Intersystem crossing-branched excited-state intramolecular proton transfer for o-nitrophenol: An ab initio on-the-fly nonadiabatic molecular dynamic simulation

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The 6SA-CASSCF(10, 10)/6-31G (d, p) quantum chemistry method has been applied to perform on-the-fly trajectory surface hopping simulation with global switching algorithm and to explore excited-state intramolecular proton transfer reactions for the o-nitrophenol molecule within low-lying electronic singlet states (S₀ and S₁) and triplet states (T₁ and T₂). The decisive photoisomerization mechanisms of o-nitrophenol upon S₁ excitation are found by three intersystem crossings and one conical intersection between two triplet states, in which T₁ state plays an essential role. The present simulation shows branch ratios and timescales of three key processes via T₁ state, non-hydrogen transfer with ratio 48% and timescale 300 fs, the tunneling hydrogen transfer with ratios 36% and timescale 10 ps, and the direct hydrogen transfer with ratios 13% and timescale 40 fs. The present simulated timescales might be close to low limit of the recent experiment results.

The excited-state intramolecular proton transfer (ESIPT) reaction is considered to be one of the most fundamental and important processes in chemistry, material and biology. The ESIPT reactions have been studied from both ab initio quantum chemistry calculations and trajectory-based nonadiabatic molecular dynamic simulations. The most of the ESIPT processes have been found to be taken place at a subpicosecond time scale. The photo-excited molecule from electronic ground singlet state (S₀) to the excited singlet states can undergo rapid internal conversion via conical intersections (CI) to the lower singlet states as well as intersystem crossings (ISC) via spin-orbital couplings (SOC) to the triplet states. By means of the radiative or nonradiative interaction with target molecule, the redistribution of electron density is essential for the transfer of a hydroxyl (or amino) proton to an oxygen or nitrogen acceptor atom on the excited states within a hydrogen bond already formed in the electronic ground state. Subsequently, the excited state molecule can decay to the ground state accompanied with a reverse proton transfer.

Fundamental mechanistic insight into photoinduced ESIPT reaction can be understood through studying prototypical representative of nitroaromatics o-nitrophenol molecule. This is because that o-nitrophenol owns two functional groups, namely nitro and the adjacent hydroxyl groups which show strong intramolecular hydrogen bond. Thus, the aci-nitrophenol can be formed via the initial intramolecular hydrogen transfer from hydroxyl group to nitro group. Differing from the other aromatic systems, the experimental and theoretical studies demonstrate peculiar behavior of the photoinduced decay process for nitrated aromatic compounds. The tautomeric aci-nitrophenol isomers via photoisomerization has been confirmed by infrared spectroscopy in low-temperature...
argon matrices\textsuperscript{23} and by laser-induced-fluorescence\textsuperscript{24}. The rich intersystem crossing network makes \(\text{o-nitrophenol}\) from the first excited singlet state decaying to the triplet states rapidly, and the process occurs in the femtoseconds to a few picoseconds time scale dependent on the intermediate states\textsuperscript{25–27}. The early experiment showed that time scale decaying to the first excited triplet state is lower than 50 ps by measuring the UV excitation of \(\text{o-nitrophenol}\) in benzene solvent\textsuperscript{28}. However, the recent experiment showed the existence of unstable aci-nitrophenol isomers and ultrafast ISC to the triplet manifold on a subpicosecond time scale by both femtosecond transient absorption spectroscopy in solution and time-resolved photoelectron spectroscopy in the gas phase\textsuperscript{29}.

Up to now, neither fluorescence nor phosphorescence has been reported for \(\text{o-nitrophenol}\) and this indicates the existence of pure ultrafast radiationless decay process\textsuperscript{29}. Therefore, an ab initio on-the-fly nonadiabatic molecular dynamics simulation is necessary to be performed to quantitatively analyze ESIP\textsuperscript{T} process of \(\text{o-nitrophenol}\) via internal conversion and intersystem crossing network. This is motivation of the present study. By using ab initio quantum chemistry method at 6SA-CASSCF(10, 10)/6-31G (\(d, p\)) and MRCI/cc-pVDZ level, we previously optimized geometries for all isomers and transition states on two low-lying singlet (\(S_0\) and \(S_1\)) and two low-lying triplet (\(T_1\) and \(T_2\)) electronic states and we computed all four-state potential energy profiles along ESIP\textsuperscript{T} coordinates\textsuperscript{30}. We confirmed the existence of unstable aci-nitrophenol isomers as observed in the early studies\textsuperscript{23,24,30}.

