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Revealing the role of excited state proton transfer (ESPT) in excited state hydrogen transfer (ESHT): systematic study in phenol-(NH$_3$)$_n$ clusters

A general model of excited state hydrogen transfer (ESHT) which unifies the excited state proton transfer (ESPT) and the ESHT mechanism is presented. It reveals the hidden role of ESPT in controlling the reaction rate of the ESHT. For this purpose, experimental and theoretical works on the excited state dynamics of phenol-(NH$_3$)$_n$ clusters and related molecular systems over the last 30 years are overviewed. The systematic change from ESHT to decoupled electron/proton transfer according to the number of solvent molecules in the clusters is rationalized by the general model of ESHT including the role of ESPT.
Revealing the role of excited state proton transfer (ESPT) in excited state hydrogen transfer (ESHT): systematic study in phenol–(NH$_3$)$_n$ clusters

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Excited State Hydrogen Transfer (ESHT), proposed at the end of the 20th century by the corresponding authors, has been observed in many neutral or protonated molecules and become a new paradigm to understand excited state dynamics/photochemistry of aromatic molecules. For example, a significant number of photoinduced proton-transfer reactions from X–H bonds have been re-defined as ESHT, including those of phenol, indole, tryptophan, aromatic amino acid cations and so on. Photo-protection mechanisms of biomolecules, such as isolated nucleic acids of DNA, are also discussed in terms of ESHT. Therefore, a systematic and up-to-date description of ESHT mechanism is important for researchers in chemistry, biology and related fields. In this review, we will present a general model of ESHT which unifies the excited state proton transfer (ESPT) and the ESHT mechanisms and reveals the hidden role of ESPT in controlling the reaction rate of ESHT. For this purpose, we give an overview of experimental and theoretical work on the excited state dynamics of phenol–(NH$_3$)$_n$ clusters and related molecular systems. The dynamics has a significant dependence on the number of solvent molecules in the molecular cluster. Three-color picosecond time-resolved IR/near IR spectroscopy has revealed that ESHT becomes an electron transfer followed by a proton transfer in highly solvated clusters. The systematic change from ESHT to decoupled electron/proton transfer according to the number of solvent molecules is rationalized by a general model of ESHT including the role of ESPT.

During my scientific career at CNRS, I worked on the photochemistry of molecules and clusters. Starting at the University Paris-Iud I developed a method to catch the intermediate state of a bimolecular photochemical reaction by freezing the collision pair in a van der Waals complex. Using molecular cluster I evidenced the role of the hydrogen loss reaction in the excited state of aromatic molecules which lead to hydrogen transfer. This hydrogen dynamics is now recognized as responsible for a lot of non-radiative processes in aromatic subsisted molecules, including protonated ones that I study now at the Aix-Marseille University. 

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1. Introduction

Excited state proton transfer (ESPT) and excited state hydrogen transfer (ESHT) play important roles in photochemistry and biology. These photochemical reactions have been explored for 30 years via the spectroscopy of molecular clusters. The initial idea was that aromatic alcohols are more acidic in the excited state than in the ground state and therefore, in a molecular cluster, it should be possible to induce a proton transfer reaction by optical excitation, i.e. ESPT. However, following developments in experimental techniques and progress in computational chemistry, controversial results were reported, particularly in phenol (PhOH) hydrogen-bonded (H-bonded) clusters, and finally ESHT took over ESPT in the description of many photochemical processes. This research topic – started in around 1985 – has since provided many surprises, with the results having had a significant impact on our understanding of the photochemistry and photophysics of substituted aromatic molecules, including biomolecules.

