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QM/MM MD insight into photodynamics of hypoxanthine: distinct nonadiabatic decay behaviors between keto-N7H and keto-N9H tautomers in aqueous solution†

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†Electronic Supplementary Information (ESI) available: Relative energy profiles along the LIIC reaction path from the S_0 state to the conical intersection between S_0 and S_1 states of keto-N9H. [The 6 optimized geometries for both tautomers from 6 sample snapshots.](#) Distributions of the $\delta(C_2-N_3)$ and $\delta(N_1-C_2)$ bond lengths of keto-N7H and keto-N9H at the $S_1 \rightarrow S_0$ hopping points. Predicted key geometric parameters in the gas phase and in aqueous solution along with the experimental crystal data. Cartesian coordinates (Å) of the QMMM-optimized representative local ground-state geometries. See DOI: 10.1039/c0xx00000x

Abstract: Extensive *ab initio* surface-hopping dynamics simulations have been used to explore the excited-state nonadiabatic decay of two biologically relevant hypoxanthine keto-N7H and keto-N9H tautomers in aqueous solution. QM/MM calculations and QM/MM-based MD simulations predict different hydrogen bonding networks around these nucleobase analogues, which influence their photodynamical properties remarkably. Furthermore, different solvent effects on the conical intersection formation of keto-N7H and keto-N9H were found in excited-state MD simulations, which also change the lifetimes of the excited states. In comparison with the gas-phase situation, the $S_1 \rightarrow S_0$ nonradiative decay of keto-N7H is slightly faster, while this decay process of keto-N9H becomes much slower in water. The presence of π -electron hydrogen bond in the solvated keto-N7H is considered to facilitate the $S_1 \rightarrow S_0$ nonradiative decay process.

1 Introduction

There is an increasing interest in characterizing the photochemistry and photophysics of electronically excited states of nucleobases.¹ As the fundamental building blocks of life, these nucleic acid bases may undergo extremely ultrafast nonradiative decays in vacuum, solution and DNA environments upon UV irradiation. It was found that the hydrogen bonding networks of the DNA bases with surrounding environments play a crucial role in their photostability.²

In recent years, the photoinduced excited-state dynamics of a rare naturally-occurring DNA nucleobase analogue hypoxanthine has been investigated by a range of experimental techniques, where the pump wavelengths are in the range between 255 to 270 nm, slightly longer than the experimentally measured absorption maximum of ~ 250 nm in aqueous solution.³⁻⁵ Temps and co-workers comprehensively studied the ultrafast electronic deactivation dynamics of hypoxanthine and its derivatives by using femtosecond time resolved absorption and fluorescence spectroscopies, and they observed that hypoxanthine returns in aqueous solution to the electronically ground state at a time constant of 0.21 ± 0.08 ps after photoexcitation.³ Peón and co-workers subsequently reported a similar excited-state lifetime ($\tau < 0.20$ ps) by means of femtosecond fluorescence up-conversion experiment.⁴ More recently, Chen and Kohler applied the femtosecond transient absorption spectroscopy to obtain a shorter time constant of 0.13 ± 0.02 ps, and it was viewed as a candidate with the shortest excited-state lifetime among the studied natural or modified nucleobases.⁵

In addition to current experimental studies, quantum chemical computations and dynamics simulations have been directed at understanding hypoxanthine's photophysics.^{4,6-10} This compound has multiple tautomers, in which the keto-N7H and keto-N9H forms are found to be dominant with approximately equal amount in methanol solution (53–54% for keto-N7H)⁴; however, there is no accurate information of their relative population in aqueous solution. According to theoretical calculations, keto-N7H is predicted to be less than 1 kcal mol⁻¹ more stable than keto-N9H in the gas phase.^{9,10} Recently, Peón and co-workers carried out the multireference configuration interaction with single excitations (MR-CIS) calculations to address a readily accessible conical intersection between the bright S₁(¹ππ*) singlet state and the S₀ ground state for both tautomers.⁴ In our previous studies, four direct “*ethylenic*” deactivation pathways from S₁(¹ππ*) to S₀ could be reached through a barrierless way, and the following *ab initio* on-the-fly dynamics simulations in the gas phase lend support to this fact.¹⁰ In addition, we also noticed that the solvatochromic effects on their vertical excitation energies are quite different, with a red-shift of 0.04 eV for keto-N7H and a blue-shift of 0.06 eV for keto-N9H.¹⁰

Despite such important contributions, the sophisticated computational characterization on the excited-state dynamics of keto-N7H and keto-N9H in aqueous solution are still unavailable to our best knowledge, and several key questions about the solvent effect nature and the detailed mechanisms for nonradiative decay processes in solution remain open. For instance: (i) What are the hydrogen-bond features for both forms at different states? (ii) And what are the solvent effects on

their excited-state lifetimes and deactivation behaviors? In an attempt to clarify these issues, we here perform the combined quantum mechanical/molecular mechanical molecular dynamics (QM/MM MD) simulations to explore the ultrafast nonadiabatic decay of two biologically relevant hypoxanthine keto-N7H and keto-N9H tautomers in aqueous solution. It was expected that the state-of-the-art surface-hopping dynamics simulations can improve the understanding of their intrinsic photophysical properties in solution.