We found total five ISC zones with SOCs at \(\sim 10\) and \(\sim 40\) wavenumbers. By performing nonadiabatic molecular dynamics simulation in the present study, we actually found one new \(S_0/T_1\) ISC which governs decay process for non hydrogen transfer. The previous computational and experimental study has proposed two possible relaxation pathways\textsuperscript{35}; one is via the ISC following (\(\text{O})\text{H-O(NO)}\) stretch and the other is via the \(S_0/S_1\) CI after overcoming potential energy barrier induced by hybrid torsion of HONO group and (\(\text{O})\text{H-O(NO)}\) stretch. Actually, the new \(S_0/T_1\) ISC is located near Frank-Condon region while \(S_0/S_1\) CI is located in configuration after hydrogen transfer. The present simulation has shown that the new \(S_0/T_1\) ISC is very effective pathway and there is almost no trajectory going via \(S_0/S_1\) CI. We can expect that there must have strong intersystem crossing-branching ESIP\textsuperscript{T}. Besides, the existence the certain potential barriers on \(S_1\) and \(T_1\) states along ESIP\textsuperscript{T} coordinate can add tunneling effects as well. The present study should reveal entire picture of photosomerization reaction and the photo-decay mechanisms including ESIP\textsuperscript{T} for \(\text{o-nitrophenol}\) molecule.

Trajectory-based ab initio nonadiabatic molecular dynamics simulations have been successfully applied to photosomerization and photoreaction processes involving intersystem crossings and conical intersections\textsuperscript{31–48}. Recently, within Tully’s fewest switching algorithm in diabatic representation dynamic simulations have been carried out for intersystem crossings\textsuperscript{43–48}. We have developed trajectory-based ab initio nonadiabatic molecular dynamics simulations without involving calculation for nonadiabatic couplings\textsuperscript{37}, and it has been applied for multiple-state azobenzene photosomerization via conical intersections\textsuperscript{38}. Now we extend our analytically global switching probability method to include intersystem crossing with spin-orbital couplings. Trajectories are calculated on on-the-fly ab initio singlet or triplet potential energy surfaces and then by applying global switching algorithm we treat trajectory surface hopping in a unified way for internal conversion and intersystem crossing processes. Intersystem-crossing dynamic simulation with global switching surface hopping algorithm is not really new with Landau-Zener switching probability; the spin-diabatic and spin-adiabatic dynamic simulations\textsuperscript{31} were performed for the model system of spin-orbital couplings that are used as diabatic coupling parameter in Landau-Zener formula, and moreover good agreement with experimental observation for the triplet to singlet branching ratio has been achieved with Landau-Zener formula for performing real dynamic simulation of O(\(\text{P}) + \text{ethylene}\textsuperscript{32}. We utilize improved Landau-Zener formula, namely Zhu-Nakamura to treat intersystem-crossing dynamic simulation in the present report. Moreover, tunneling effects are treated by one-dimensional semiclassical method along ESIP\textsuperscript{T} coordinate\textsuperscript{39}.

**Results**

**Global switching algorithm.** Global switching algorithm makes trajectory hopping at the time \(t\) where \(d(t) = |U_2(t) - U_1(t)|/|V_2(t) - V_1(t)|\) reaches local maximum (\(U_2\) and \(U_1\) are adjacent two adiabatic potential energy surfaces, while \(V_2\) and \(V_1\) are corresponding two diabatic potential energy surfaces). Analytical switching probability can be generally expressed in terms of \(d(t)\) as\textsuperscript{50}

\[
\sinh\left(\frac{d^2 \Delta \delta}{2}\right) \exp\left(-\frac{\Delta \delta}{2}\right) \exp\left(-\frac{\Delta \delta}{2}\right),
\]

where

\[
\Delta \delta = \pm \frac{1}{2} \Delta \delta\]

in which \(\Delta \delta\) is estimated from two potential energy surfaces and its gradients at hopping spot along on-the-fly running trajectory. In the case of \(\Delta \delta \gg 1\), it basically goes to Landau-Zener or improved version Zhu-Nakamura formula\textsuperscript{31,32}.

