>
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<td>40.9</td>
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<tr>
<td>7AI–(H2O)2 CASPT2</td>
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<td>105.6</td>
<td>126.7</td>
<td>92.6</td>
<td>77.0</td>
<td></td>
</tr>
</tbody>
</table>

* Minimum energy conical intersection point.

**Figure 3.** Time evolution of (a) the relative energy of S0 and S1 states and (b) the interatomic distances related to the H transfer in the AIMD simulation for 7AI–H2O. Solid and dashed lines denote changes in the N–H and O–H distances, respectively.
Excited-State Hydrogen Transfer of 7-Azaindole


The SA-CASSCF energies for the $S_1$ state were calculated at linearly interpolated points between these two structures and verified that no obstructive barrier exists along the interpolated path. Therefore, when an H is transferred from the out-of-plane side to the six-membered ring N atom, 7AI could easily reach this conical intersection point, suggesting a nonradiative decay process after the phototautomerization in an isolated 7AI molecule is much higher than that in 7AI-$\text{H}_2\text{O}$ since, without a bridge of water molecules, an H needs to be directly transferred from the five-membered ring N atom to the six-membered ring N atom.\(^{24}\) As indicated already, the energy profiles for the tautomerization in 7AI-$\text{H}_2\text{O}_2$ are almost the same as those in 7AI-$\text{H}_2\text{O}$ as shown in Table 1. Also, the activation barrier for tautomerization in an isolated 7AI molecule is much higher than that in 7AI-$\text{H}_2\text{O}$ since, without a bridge of water molecules, an H needs to be directly transferred from the five-membered ring N atom to the six-membered ring N atom.\(^{24}\) Note (Table 1) that the addition of dynamic correlation via CASPT2 significantly reduces the activation barrier for the reaction in both the $S_0$ and $S_1$ states. In particular, the CASPT2 activation barrier for 7AI-$\text{H}_2\text{O}$ in the $S_1$ state is predicted to be only 2.5 kcal/mol at the SA-CASSCF geometries. This trend was previously pointed out by Chaban and Gordon.\(^{24}\) This small activation barrier indicates that at room or higher temperatures, the H-transfer reaction does not require quantum tunneling to proceed.

### III.B. Solvent Effects on the ESHT Dynamics for the 7AI-$\text{H}_2\text{O}_n$-EFP System

Now, consider aqueous solvation effects on the ESHT dynamics of 7AI-$\text{H}_2\text{O}$ and 7AI-$\text{H}_2\text{O}_2$, using the EFP method. The accuracy of the EFP method was evaluated by comparing the hydrogen-bond distance and the binding energy of an EFP water to the 7AI-$\text{H}_2\text{O}_n$ clusters with those obtained by fully ab initio calculations. CASSCF calculations were performed for the 7AI-$\text{H}_2\text{O}_n$ ($n = 1, 2$) clusters with one solvent water molecule (using an EFP or an ab initio water) shown in Figure 6. The EFP hydrogen-bond distances are $r_1 = 2.037$ Å, $r_2 = 2.191$ Å, and $r_3 = 2.545$ Å in the $S_0$ state, whereas $r_1 = 2.036$ Å, $r_2 = 2.192$ Å, and $r_3 = 2.555$ Å in the $S_1$ state. The corresponding ab initio bond distances are $r_1 = 2.073$ Å, $r_2 = 2.154$ Å, and $r_3 = 2.476$ Å in the $S_0$ state and $r_1 = 2.072$ Å, $r_2 = 2.152$ Å, and $r_3 = 2.489$ Å in the $S_1$ state. The binding energy comparison was performed for the EFP-optimized structures. For the 7AI-$\text{H}_2\text{O}_2$ complex, the binding energy is 4.7 (EFP) and 5.0 kcal/mol (ab initio) in the $S_0$ state and 4.7 (EFP) and 5.0 kcal/mol (ab initio) in $S_1$. For the 7AI-$\text{H}_2\text{O}_2$ complex, the binding energy is 6.0 (EFP) and 6.7 kcal/mol (ab initio) in the $S_0$ state and 6.0 (EFP) and 6.80 kcal/mol (ab initio) in $S_1$. Note that the ab initio binding energies may suffer from the basis set superposition error, whereas this is not relevant for the EFP model potential.