2. Computational details

The geometry optimizations of keto-N7H and keto-N9H tautomers in aqueous solution were carried out with the CHEMSHELL 3.5¹¹ package integrating the AMBER 12¹² and TURBOMOLE 6.4¹³ programs. The model was constructed by placing each tautomer into a cubic box of TIP3P¹⁴ water molecules with more than 24 Å length of a side. Herein, the charge and force field parameters were treated by the restrained electrostatic potential (RESP)¹⁵ and the AMBER GAFF¹⁶ schemes, respectively. The 10 Å cutoff was applied for the electrostatic interaction calculations. Following the energy minimization procedure, MD simulations were performed. In particular, the system was heated to 300 K, and then 1 ns equilibration was carried out along with the constant pressure periodic boundary conditions. The SHAKE^{17,18} algorithm was used to constrain all of the bonds related to hydrogen atoms. The 6 sample snapshots extracted from the equilibrated configurations of each solvated base analogue by MD simulations were considered as the starting structures for following

QM/MM optimizations. Here, the QM region including keto-N7H or keto-N9H was computed at the B3LYP/def-SVP level, while the MM region was depicted by the AMBER force field. The hybrid delocalized coordinates (HDLC)¹⁹ technique was finally employed during the geometry optimization.

The hybrid cluster-continuum model was used to calculate the vertical excitation energies of two low-lying excited singlet states, in which the small cluster model containing the base analogue and several explicit water molecules was extracted from the QM/MM optimized equilibrium configuration in aqueous solution. The polarizable continuum model (PCM) was used to explore the bulk solvent effect. In calculations of time-dependent density functional theory (TD-DFT), three functionals including B3LYP, and X3LYP and M06-2X along with an aug-cc-pVDZ basis set were employed. TD-DFT calculations were performed using the TURBOMOLE 6.4¹³ and GAUSSIAN 09²⁰ programs.

Photoinduced nonadiabatic dynamics was performed for the two species by the fewest switches trajectory surface-hopping method²¹, as implemented with the NEWTON-X^{22,23} program integrating the COLUMBUS²⁴⁻²⁶ and TINKER²⁷ packages. The *ab initio* calculations (QM region) were employed at the state-averaged complete active space self-consistent-field (SA-CASSCF) level. Here CAS was composed of ten electrons distributed into eight π -orbitals, namely, CAS(10e/8o). State-averaging over two states was considered, and a 6-31G(d) basis set was employed for all atoms. In order to approximately evaluate the CASSCF performance for the potential energy surface of the low-lying states, the relative energy profiles along the linearly

interpolated internal coordinate (LIIC) reaction path from the S_0 state to the S_0/S_1 conical intersection of keto-N9H were predicted by CASSCF and CASPT2, and a reasonable agreement between both calculations has been found (see Fig. S1 of the ESI†). The molecular mechanics (MM) region was described by the OPLS-AA force field.²⁸ The TIP3P model was chosen for the solvent water, and 833 molecules were included in calculation. One hundred trajectories starting at the S_1 excited state were computed for each tautomer and the maximum simulation time is 400 fs. For keto-N7H, however, trajectories that evolved into the S_0 state and stayed there for more than 100 fs were automatically terminated. The classical equations were integrated using the Velocity-Verlet algorithm²⁹ with a time step of 0.5 fs. Further technical details on the dynamics simulations can be found in previous studies.^{30–33}

3. Results and discussion

3.1 Ground-state structures in water

The representative QM/MM-optimized structures of solvated hypoxanthine keto-N7H and keto-N9H tautomers in their S_0 states are partially plotted in Figs. 1 and 2, respectively. [The 6 optimized geometries for both tautomers from 6 sample snapshots are shown in Figures S2 and S3](#), and the key structural parameters as well as corresponding Cartesian coordinates can be found in Tables S1–S4 (see the ESI†), together with the available experimental crystal data³⁴ for a direct comparison.