\[
P = \begin{cases} 
\exp(-2\Delta \delta) & \text{between two adiabatic potentials} \\
1 - \exp(-2\Delta \delta) & \text{between two diabatic potentials} 
\end{cases}
\]

(2)

where

(3)

in which the effective coupling parameter \(a^2\) and effective collision energy \(b^2\) are given by
µ = \frac{-aFF - a}{V_{\text{E}E} FF (2V_{12})^2} \quad \text{and} \quad b^2 = \frac{(E_1 - E_X)}{\sqrt{2F_1 FF (2V_{12})}}

where \( F_1 \) and \( F_2 \) are forces on two diabatic potential energy surfaces, \( V_{12} \) is diabatic coupling, \( \mu \) is reduced mass and \( E_X \) is energy at crossing point and \( E_t \) is potential energy plus kinetic energy component in direction of hopping direction. All those quantities in equation 4 are calculated along on-the-fly running trajectory at hopping spot 37 and its details are also given in the Supplementary Note 1. The hopping direction in the present theory is defined based on sort of the local modes and this agrees with the normal modes constructed from the regularized diabatic states 53. Global switching algorithm has been extensively compared with the Tully’s fewest switching algorithm 37, 40 and two algorithms are basically similar for highly averaged quantities like quantum yields and lifetimes. In the present on-the-fly simulation, along three consecutive time steps we detect minimum energy separation between two singlet-states (\( S_1 \) and \( S_0 \)) or two triplet-states (\( T_2 \) and \( T_1 \)) and this gap is considered as diabatic coupling \( V_{12} \) in equation 4. In this case, the switching is considered as in between two adiabatic potential energy surfaces. On the other hand, we detect the local maximum \( d(t) \) between the singlet and triplet states. The diabatic coupling is considered as spin-orbital coupling as \( V_{12} = \text{SOC} \) in equation 4 as proposed in ref. 43., and in fact, singlet state \( V_1 \) and triplet state \( V_2 \) can be transformed into adiabatic representation according to

\[
U_{1,2} = \frac{V_1 + V_2}{2} \pm \frac{1}{2} \sqrt{(V_1 - V_2)^2 + 4V_{12}^2}
\]

However, in this case, we can do the switching directly between the singlet and triplet states as the spin-orbital coupling is known.

Intersystem crossings with ab initio dynamics. We have previously performed ab initio quantum chemistry calculations at 6SA-CASSCF(10, 10)/6-31G (d, p) level and have compared with MRCI/cc-pVDZ energy corrections for two singlet states (\( S_0 \) and \( S_1 \)) and two triplet states (\( T_1 \) and \( T_2 \)). It is well-known that the CASSCF method is lack of dynamic correlation. Therefore, we carried out energy correction by multi-reference configuration interaction (MRCI) calculation at CASSCF optimized geometries. Both vertical excitation energies (see Fig. S1(a) at CASSCF and Fig. 1S(b) at MRCI given in Supplementary Note 2) and adiabatic energies (see Fig. S2(a) at CASSCF and Fig. 2S(b) at MRCI given in Supplementary Note 2) at all key geometries show the same energy sequences. Relative energy differences between CASSCF and MRCI are mostly smaller than 0.1 eV, especially for energy gaps at six intersystem crossings and one conical intersection being even smaller (see Table S1). Therefore, we think that the present on-the-fly dynamical simulation based on CASSCF level should present the almost same results as it is based on MRCI level. We have confirmed 6SA-CASSCF(10, 10)/6-31G (d, p) is suitable choice for the present dynamic simulation.