The agreement between the ab initio and EFP results are reasonable, indicating that the EFP approach is suitable for the present system.

To determine the initial positions and velocities of 100 EFP waters for the examination of excited-state dynamics, preliminary RHF AIMP simulations were carried out for the 7AI-$\text{H}_2\text{O}_n$-EFP ($n = 1, 2$) ground state. In this simulation, velocity scaling was applied for a time interval of 0.25 ps, to control the temperature of EFP waters at 300 K. The temperature shows a small fluctuation around 300 K for $t > 1.0$ ps. Twenty points from the trajectory were chosen as the initial conditions for the excited-state AIMD simulations. During the simulations, each EFP water maintains a hydrogen bond with other EFP or

**Figure 4.** Time evolution of (a) the relative energy of $S_0$ and $S_1$ states and (b) the interatomic distances related to the H transfer in the AIMD simulation for 7AI-$\text{H}_2\text{O}_2$ where solid, dashed, and dotted lines denote changes of the $\text{H}^\cdots\cdot\text{O}^1$, $\text{H}^\cdots\cdot\text{O}^2$, and $\text{H}^\cdots\cdot\text{N}$ distances, respectively. A longer time evolution of the relative energy of $S_0$ and $S_1$ states for 1000 fs is given in (c).

The SA-CASSCF energies for the $S_1$ state were calculated at the energetics of the reactions for 7AI-$\text{H}_2\text{O}_2$ and $\text{H}_2\text{O}$ surfaces, however, geometries are determined at the respective SA-CASSCF and CASPT2 computational levels. The geometry at the MECI point on the CASPT2 potential energy surface was determined by utilizing the SA-CASSCF analytical energy gradients and the CASPT2 energy difference, assuming that the SA-CASSCF potential energy surface is parallel to the CASPT2 one. Of course the CASPT2 energy gradients are not strictly equal to the SA-CASSCF energy gradients, but the CASPT2 MECI point was located to within 0.2 kcal/mol by several trial calculations. The CASPT2 MECI geometry is shown in Figure 5b, which is very close to the SA-CASSCF MECI geometry.
ab initio waters. When one hydrogen bond breaks, a new hydrogen bond forms.

In the AIMD simulations for 7Al–H₂O–EFP, SA-CASSCF does not converge in one trajectory around \( t \sim 45 \) fs, so this trajectory was eliminated from the subsequent analyses. Among the remaining 19 trajectories examined for 7Al–H₂O–EFP, the ESHT reaction occurs in 15 trajectories, while no reaction occurs in 3 trajectories. In one trajectory, the ESHT reaction occurs once, but promptly the system returns to the normal-form region. This is an example of transition state recrossing. Figure 7 shows the time evolution of (a) the relative energy of S₀ and S₁ states and (b) the interatomic distances related to the H transfer in the AIMD simulation for 7Al–H₂O–EFP where solid and dashed lines denote changes of the N···H and O···H hydrogen-bond lengths, respectively.

AIMD simulations were also carried out for 7Al–(H₂O)₂–EFP starting from 20 different initial conditions. The triple-H-transfer occurs in 14 trajectories, while no reaction occurs in 5 trajectories. Recrossing behavior is seen in one trajectory. Unlike 7Al–H₂O–EFP, the surrounding solvent waters affect the ESHT dynamics more significantly, resulting in different H-transfer mechanisms, depending on the initial solvent configurations. Table 2 summarizes the results of AIMD simulations for 7Al–H₂O–EFP and 7Al–(H₂O)₂–EFP. In 12 trajectories, the ESHT dynamics proceeds asynchronously via an H₂O–OH⁻–7Al–H⁺ species in nearly half of the trajectories, while the other half of trajectories exhibit an
TABLE 2: Branching Numbers of Reaction Patterns for AIMD Simulations on 7AI–H2O–EFP and 7AI–(H2O)2–EFP Systems