We note that both keto-N7H and keto-N9H have planar structures for their S_0 states in the gas phase.¹⁰ However, in aqueous solution, their aromatic rings are found

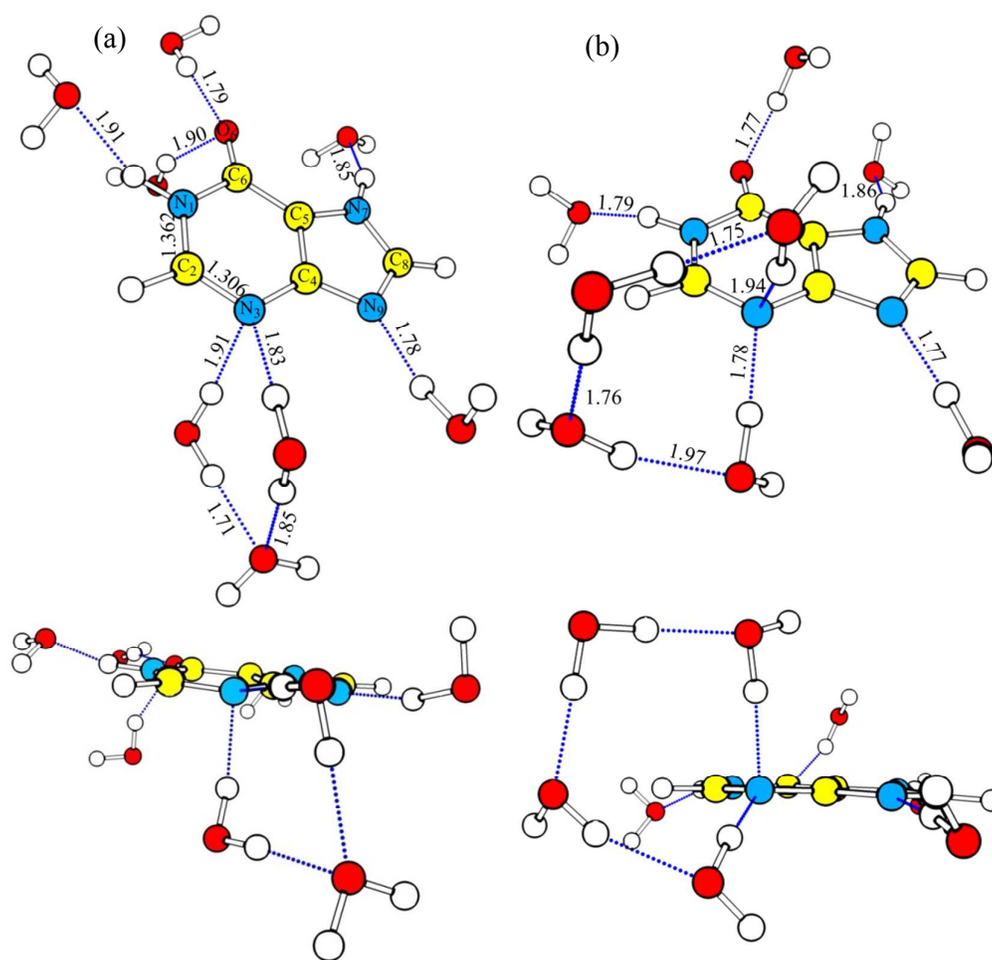


Fig. 1 QM/MM-optimized two representative local structures and atom numbering of hypoxanthine keto-N7H tautomer and only key hydrogen-bond interactions are shown.

Upper panel: top view; lower panel: side view.

to show a slight distortion, with the $\phi(\text{C}_6\text{-C}_5\text{-C}_4\text{-N}_9)$ dihedral angles of 178.0/178.3 and 177.9°, respectively. Structurally, the solvation leads to the elongation of $\text{C}_2\text{-N}_3$ bond by 1.3/1.4 pm for keto-N7H and 0.7 pm for keto-N9H (cf. Table S1). Such notable lengthening in keto-N7H can be attributed to the fact that the π -electron of N_3 atom was involved in the formation of H-bonds with surrounding water molecules as shown in Fig. 1. On the contrary, the $\text{N}_1\text{-C}_2$ bond was shortened by 1.5/2.2 pm for

keto-N7H and 0.7 pm for keto-N9H in solution. It is obvious that present QM/MM calculations reasonably reproduce the experimental structural data (cf. Table S1), and these optimized structures partially displayed in Figs. 1a and 2 can be used to generate the 100 initial configurations through the random-velocities method for the subsequent nonadiabatic dynamics simulations in aqueous solution.