We found there are two intersystem crossings between \( S_1 \) and \( T_1 \) (\( S_1T_1\text{-IC1} \) and \( S_1T_1\text{-IC2} \)) around which SOC can be almost considered as constant 10 cm\(^{-1}\), one between \( S_1 \) and \( T_2 \) (\( S_1T_2\text{-IC} \)) around which SOC is 40 cm\(^{-1}\), three between \( S_0 \) and \( T_1 \) (\( S_0T_1\text{-IC1}, S_0T_1\text{-IC2} \) and \( S_0T_1\text{-ICX} \)) around which SOCs are close to 40 cm\(^{-1}\). The \( S_0T_1\text{-ICX} \) is newly found in the present study by trajectory surface hopping dynamic simulation, and the other five were optimized previously 30. Furthermore, the present dynamic simulation shows that only three of six are active for ESIPT reaction, namely \( S_1T_1\text{-IC1}, S_1T_1\text{-ICX}, \) and \( S_0T_1\text{-IC1} \) as shown in Fig. 1. The \( S_1T_1\text{-IC} \) is less active, and \( S_1T_2\text{-IC} \) and \( S_1T_2\text{-IC2} \) are not active at all (key geometries and Cartesian coordinates for all six intersystem crossings are given in Supplementary Note 2 and Note 4). Figure 1 shows that the dihedral angels \( C_4C_5O_11H_12 \) are zero for \( S_0T_1\text{-ICX} \) and \( S_1T_1\text{-IC1} \), and the \( O_11H_12 \) bond lengths are 0.945 Å and 0.955 Å, respectively. However, unlike to the planar geometry of \( S_1T_2\text{-IC1} \), the nitro group of \( S_0T_1\text{-ICX} \) and \( S_1T_1\text{-IC1} \) show a similarity although the \( H_12 \) bonded to \( O_11 \) in \( S_0T_1\text{-ICX} \) instead of \( O_{15} \) in \( S_1T_1\text{-IC1} \). The nitro group of them is out of the aromatic skeleton. Those geometry differences essentially govern the three intersystem crossings undergoing distinct ESIPT reaction pathways. Interestingly, the structure of \( S_1T_1\text{-ICX} \) is close Frank-Condon geometry and the structure of \( S_0S_1\text{-CI} \) is close to \( S_0T_1\text{-IC1} \) geometry (see Fig. S3 given in Supplementary Note 2). The \( S_0T_1\text{-ICX} \) is responsible for...
relaxing pathway of non-hydrogen transfer, while the \( S_0 T_1 - IC1 \) responsible for relaxing pathway of after-hydrogen transfer.

The initial condition of trajectories is started from Franck-Condon region of \( \sigma \)-nitrophenol. We perform frequency calculation at the ground state equilibrium geometry of \( S_0 \) to obtain the normal mode coordinates. Initial normal-mode coordinates and velocities of trajectory are selected according to the Wigner distribution on \( S_0 \) state. Finally, these initial normal-mode coordinates and velocities are converted into Cartesian coordinates and velocities plus vertical excitation energy to excited \( S_1 \) state. The thermal kinetic energy with \( T = 300 \text{K} \) is added to all sampling trajectories with randomly distributing into initial Wigner velocities. However, such equally distributed initial conditions are not suitable for stimulating ESIPT reaction, there are the certain vibronic modes enhanced more than the others as molecule absorbed light to be excited to \( S_1 \) state. We found that four normal modes (\( O_1 H_{12} \) stretch (4172 cm\(^{-1}\)), \( C_6 O_1 H_{12} \) bend (1473 cm\(^{-1}\)), and two \( C_6 O_1 H_{12} - C_5 N_{13} O_{15} \) scissor (408 cm\(^{-1}\) and 310 cm\(^{-1}\)) involved in the ESIPT reaction path have stronger vibronic couplings than the others. Therefore, we added extra kinetic energy (equivalent to \( T = 500 \text{K} \)) to these four normal modes. Along an on-the-fly running trajectory, the nuclear coordinates and velocities in Cartesian coordinates are propagated by numerically integrating the Newtonian equation of motion with the velocity-Verlet method\(^{54}\). We determine the minimum potential energy gap between \( S_1 \) and \( S_0 \) states, and between \( T_1 \) and \( T_2 \) states at the conical intersection zones, and determine maximum \( \delta \) between singlet and triplet states at intersystem crossing zones, at which we compute the effective coupling parameter \( \alpha^2 \) and the effective collision energy \( \beta^2 \). We found SOC varies very slowly at intersystem crossing zones, so that we choose constants accordingly \( 10 \) and \( -40 \text{cm}^{-1} \) in simulation (when the energy gap is smaller than 0.25 eV, we start to check attempted trajectory surface hopping). We have run several expensive on-the-fly SOC trajectories in comparison with fixed SOC trajectories and results show small difference. We have checked for four active intersystem crossings, relative spin-orbital coupling variations (\( \delta \text{SOC/SOC} \)) are about less than 2% around crossing zones in which trajectory hops take place. This fixed SOC technique was also utilized for performing real dynamic simulation of \( \text{O}(\text{P}) + \text{ethylene} \).\(^{32}\)