<table>
<thead>
<tr>
<th>Reaction Pattern</th>
<th>Asynchronous</th>
<th>Recrossing</th>
<th>No Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>7AI–H2O</td>
<td>0</td>
<td>15</td>
<td>0</td>
</tr>
<tr>
<td>7AI–(H2O)2</td>
<td>2</td>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

H2O–H⁺⁺–H₂O⋯7AI⁰⁻ species. In two trajectories, an almost-synchronous H transfer is observed. Figure 8 shows the time evolution of (a) the relative energy between the S₀ and S₁ states and (b) the interatomic distances along one of 14 reactive trajectories for 7AI–(H₂O)₂–EFP. The changes in the energy and the interatomic distances are similar to those in the 7AI–(H₂O)₂ trajectory (Figure 4, parts a and b) in some respects: (1) the relative energy shows fluctuations with a period of 11 fs in the initial stage; (2) the interatomic distances show complicated changes in the time period t = 40 to 60 fs with a gradual decrease of the relative energy. The fluctuations in the relative S₀ – S₁ energy with an approximately 20 fs cycle, observed later in the trajectory, may be caused by the vibrational motions of the 7AI ring deformation. As for 7AI–H₂O–EFP, the water molecules departing from 7AI are prevented from moving further by the surrounding EFP solvent waters.

If the transition state has some charge transfer character, polar solvents such as water should lower the activation barrier, and increase the reaction rate. The present simulations show that ESHT reactions of 7AI–H₂O and 7AI–(H₂O)₂ proceed via polar configurations caused by the asynchronous H transfer. A continuum model for solvation assumes that the reorientation of solvent molecules is very fast compared to the motion of solute molecule, whereas the EFP approach takes into account the reorientation process of solvent molecules explicitly. It is possible that the EFP waters do not have sufficient time to relax during the ESHT processes.

It is important to mention the limitations of the present simulation. For example, this work reports classical dynamics simulations that do not include nuclear quantum effects, such as tunneling, which could play an important role in the ESHT process. Experimentally Sakata and co-workers⁸,¹⁰ suggested that quantum tunneling is a dominant mechanism for ESHT in a 7AI–(solvent) cluster in the gas phase at low temperature. One needs a quantum dynamical approach to include such quantum effects in the excited-state dynamics simulation.⁵² In solution at standard thermodynamic conditions, however, the system consists of a large number of solute and solvent molecules and is expected to have sufficient energy to surpass the activation barrier upon photoexcitation. Although the present results are obtained from classical dynamics simulations, they nonetheless provide useful information related to ESHT reaction dynamics.

The foregoing 7AI–(H₂O)₂–EFP AIMD simulations were initiated from configurations in which one or two water molecules bridge N (six-membered ring) and H (five-membered ring) atoms of 7AI to assist the tautomerization. It is interesting to examine the number of waters that enter the 7AI hydrogen-bonded region under thermodynamic equilibrium conditions. Additional AIMD simulations were performed for 7AI + 101 EFP waters (trajectory 1) and 7AI + 102 EFP waters (trajectory 2) in the ground state using RHF/STO-3G at 300 K with a time step of 1 fs over 60 ps (60 000 steps) and with a fixed internal 7AI geometry. The initial geometries and velocities were taken from those used for the previously discussed AIMD simulations on 7AI–H₂O–EFP (trajectory 1) and 7AI–(H₂O)₂–EFP (trajectory 2). Ab initio waters were replaced by EFP waters for these simulations. Along the trajectories, the number of configurations was counted each time an EFP water O atom entered a sphere whose center was located at the equilibrium position of the O atom of isolated 7AI–H₂O and 7AI–(H₂O)₂ clusters. Figure 9 illustrates the cumulative average of the probability that an O atom is inside the sphere of single-water-bridged configuration of 7AI–H₂O (case 1) and that two oxygen atoms are within the spheres of double-water-bridged configuration of 7AI–(H₂O)₂ (case 2). The radius of the spheres is set equal to 0.7 (Figure 9a) and 1.0 Å (Figure 9b). As seen in the figure, the spheres of the 7AI–H₂O and 7AI–(H₂O)₂ systems overlap appreciably; thus, when an O atom enters this overlap region and simultaneously another oxygen atom is found in the other sphere, it is counted as case 2 and not counted as case 1 in order to avoid double-counting of the configuration. One can see that converged results are obtained in case 2 for both trajectories, whereas slow convergence is seen in case 1. The relative ratio of case 2 and case 1 indicates that the single-water-bridged configurations are observed roughly twice as much as the double-water-bridged ones under the equilibrium condition of 300 K.