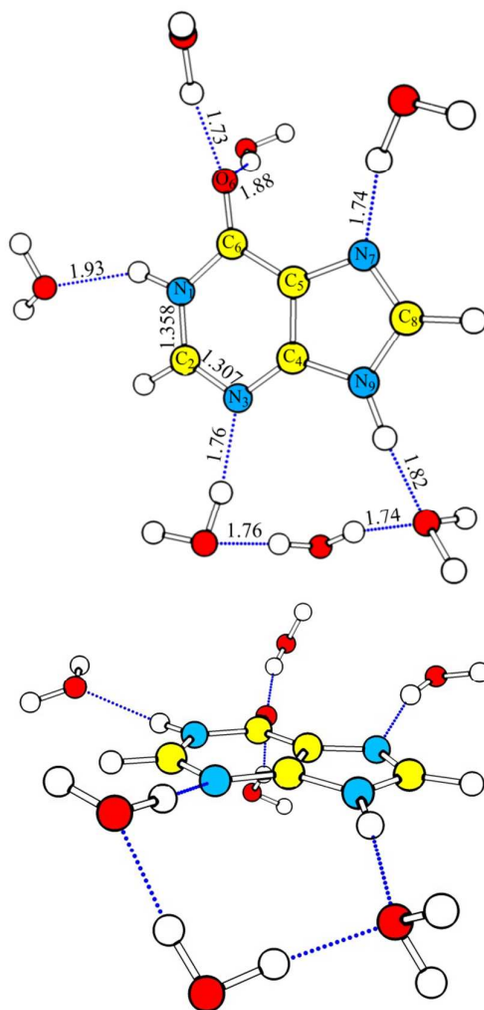


Fig. 2 QM/MM-optimized one representative structure and atom numbering of hypoxanthine keto-N9H tautomer and only key hydrogen-bond interactions are shown.

Upper panel: top view; lower panel: side view.

As shown in Figs. 1 and 2, the H-bond interactions around keto-N9H are slightly stronger than those around keto-N7H, and remarkable differences of the H-bond networks around both base analogues occur at the N₃ site. In keto-N7H, there are two representative local ground-state structures from the QM/MM optimizations based on MD simulations. One structure has two relatively weak H-bonds, $\delta_1(\text{N}_3\cdots\text{H}-\text{O})$ (1.83 Å) and $\delta_2(\text{N}_3\cdots\text{H}-\text{O})$ (1.91 Å), and both are approximately parallel and perpendicular to the ring plane, respectively (cf. Fig. 1a). Similarly, the other structure has also two relatively weak H-bonds, $\delta'_1(\text{N}_3\cdots\text{H}-\text{O})$ (1.78 Å) and $\delta'_2(\text{N}_3\cdots\text{H}-\text{O})$ (1.94 Å), with the structural similarity to $\delta_1(\text{N}_3\cdots\text{H}-\text{O})$ and $\delta_2(\text{N}_3\cdots\text{H}-\text{O})$, respectively (cf. Fig. 1b). It should be mentioned that both $\delta_2(\text{N}_3\cdots\text{H}-\text{O})$ and $\delta'_2(\text{N}_3\cdots\text{H}-\text{O})$ H-bonds locate below or above the ring plane, respectively. These H-bonds in keto-N7H may arise from the contribution of the lone-pair (n_{N}) and π electrons localized at the N₃ atom. Presumably, the presence of $\delta_2(\text{N}_3\cdots\text{H}-\text{O})$ or $\delta'_2(\text{N}_3\cdots\text{H}-\text{O})$ in keto-N7H model may affect distortion of the adjacent C₂-H₂ bond as can be seen in the following nonadiabatic decay process (vide infra). For keto-N9H, a different H-bond network was found among N₃, N₉(H) and three water molecules, with a $\delta_3(\text{N}_3\cdots\text{H}-\text{O})$ H-bond length of 1.76 Å (cf. Fig. 2), which is structurally similar to $\delta_1(\text{N}_3\cdots\text{H}-\text{O})$ of keto-N7H. This clearly indicates that both tautomers of hypoxanthine are inclined to form the different H-bond networks in aqueous solution, and such differences in H-bond interactions may modify their excited-state properties to a certain extent.