Due to the present dynamics involving \( \text{OH} \) stretch vibration (over 3500 cm\(^{-1}\)) along ESIPT reaction path, we made test runs with time steps of 0.05 fs, 0.1 fs and 0.15 fs, and finally we set up the 0.1 fs time step from the beginning to the end for the entire dynamic simulation. The dynamics simulation time is set up as 500 fs and as we increase up to 1000 fs, there is no notable difference. All the quantum chemical calculations of on-the-fly potential energy surfaces and its gradients are carried out at 6SA-CASSCF(10, 10)/6-31G (d, p) level by using the quantum chemistry package MOLPRO 2009.1.\(^{35}\) and the dynamic simulation is carried by our own code.

**On-the-fly trajectory analysis.** We have run total 280 trajectories and the outcomes show that 6 stay on \( S_1 \) state regarded as resonance, 6 go via \( S_1 T_1 - IC1 \) as direct hydrogen transfer via \( T_1 \) state, and 268 go via \( S_1 T_2 - IC1 \) initially as shown in Fig. 2 (this is basically Fig. 7 in ref. 30). After going via \( S_1 T_2 - IC1 \), 265 trajectories hop to \( T_1 \) states via conical intersection between \( T_1 \) and \( T_2 \) states, and 3 go direct hydrogen transfer via \( T_1 \) state. Further three bifurcations take place for the 265 trajectories on \( T_1 \) state, 101 regarded as resonance on \( T_1 \) state, 133 going along this path and the other two (\( S_1 T_1 - IC2(SOC = 40 \text{cm}^{-1}) \) and \( S_1 T_1 - IC2(SOC = 10 \text{cm}^{-1}) \)) are far away. Number in parentheses associated with each intersystem crossing represents trajectories going via it. The small widow stands for potential energy profiles for newly found \( S_0 T_1 - ICX \). The Cartesian coordinates of six ICs are summarized in the Supporting Information.

![Figure 2. Potential profiles along ESIPT reaction coordinates.](image-url)

**Figure 2. Potential profiles along ESIPT reaction coordinates.** Four intersystem crossings (\( S_1 T_1 - IC1(SOC = 10 \text{cm}^{-1}) \), \( S_1 T_1 - IC(SOC = 40 \text{cm}^{-1}) \), \( S_1 T_1 - IC1(SOC = 40 \text{cm}^{-1}) \) and \( S_1 T_1 - ICX(SOC = 40 \text{cm}^{-1}) \)) are approximately along this path and the other two (\( S_1 T_1 - IC2(SOC = 40 \text{cm}^{-1}) \) and \( S_1 T_1 - IC2(SOC = 10 \text{cm}^{-1}) \)) are far away. Number in parentheses associated with each intersystem crossing represents trajectories going via it. The small widow stands for potential energy profiles for newly found \( S_0 T_1 - ICX \). The Cartesian coordinates of six ICs are summarized in the Supporting Information.
states for a while. On the other hand, the present ab initio dynamic simulation confirms that intersystem crossing zone between the $S_1$ and $T_2$ states with small SOC (10 cm$^{-1}$) are much wider than intersystem crossing zone between the $S_1$ and $T_1$ states with large SOC (40 cm$^{-1}$). Therefore, without performing ab initio dynamic simulation, we could not realize that $S_1 \rightarrow T_1$ intersystem crossing (ISC-$S_1T_1$) channel is paramount relaxing pathway. Actually, similar situation is found that the efficient ultrafast ISC dynamic can still be taken place with a rather small spin-orbit coupling due to the counterbalance mechanism. Further detailed distribution is shown in Fig. 3b for the sampling trajectories passing through the isomerization of the o-nitrophenol and aci-nitrophenol.