Due to the importance of this topic, as will be shown below, a substantial number of papers on ESHT and ESPT have been published. Despite all that has been published on this subject over a period of around forty years, it seems that only the most recent studies have given rise to an accurate understanding of ESHT. It should also be noted that the understanding of the experimental results is now facilitated by ab initio calculations of excited states, a technique that has made considerable progress in recent years. Theoretical calculations well reproduce not only the energy of the electronic transitions but also the intensities of the active vibrational modes in spectroscopy. From the theoretical calculations and advanced experimental results, we can understand the relation between ESHT and ESPT, which has not been previously discussed in a systematic manner. In this review, the role of the ESPT underlying the ESHT process will be discussed using the benchmark molecular system PhOH–(NH₃)ₙ (n = 1–5). The review will begin by giving an overview of progress in this research field from the initial idea of ESPT to the establishment of ESHT.

2. Research history from ESPT to ESHT

2.1 Proton transfer in the condensed phase

Since the pioneering work of Förster,¹ the protolytic processes (displacement of the acid–base equilibrium or proton transfer in hydrogen-bonded complexes) induced by optical electronic excitation have been studied in the condensed phase.² Emission from neutral photoexcited 1-naphthol (1-NpOH⁺) is shifted from its gas phase value at 350 nm in the near ultraviolet (UV) to the visible (blue) in an aqueous solution. The blue emission after UV excitation was initially interpreted as the formation of the naphtholate anion³ and accompanying proton release from the excited state, i.e. ESPT. This implies a dramatic decrease in pKₐ (9.1 (S₀) to 0.5 (S₁)) upon photoexcitation to the first excited state of 1-NpOH. The decrease of pKₐ was also found in aromatic hydroxy, amino, and carboxy derivatives, and thus ESPT was established in the condensed phase.

2.2 Gas phase PT

Since the 1980s, molecular beam laser spectroscopy has been applied to aromatic molecular clusters and ESPT has been studied in both the frequency and time domains. A combination of several factors explains the success of this technique. Collisional relaxation can be neglected in a molecular beam. The number of molecules in the cluster is clearly specified when coupling with mass-selected multiphoton ionization detection. The relative orientations and geometries of molecules in the cluster are well defined because of the low temperature. Furthermore, an excited state reaction implies that the optical excitation is triggering the reaction, thus the reaction dynamics can be followed in real time, from a well-defined t = 0 of the reaction. Due to these advantages, the ESPT mechanism can be revealed at the molecular level by using molecular beam laser spectroscopy.

2.3 ESPT in naphthol/phenol–ammonia clusters and the contradictions

Leutwyler and coworker⁴ provided the first example of ESPT in the gas phase by the study of the fluorescence spectra of AH–B clusters [AH = 1-NpOH, B = (NH₃)ₙ, (piperidine)ₙ]. A red-shifted emission assigned to the proton-transfer process AH⁺–B → A⁺(–HB) was observed only from the excited state of sufficiently large clusters (n ≥ 4 or 2 for ammonia or piperidine, respectively). The observation of a threshold number of solvent molecules for an intracluster ESPT attracted the research groups specializing in time-resolved spectroscopy combined with the mass-selected multiphoton ionization technique. The dynamics in S₁ were measured by picosecond (ps) pump–probe spectroscopy of NpOH–(NH₃)ₙ. However, not only n = 4 but also n = 3 were reported to be the threshold size of ESPT in NpOH–(NH₃)ₙ.⁵,⁶ High-resolution time-of-flight mass spectroscopy revealed the influence of NH₃ evaporation after ionization in

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these experiments, and eventually it was concluded that the threshold size was \( n = 5 \). This threshold size was also supported by recent theoretical calculations.

It was expected that solvated clusters of PhOH and NpOH show similar behavior as far as ESPT is concerned. For PhOH solvated clusters, ESPT was postulated from the lowering of the ionization potential which was measured by the two-color resonance-enhanced multiphoton ionization (REMPI) technique. Since a strong H-bond lowers the ionization potential of the proton donor (acid), one can expect a large shift of the ionization thresholds when the ion pair is formed in the clusters. As expected, a strong lowering of the IP was found in PhOH–(NH\(_3\))\(_n\) at \( n = 4 \) and was interpreted as the ionization from the proton-transferred structure (PT structure).