3.2 Vertical excitation energies

TD-DFT-calculated vertical excitation energies of the $^1(\pi\pi^*)$ and $^1(n\pi^*)$ excited states of hypoxanthine keto-N7H and keto-N9H tautomers in the gas phase and in aqueous solution are tabulated in Table 1, together with the available experimental findings and our previous CASPT2 results.^{4,10,35} It is worth to mentioning that the TD-X3LYP/PCM calculations have been successfully applied to five naturally-occurring nucleic acid base systems and the maximum deviation of vertical excitation energies between theory and experiment is less than 0.2 eV.³⁶

Table 1 Vertical excitation energies (in eV) and oscillator strengths (in parentheses) of the $^1(\pi\pi^*)$ and $^1(n\pi^*)$ states of hypoxanthine keto-N7H and keto-N9H tautomers in vacuum and water by different theoretical approaches and experiments

Methods	States	Vacuum ^a		Water ^b	
		keto-N7H	keto-N9H	keto-N7H	keto-N9H
TD-B3LYP ^c	$^1(\pi\pi^*)$	4.90 (0.077)	4.71(0.136)	5.12(0.207)	5.03(0.116)
	$^1(n\pi^*)$	5.07	5.20	5.35	5.49
TD-B3LYP ^d	$^1(\pi\pi^*)$	4.91(0.077)	4.72 (0.136)	5.01 (0.101)	4.95 (0.150)
	$^1(n\pi^*)$	5.07	5.22	5.36	5.50
TD-X3LYP ^d	$^1(\pi\pi^*)$	4.93(0.079)	4.74 (0.138)	5.03 (0.113)	4.98 (0.151)
	$^1(n\pi^*)$	5.12	5.18	5.41	5.54
TD-M06-2X ^d	$^1(\pi\pi^*)$	5.18(0.101)	5.07 (0.149)	5.25 (0.206)	5.31 (0.125)
	$^1(n\pi^*)$	5.36	5.35	5.63	5.71
CASPT2 ^e	$^1(\pi\pi^*)$	4.79 (0.122)	4.58 (0.167)	4.75 (0.128)	4.64 (0.175)
Expt. ^f				4.98	
Expt. ^g			4.41		4.59

^a B3LYP/def-SVP geometry. ^b B3LYP/def-SVP/MM geometry. ^c QM(molecular + explicit waters)/MM. ^d QM (molecular + explicit waters)/PCM. ^e CCSD/6-31G(d) geometry.¹⁰ ^f Experimental value of hypoxanthine in water at PH = 5.⁴ ^g Experimental values in the vapor gas and water at PH = 6.1 from 9-methyl-hypoxanthine.³⁵

It can be seen from Table 1 that the vertical excitation energies predicted by the TD-B3LYP and TD-X3LYP methods are close to each other, while TD-M06-2X approach overestimates the vertical excitation energies by about 0.22–0.33 eV, compared to the TD-X3LYP results. At the TD-X3LYP level of theory, the vertical

excitation energies of the $^1(\pi\pi^*)$ states of keto-N7H and keto-N9H in water lie at 5.03 and 4.98 eV, respectively, showing good agreement with the available experimental value of 4.98 eV.⁴ This indicates that the maximum absorption band observed in experiment may arise from the superposition of the $^1(\pi\pi^*)$ states of the two dominant hypoxanthine tautomers, which is in accordance with the previous calculations.^{4,6} On the other hand, TD-X3LYP-predicted $^1(\pi\pi^*)$ states of keto-N7H and keto-N9H in aqueous solution are blue-shifted by 0.10 and 0.24 eV, respectively, relative to the gas phase, whereas both $^1(n\pi^*)$ states exhibit the more remarkable blue shift (0.29 and 0.36 eV, respectively). We note that such a blue shift of hypoxanthine keto-N9H tautomer is nicely consistent with that of the 9-methyl-hypoxanthine analogue (0.18 eV).³⁵ Considering that the relatively high $^1(n\pi^*)$ state is generally less accessible optically, it could be not expected in the subsequent nonadiabatic decay process. In the following, we will focus only on the nonadiabatic dynamics of the optically bright $^1(\pi\pi^*)$ states of keto-N7H and keto-N9H in aqueous solution (vide infra).

3.3 Surface hopping excited-state dynamics

Fig. 3 shows the time evolution of average occupations of the S_1 states from respective 100 trajectories of keto-N7H and keto-N9H in aqueous solution. It can be clearly seen that after photoexcitation into the optically bright state, a faster $S_1 \rightarrow S_0$ deactivation takes place for keto-N7H.