In the present simulations, we defined the aci-nitrophenol isomers as aci-isomer1 and aci-isomer2, respectively. For aci-isomer1, it still exhibits intramolecular hydrogen bond with ortho oxygen atom of benzene and while for aci-isomer2, the hydrogen rotates with nitro group and far away from the ortho oxygen atom. Diverse distributions are shown, for instance, that o-nitrophenol overcomes barrier of hydrogen transfer and reaches the aci-isomer1 (13.33%), after getting over the hydrogen transfer barrier it passes to aci-isomer1 and then back to o-nitrophenol (42.22%), the o-nitrophenol converts to aci-isomer1 and then continue to produce aci-isomer2 with the inversion of $O_3H_{12}$ (13.33%), the o-nitrophenol changes to aci-isomer1 and then regenerated again after going via aci-isomer2 (13.33%), and so on. Starting from Frank-Condon region on $S_0$ state, among 280 sampling trajectories there are 3, 37, and 5 trajectories undergoing direct hydrogen transfer on $T_2$, $T_1$, and $S_0$ states, respectively. However there are 6 and 101 trajectories undergoing tunneling hydrogen transfer on $S_0$ and $T_1$ states, respectively. Although direct hydrogen transfer on $T_1$ is rare case, it presents a new perspective for general ESIPT reactions. In most of situation, the $T_2$ state looks more like a "hub", trajectories stop on it for a moment and then quickly decay to the reactive $T_1$ state where is major channel for direct hydrogen transfer. The barrier for hydrogen transfer on $T_1$ state is lower than that of on $S_0$ state as shown in Fig. 2. On the other hand, the present dynamic simulation shows that kinetic energy from the four vibartional normal modes responding for hydrogen transfer easily dissipates to the other vibrational modes on $S_0$ state much faster than on $T_1$ state. This is part of reason that direct hydrogen transfer does not occur on $S_1$ state. Hydrogen transfer has a peculiar property on $S_0$ state which differs from that of on $T_1$ state, trajectory reaches to $S_0$ state via multi-steps continuous hops in a relatively short period of time and then it runs on the ground state, eventually the hydrogen transfer reaction occurred and aci-isomer formed. It should be noted that the aci-isomer can undergo backward hydrogen transfer reaction to o-nitrophenol due to very low barrier (nearly barrier less). This is reason that hydrogen transfer occurs on $S_0$ state not very often. The newly found intersystem crossing $S_0T_1$-ICX has its geometry close to $S_0$-NP, so that 133 out of 280 go via it and those trajectories relax to ground state by non-hydrogen transfer pathway. In brief summary, we conclude that $S_0T_1$-ICX serves as the center player of intersystem crossing for three competing processes, 45 trajectories results in direct hydrogen transfer, 107 in tunneling hydrogen transfer, and 128 in non-hydrogen transfer.

Although every trajectory has its own lifetime when surface hopping occurs after photoexcitation, and the final result is estimated by the average assemble of all sampling trajectories. Figure 4 shows average population decay with respect of time from excited states $S_1$, $T_2$, and $T_1$, respectively. A single exponential function curve fitting is applied to calculate average lifetimes on those excited states. The lifetime of $S_1$ state was estimated to be about 8 fs as an ultrafast decay process and population is almost diminished after 25 fs. This is understandable in common sense that the first excited state decay more likely undergoes via intersystem crossing rather than vibrational cooling. The decaying process is also an ultrafast on $T_1$ state and the lifetime of $T_1$ state is estimated to be 14 fs. The population on $T_2$ state is quickly raised first and it reaches maximum at 10 fs and then it almost diminishes after 40 fs. The most of population via $T_1$ state transfers very quickly to $T_2$ state via conical intersection between $T_1$ and $T_2$ states. The population on $T_2$ state reaches its maximum (~93%) after 40 fs and the lifetime of $T_2$ state is estimated to be around 1000 fs. This is the most likely to contribute to the long-lived excited species that...
should be observed in both gas and liquid phase because the population on T1 state decays very slowly. However, the real lifetimes on excited states from experimental observation might be longer than the present estimation due to some extra kinetic energy being initially put in four vibrational modes which are enhancing hydrogen transfer dynamics.