There was, however, a notable difference between PhOH–(NH\(_3\))\(_n\) and NpOH–(NH\(_3\))\(_n\) clusters: free NH\(_4\)(NH\(_3\))\(_{n-1}\) clusters formed after photoexcitation were detected only for PhOH–(NH\(_3\))\(_n\) clusters and not for NpOH–(NH\(_3\))\(_n\) clusters. Moreover, a ps rise time of the NH\(_4\)(NH\(_3\))\(_{n-1}\) signal was similar to the decay signal of the parent PhOH–(NH\(_3\))\(_n\) cluster. The formation of NH\(_4\)(NH\(_3\))\(_{n-1}\) clusters after the ionization of the PhOH–(NH\(_3\))\(_n\) cluster was observed, and the energy dependence of the ps dynamics was probed. The observation of NH\(_4\)(NH\(_3\))\(_{n-1}\) implies the presence of a high barrier in the ionic state at a size where the barrier is very small in the excited state. However, it is well established that the acidity of ionic species is larger than that of the neutral species, hence the barrier should be lower than that in S\(_1\). In addition, Schmitt \textit{et al.} found that the action spectra of PhOH–(NH\(_3\))\(_n\) clusters obtained by monitoring NH\(_4\)(NH\(_3\))\(_{n-1}\) show well-resolved vibronic structures in S\(_1\). This posed a problem, as no ESPT is expected for small cluster sizes \((n \leq 3)\), whereas the appearance of protonated fragments was supposed to be a signature of the proton transfer mechanism either in the excited or in the ionic state. These contradictions suggest the presence of dynamics other than ESPT. A full discussion of these contradictions can be found in two reviews.\(^{15,16}\)

### 2.4 Discovery of ESHT

These contradictions were resolved upon the discovery of ESHT at the end of the 20\(^{1}\) century. The first key experiment was the observation of the NH\(_4\)(NH\(_3\))\(_n\) hypervalent cluster produced by excitation of the S\(_1\) state of PhOH–(NH\(_3\))\(_n\).\(^{16,17}\) In 1999, Pino \textit{et al.} observed the formation of a long-lived species from the S\(_1\) excited state by a delayed ionization of PhOH–(NH\(_3\))\(_n\) clusters.\(^{16}\) Serendipitously, they measured the pump–probe photoionization mass spectra not only at a short delay (\(\Delta t = 0\)) but also at very long delay (200 nanosecond (ns)), which is obviously longer than the lifetime of the parent cluster in S\(_1\) (see Fig. 1). When they measured the mass spectra without delay \(\Delta t = 0\), both parent ions PhOH–(NH\(_3\))\(_n\) and fragment ions NH\(_4\)(NH\(_3\))\(_{n-1}\) were detected (see Fig. 1). The typical lifetime of hydrogen-bonded PhOH clusters is \(\sim 10\) ns therefore all the signals should be very weak with a delay of 200 ns. Indeed, the mass peaks of parent ions PhOH\(^+\)(–NH\(_3\))\(_n\) become significantly weaker. However, even with such a long delay, the mass peaks of the fragment ions remain. This indicates that long-lived neutral species are generated from the photoexcited PhOH–(NH\(_3\))\(_n\), and photoionization of these species produces NH\(_4\)(NH\(_3\))\(_{n-1}\). From the various possibilities of long-lived species (including a triplet state of the clusters), they concluded that these neutral species are hydrogenated ammonia clusters generated by neutral cleaving of the OH bond, followed by a successive H atom transfer to the NH\(_3\) moiety: PhOH–(NH\(_3\))\(_n\) \(\rightarrow\) PhOH + 'NH\(_4\)(NH\(_3\))\(_{n-1}\). This result was soon confirmed\(^{18}\) by studying the electronic spectra of the reaction products using UV-near infrared (NIR)/IR-UV three-color ion dip spectroscopy. The NIR transition is the signature of a surplus electron in 'NH\(_4\)(NH\(_3\))\(_{n-1}\), which occupies the 3s orbital and gives rise to Rydberg transitions, such as 3p–3s in the NIR region. Fig. 2 displays the NIR spectra obtained by UV-NIR-UV ion dip spectroscopy of PhOH–(NH\(_3\))\(_n\) \((n = 2–5)\). Regardless of the number of ammonia molecules in the cluster, broad NIR transitions are observed. The spectra are very similar to the NIR spectra of NH\(_4\)(NH\(_3\))\(_{n-1}\) generated by photodissociation of pure NH\(_3\) clusters.\(^{19,20}\) which are traced in the same figure in dotted lines. This strongly supports the formation of 'NH\(_4\)(NH\(_3\))\(_{n-1}\) and neutral cleaving of the OH bond after photodissociation, \(i.e.\) ESHT. The infrared (IR) spectra measured by UV-IR-UV spectroscopy also confirmed the formation of 'NH\(_4\)(NH\(_3\))\(_{n-1}\).\(^{21}\)