To estimate the excited-state lifetime of keto-N7H, we have fitted its occupation of the S_1 state using a mono-exponential model within the timescale of 200 fs:

$$y = y_0 + (1 - y_0)\exp[-(t - t_1)/t_2]$$

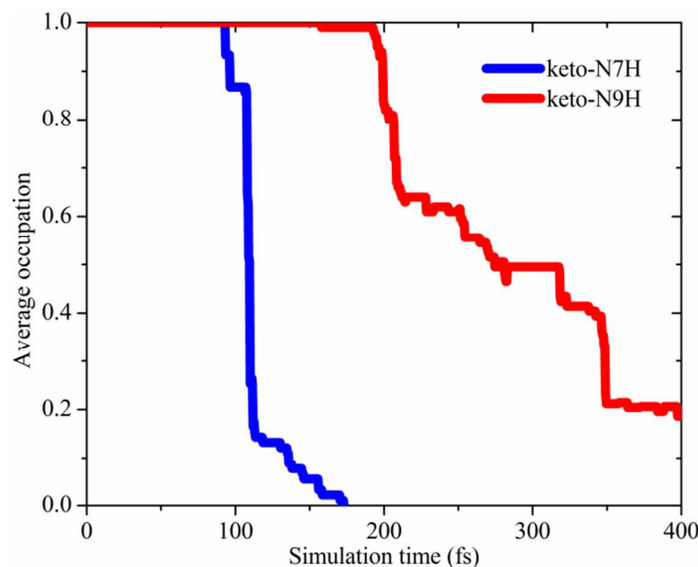


Fig. 3 Time evolution of average occupations of the S_1 states of keto-N7H and keto-N9H in aqueous solution.

where y_0 is the fraction of the population which does not relax to the S_0 state, t_1 is the initial delay or latency time, and t_2 is the exponential time constant. The excited-state lifetime (τ) is the sum of t_1 and t_2 . The fitting process leads to $y_0 = 0$, $t_1 = 96.1$ fs, $t_2 = 16.5$ fs, and $\tau = 112.6$ fs (correlation coefficient $R^2 = 0.88$), suggesting that the dynamics of keto-N7H in aqueous solution shows a latency time of 96.1 fs before starting an ultrafast and exponential nonadiabatic decay with a decay constant of 16.5 fs. We note that the excited-state lifetime of keto-N7H in water is slightly shorter than the gas-phase result of 137.7 fs,¹⁰ and such a similarity at two different environments shows good agreement with the available experimental findings.³⁻⁵ It is worthy of note that the dynamics of keto-N7H in the S_1 state is not strongly affected by steric interaction with solvent molecules.

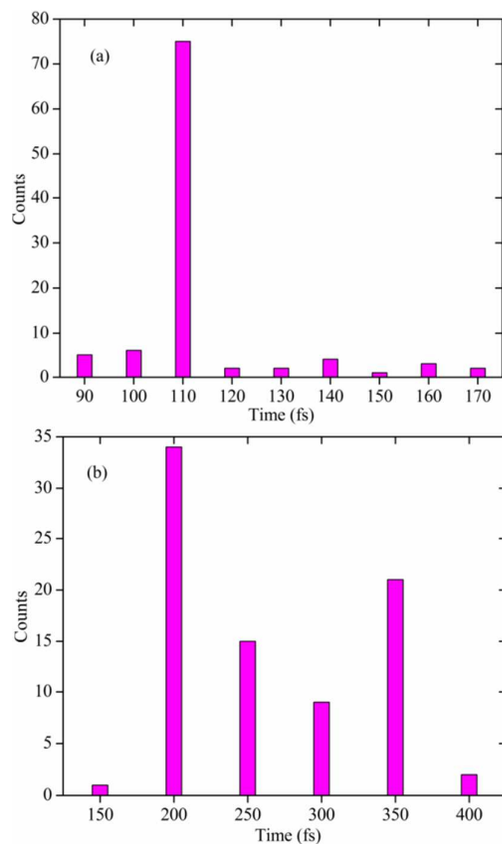


Fig. 4 Time distributions of keto-N7H (a) and keto-N9H (b) at the $S_1 \rightarrow S_0$ hopping points.

Unfortunately, such single exponential model cannot be applied to the fitting of the population of the S_1 state for keto-N9H. To obtain an in-depth insight into this discrepancy between photodynamical behaviors of keto-N7H and keto-N9H in water, we have, in Fig. 4, plotted both time distributions at the $S_1 \rightarrow S_0$ hopping points. We note that in calculated 100 trajectories of keto-N7H, more than 70 ones switch to their S_0 states with time constants of approximate 110 fs, in accordance with the above estimated lifetime of 112.6 fs. However, the time distribution of keto-N9H differs from that of keto-N7H. It basically locates at two time ranges from 200 to 250 fs and from 300 to 350 fs, and thus keto-N9H exhibits a longer excited-state lifetime in

aqueous solution, compared to keto-N7H, which is still close to the experimentally observed results mentioned above.³⁻⁵ It is a reasonable assumption that keto-N7H and keto-N9H tautomers are in a thermodynamic equilibrium in aqueous solution, both of which may be responsible for the ultrafast nonradiative deactivation of photoactive hypoxanthine observed in experiment.³⁻⁵