The triplet T1 state plays an essential role for intersystem crossing-branched ESIPT photoisomerization dynamics in which 97% sampling trajectories bifurcate into three major distinct decay channels. The first branch is the non-hydrogen transfer that counts for 47% sampling trajectories, and timescale is about 100 fs to 500 fs (average is about 300 fs). For trajectories running on T1 state, once they form out-of-plane synchronous motion by O11H12 vibrating around C5O11 bond and the nitro group vibrating around C4N13 bond simultaneously, trajectories can reach S0T1-ICX to decay to ground S0 state. The present simulation shows if this out-of-plane synchronous motion form coherent motion before trajectories reach S0T1-ICX (we checked those trajectories from 500 fs to 1000 fs), they stay on T1 state as resonance trajectories. This is the second branch as the tunneling hydrogen transfer that counts for 36% sampling trajectories. We have roughly estimated thermal and microcanonical rate constants of nonadiabatic chemical reaction along ESIPT coordinate as shown in Fig. 2 49,

$$k = \frac{1}{2\pi Z_r} \int \exp(-\beta E) P(E) dE$$

(6)

in which $\beta$ is thermal energy (temperature is set up T = 300 K), $Z_r$ is reactant partition function and $P(E)$ is nonadiabatic transmission probability for the tunneling trajectories. The average tunneling timescale estimated from equation 6 is about 10 ps. This tunneling timescale is about the same as tunneling proton transfer reaction for tropolone molecule60. The third branch is the direct hydrogen transfer that counts for 13% sampling trajectories, and timescale is from 20 fs to 100 fs (average is about 40 fs). In this case, trajectories can go the direct hydrogen transfer before the out-of plane synchronous motion occurs. Then, the trajectories continue to run on the T1 state until they reach S0T1-IC1 accompanying with O14N13O15H12 group out-of-plane rotating and finally hop to S0 state.

**Typical trajectories.** Figure 5 shows the non-hydrogen transfer for a trajectory that hops to the S0 state via S0T1-ICX at 478.2 fs (this is trajectory staying the longest time on T1 state among all non-hydrogen transfer sampling trajectories). Starting from Frank-Condon region on S1 state, the trajectory experiences the following consecutive processes, it hops from the S1 to T2 state via S1T2-IC1 at the 4.8 fs and from the T2 to T1 state via conical intersection at the 10.1 fs. During the evolution period of time, O11H12 and O15H12 have small and large vibrations, respectively as shown in the second panel of Fig. 5. Two dihedral angles oscillate smoothly while four bond angles oscillate drastically as shown in the third and fourth panels of Fig. 5, respectively. This trajectory propagates a long time on the T1 state accompanying with the twists around the C6C7O11H12 and C6C7C8O15 dihedral angles. At 478.2 fs, it hops to ground S0 state. In the overall process, the out-of-plane synchronous motion by O11H12 vibrating around C6O11 bond and the nitro group vibrating around C6N13 bond and this reduces the possibility of hydrogen migration. We show the tunneling hydrogen transfer on T1 state for a typical trajectory in Fig. S5 of Supplementary Note 3. This case is similar to non-hydrogen transfer as shown in Fig. 5 except that O11H12 and O15H12 bonds oscillate coherently which differs from the non-hydrogen transfer case.

Figure 6 shows the direct hydrogen transfer on T1 state for a trajectory that takes the aci-nitrophenol isomerization in the ground state finally. This trajectory hops from the S1 to T2 state via S1T2-IC1 at the 16.6 fs and from the T2 to T1 state via conical intersection at the 18.2 fs. Sequentially, the hydrogen migration is occurred around
Figure 5. Typical trajectory. The first panel is for potential energy profiles along non-hydrogen transfer channel against the time. The second pane is evolution for bond lengths O$_1$H$_{12}$ and O$_{15}$H$_{12}$, the third is for dihedral angles C$_4$C$_5$O$_{11}$H$_{12}$ and C$_5$C$_4$N$_{13}$O$_{15}$ and the fourth is for bond angles C$_4$C$_5$O$_{11}$, C$_5$O$_{11}$H$_{12}$, C$_5$C$_4$N$_{13}$, and C$_4$N$_{13}$O$_{15}$. 
36 fs on the $T_1$ state accompanied by a shortening of the $O_1H_2$ bond and an elongation of the $O_2H_3$ bond. After 135.0 fs, this trajectory hops from the $T_1$ to $S_0$ state via $S_0T_1\text{IC}_1$. In the initial stage of evolution on the $S_0$ state,
this trajectory seems attempted back hydrogen transfer, but the reverse process is not successfully taken place. The aci-nitrophenol isomerization reaction appears in the rest of evolution time. The dihedral angle of O_{14}N_{13}O_{15}H_{12}

Figure 7. The same as Fig. 5 except for the direct hydrogen transfer trajectory on T_{1} state and back hydrogen transfer on S_{0} state.
rotates slightly around the C4N13 bond and the transferred H12 turns to away from the original O11, finally the aci-isomer2 is observed in this trajectory.