IV. Concluding Remarks

AIMD simulations have been performed to explore the ESHT dynamics of 7AI–(H₂O)ₙ clusters in the gas phase and in water solution, where in the latter case the surrounding water molecules are represented by the EFP model. The AIMD simulations for the 7AI–H₂O cluster show that a hydrogen is transferred from H₂O to 7AI at t ~ 50 fs after the photoexcitation, and the second H transfer occurs promptly from NH on the five-membered ring (asynchronous concerted mechanism). Since the tautomer form is more stable compared to the normal form in the S₁ state, H₂O and 7AI fragments depart from each
indicating an asynchronous concerted mechanism. The relative H-transfer relay also occurs very quickly in a correlated way, subsequently the second and third H transfers occur. This triple-

Figure 9. Cumulative average of the probability that an oxygen atom is inside the sphere of a single-water-bridged configuration of 7AI—(H2O) (red circle) and that two oxygen atoms are within the spheres of double-water-bridged configuration of 7AI—(H2O)2 (blue circles), where a radius is set equal to (a) 0.7 Å and (b) 1.0 Å. (1,s) represents trajectory 1 and case 1 (single-water-bridged) and (1,d) indicates trajectory 1 and case 2 (double-water-bridged).

other just after the ESHT. For the 7AI—(H2O)n cluster, a hydrogen on the five-membered ring moves to H2O, and subsequently the second and third H transfers occur. This triple-

H-transfer relay also occurs very quickly in a correlated way, indicating an asynchronous concerted mechanism. The relative energy between the $S_0$ and $S_1$ states after the ESHT reaction indicates the existence of a conical intersection in the region of the tautomer form. The MECI point was determined using the SA-CASSCF and CASPT2 methods. The present results suggest the possibility of a nonradiative decay after the ESHT.

To investigate the solvent effects on the ESHT dynamics in 7AI—(H2O)n ($n = 1, 2$), AIMD simulations were performed for 7AI—(H2O)2 plus 100 surrounding EFP water molecules. Twenty different sets of positions and velocities were taken as the initial conditions for solvent molecules from a constant temperature MD simulation at 300 K. ESHT occurs in 15 trajectories for 7AI—(H2O)2—EFP and in 14 trajectories for 7AI—(H2O)2—EFP. In the former case, the ESHT reaction occurred in a similar manner to that found in the 7AI—H2O cluster, in which the H transfer takes place asynchronously via 7AI—H5+···OH9−. In contrast, in the AIMD simulations for 7AI—(H2O)2—EFP, three different mechanisms are observed. The ESHT reaction goes asynchronously through H2O—H5+—H2O···7AI9− configurations in six trajectories, whereas in six other trajectories the process goes through H2O—OH9−···7AI9−; in two trajectories a synchronous H transfer is observed. This result indicates that, in 7AI—(H2O)2—EFP, the ESHT dynamics is more affected by the surrounding solvent waters.

The simulations discussed here were limited to 20 trajectories for the 7AI—(H2O)2—EFP systems due to the expensive computational costs (~100 h for each 7AI—(H2O)2—EFP trajectory on a Xeon 5160 (3.0 GHz) processor). To obtain a dynamical property, such as a reaction probability or the energy distribution of the products from statistical analyses of AIMD simulations, one would need to run many more trajectories, starting from various initial conditions. The electronic transition should be incorporated into the simulation in order to account for the possibility of nonradiative decay. This could be possible in combination with a semiclassical scheme, such as the surface-hopping algorithm.

The present AIMD code has been implemented in a general ab initio program package. GAMESS.34,35 The statistical analyses of the ESHT dynamics including electronic transition from AIMD simulations will become a tractable task in the near future.

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References and Notes

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