Quite surprisingly, the $S_1 \rightarrow S_0$ decay of keto-N9H in solution is much slower than its gas-phase counterpart with the lifetime of 85.5 fs,¹⁰ which is different from keto-N7H and 9H-gaunine analogues.³⁷ We note that the C_2-N_3 bond lengths of keto-N7H at the state-state hopping points mostly distribute below 1.4 Å, whereas those in keto-N9H generally spread above 1.4 Å, and a similar phenomenon can be also found for the N_1-C_2 bond (cf. Fig. S4 in ESI†). Such structural features indicate that at the $S_1 \rightarrow S_0$ hopping points keto-N9H has much more expanding geometry than keto-N7H. Additionally, in consideration of a shorter N_1-C_2 bond of keto-N9H in water, its conical intersection structure is less accessible, compared to keto-N7H. Therefore, keto-N9H has to spend more time to evolve into the hopping point once it is promoted to the Frank-Condon region, resulting in a long excited-state lifetime.

Fig. 5 shows the distributions of the $\phi(C_4-N_3-C_2-H_2)$ dihedral angles at the $S_1 \rightarrow S_0$ hops of keto-N7H and keto-N9H. Interestingly, each of the trajectories in keto-N7H decays to the S_0 state via an S_1/S_0 conical intersection with an upward distortion puckering at the C_2 atom. The ϕ -values of more than one half of the trajectories lie around -110° , partially around -90° (cf. Fig. 5a), which are significantly different from those in the gas phase where two slightly different kinds

of conical intersections were found with approximately equal branching ratios and the corresponding ϕ -values are -82.2 and -111.5° , respectively.¹⁰ This means that one decay channel would dominate the ultrafast nonadiabatic decay process of keto-N7H in aqueous solution. Such a behavior of keto-N7H in solution may be ascribed to the H-bonding interaction contributed by π -electron at N₃ site (cf. Fig. 1a). Presumably, the presence of such π -electron H-bond below the ring plane may facilitate the C₂-H₂ upward out-of-plane distortion to the hopping point. This is one reason why the S₁ → S₀ decay of keto-N7H in solution is slightly faster than that in vacuum.

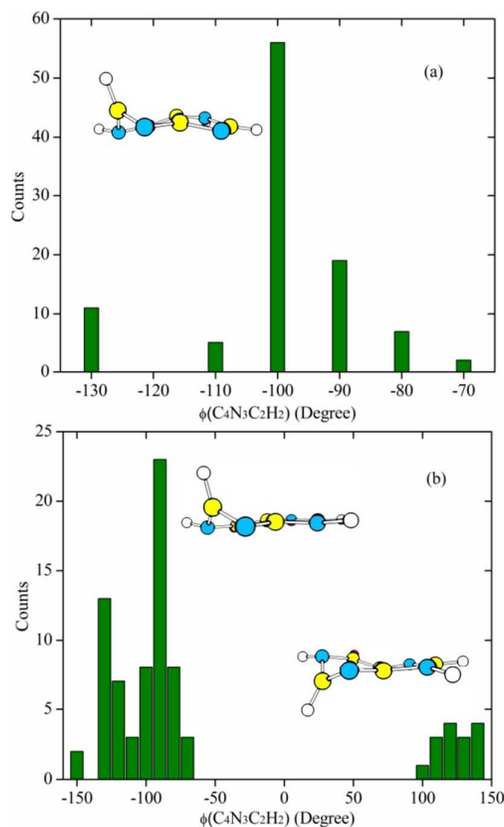


Fig. 5 Distributions of the $\phi(\text{C}_4\text{-N}_3\text{-C}_2\text{-H}_2)$ dihedral angles at the S₁ → S₀ hops of keto-N7H (a) and keto-N9H (b). The geometries shown as insets are the representative hopping structures.

As mentioned above, there is another representative structure with the π -electron H-bond above the ring plane of keto-N7H (cf. Fig. 1b). Presumably, the presence of such π -electron H-bond of keto-N7H may promote the downward distortion puckering at the C₂ atom during evolution to the conical intersection, like its counterpart (cf. Fig. 1a). Accordingly, both representative solvated structures of keto-N7H shown in Fig. 1 have similar excited-state dynamical behaviors.

For keto-N9H, 82 out of 100 trajectories can relax back to the S₀ states, where the conical intersection structures of 67 ones show an upward deformation puckering at the C₂ atom, while 15 ones are predicted to have a downward distortion at the same position. The remaining 18 are trapped in the S₁ states. As displayed in Fig. 5b, its ϕ -values mainly locate at two regions between 100 and 140° and between -80 and -130°, respectively. Obviously, keto-N9H exhibits more patterns of conical intersections than keto-N7H in our present dynamics simulations, due to the absence of π -electron H-bond.