Figure 7 shows that the trajectory makes the hydrogen transfer on the T1 and back hydrogen transfer on S0 state. Starting from Frank-Condon region on S1 state, the trajectory hops from the S1 to T1 state via S1T1-IC1 at the 7.5 fs and from the T1 to T1 state via conical intersection at the 9.1 fs. Then, the hydrogen transfer is complete and the aci-nitrophenol species is formed at 35 fs on T1 state. During trajectory evolution, the distance between O15 and H12 is shortened gradually until bond forming, and bond angles of C5O11H12 and C5C4N13 change drastically while dihedral angles of C4C5O11H12 and C5C4N13O15 fluctuate smoothly. At 86.1 fs, trajectory decays to S0 state via S0T1-IC1. Soon after 130.0 fs, the back hydrogen transfer takes place on S0 state and regenerates the o-nitrophenol. In the rest of evolution, the trajectory shows a periodic fluctuation with H12 vibrates around the O11H12 bond. We have plotted more cases in Supplementary Note 3 for various more complicated fast and slow hydrogen transfer processes.

The present trajectory simulation indicates that the aci-nitro isomers are mostly generated on the triplet states, especially on T1 state and this confirms the previous assumption in which the HONO split-off motion is taken place in the triplet manifold. The present simulation also indicates that the o-NP dynamic decay via ISC ESIPT is mostly in time scale of subpicosecond (expect tunneling ESIPT) and this is in close agreement with the observations for a large number of nitrated polycyclic aromatic compounds. The present simulation shows that there excites the wide Franck-Condon region with very small energy gap between the first excited singlet S1 and the second triplet T2 states, and this can facilitate the fast ISC radiationless process. Therefore, trajectories hop from S1 to T2 state along the (O)H-O(NO) barrierless stretching pathway and in the tunneling zone the ESIPT pathways can generally have lower barrier via triplet states than via S1 state. The present simulation shows that the (O)H-O(NO) stretch related vibrational modes can enhance ESIPT and the torsion motion of nitro group would hinder the hydrogen transfer reaction.

Conclusion

By applying ab initio on-the-fly nonadiabatic molecular dynamic simulation for intersystem crossing-branched ESIPT for o-nitrophenol with use of global switching trajectory surface hopping algorithm, we have estimated reaction branch ratios and timescales of various ESIPT processes. As is summarized in Fig. 8, there are three decay branches, the first one is non-hydrogen transfer via newly found S0T1-ICX with ratio 47.5% and average timescale 300 fs, the second branch is tunneling hydrogen transfer with ratios 36.08% and 2.14% taken place on T1 and S1 states, respectively and timescale 10 ps, and the third branch is direct hydrogen transfer with ratios 13.21% and 1.07% taken place on T1 and T2 states, respectively and timescale 40 fs. However, the real lifetimes on excited states from experimental observation might be longer than the present simulated timescales due to some initial extra kinetic energy maybe enhance hydrogen transfer dynamics. Finally, we believe that the present trajectory-based on-the-fly nonadiabatic molecular dynamic simulation method can be generally applied to ultrafast photophysical and photochemical processes in a unified way involved in conical intersections and intersystem crossings for large-scale simulation.

Figure 8. One-slide summary of simulated branch ratios and its time scales for intersystem crossing-branched ESIPT reactions of o-nitrophenol.
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Author Contributions
C.X., L.Y. and C.Z. designed the simulations and the framework for trajectory surface hopping involving in both internal conversions and intersystem crossings. C.X. and L.Y. made the code for dynamic part and C.X. performed the simulations. C.X., L.Y., C.Z., J.Y. and Z.C. interpreted the data and analyzed excited-state intramolecular proton transfer for o-Nitrophenol. C.X. and C.Z. wrote the paper.

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