Fig. 6 presents the distributions of four key H-bonds at the S₁ → S₀ hopping points of keto-N7H and keto-N9H. It is apparent that all S₁ → S₀ hops in keto-N7H lead to the shortening of $\delta_2(\text{N}_3\cdots\text{H}-\text{O})$ and simultaneously the lengthening of $\delta_1(\text{N}_3\cdots\text{H}-\text{O})$ H-bond lengths (cf. Fig. 6a). The significantly enhanced $\delta_2(\text{N}_3\cdots\text{H}-\text{O})$ H-bond can promote the evolution of keto-N7H at the S₁ state to the conical intersection, with the upward distortion as mentioned above. On the contrary, the $\delta_3(\text{N}_3\cdots\text{H}-\text{O})$ H-bond of keto-N9H becomes much weaker at most of S₁/S₀ hopping points (61 trajectories) than in the ground state (1.76 Å) (cf. Fig. 6b). Clearly, the

remarkable differences in intermolecular H-bonding interactions around keto-N7H and keto-N9H account for their distinct excited-state dynamical behaviors such as excited-state lifetimes and deactivation pathways in water.

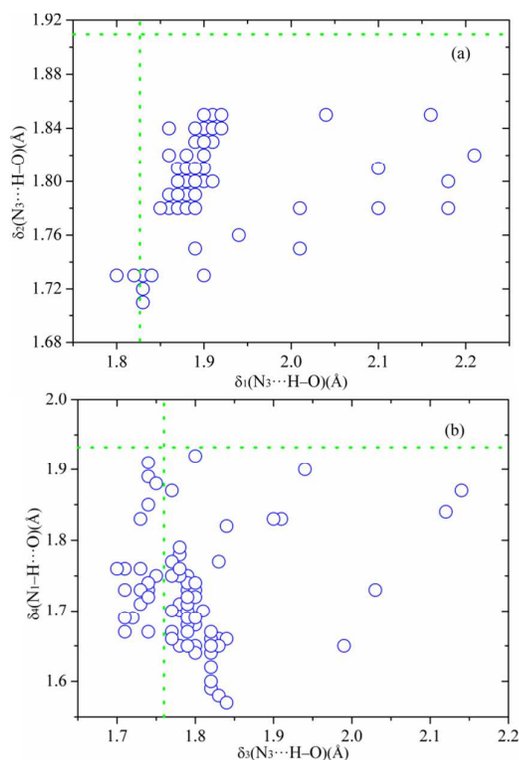


Fig. 6 Distributions of key H-bond lengths at the $S_1 \rightarrow S_0$ hopping points of keto-N7H (a) and keto-N9H (b). Green lines represent the corresponding values in their ground states: (a) 1.83 and 1.91 Å; (b) 1.76 and 1.93 Å.

4. Conclusions

The excited-state dynamics of two most stable hypoxanthine tautomers (keto-N7H and keto-N9H) in aqueous solution has been investigated by means of the hybrid QM/MM MD simulations. The present results suggest that the $S_1 \rightarrow S_0$ decay of keto-N7H in water is slightly faster, while the evolution of keto-N9H to S_0 state is

much slower than in vacuum. In aqueous solution, there is one dominant decay pathway for keto-N7H, but the decay channels of keto-N9H are less sensitive to the solvent effect. Such distinct photodynamical behaviors can be ascribed to the H-bond network differences around keto-N7H and keto-N9H in water environment. The presence of π -electron H-bond at N₃ site and its evolution along the conical intersection formation play an important role in the photodynamics of keto-N7H in water. The structural features of S₁/S₀ hopping configurations and their accessibilities provide a basis for understanding of their excited-state lifetimes and possible nonradiative decay channels in aqueous solution.

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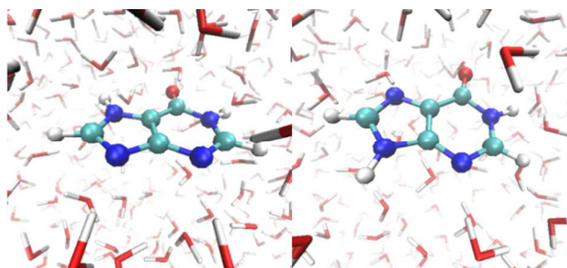
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Graphical Abstract

QM/MM MD simulations reveal different H-bond networks around DNA base analogues, and the π -electron H-bond in the solvated keto-N7H may facilitate its $S_1 \rightarrow S_0$ nonadiabatic decay.