### 2.5 The S.D.D.J. model

Theoretical approaches also support the existence of ESHT. \textit{Ab initio} calculations were performed on PhOH–(H\(_2\)O)\(_n\)\(^{22,23}\) with a diffuse function on the H atom in order to properly describe...
the Rydberg character of the $^1\pi\sigma^*$ state (see Fig. 3 for the orbital shapes). The calculations revealed a new predissociation pathway from the $S_1$ ($\pi\pi^*$) state to a low-lying $^1\pi\sigma^*$ state through a conical intersection, which leads to a concerted electron- and proton-transfer reaction from the chromophore to the solvent. Theoretical calculations also show that the excited-state reaction is endothermic in PhOH–(H₂O) complex but exothermic in the PhOH–NH₃ complex. Thus, ESHT in PhOH–(NH₃)ₙ clusters is also validated by the theoretical results. Combining experimental and theoretical results, a general model – the SDDJ model²³ – was proposed in 2002. In this model, a key role is played by excited singlet states of $\pi\sigma^*$ character, which have repulsive potential-energy functions with respect to the stretching of OH or NH bonds (see Fig. 4). The $^1\pi\sigma^*$ potential-energy functions intersect not only the bound potential-energy functions of the $^1\pi\pi^*$ excited states, but also that of the electronic ground state. The $S_1$ surface typically exhibits a local minimum of $^1\pi\pi^*$ character in the vicinity of the equilibrium geometry of the ground state. The second excited state is a dissociative $^1\pi\sigma^*$ which induces a barrier on the $S_1$ surface by its conical intersection with the $^1\pi\pi^*$ potential surface. The excited state lifetime is governed by the rate of tunnelling through the barrier from the $\pi\pi^*$ to the $\pi\sigma^*$.²³–²⁵ The barrier is also responsible for pronounced isotope effects on the fluorescence lifetime and quantum yield.²⁶ The $\pi\sigma^*$ state crosses the ground state at a greater distance, leading to the second conical intersection. Classically, at this point, the H atom can pursue the $\pi\sigma^*$ potential curve and will retain quite a high kinetic energy or be trapped on the ground state surface leading to a hot molecule which undergoes statistical evaporation of the H atom. From the theoretical point of view, the potential energy surfaces calculated by different methods are quite similar and vary only in details.²⁷–³¹

Fig. 2  Electronic spectra of the reaction products from photo-excited PhOH–(NH₃)ₙ are shown in color.¹⁸ Electronic spectra of the reaction products from photo-excited pure (NH₃)ₙ clusters assigned to the excitation of the $^1\text{NH}_2$(NH₃)ₙ radical cluster are shown in black.¹⁹,²⁰ Reproduced from ref. 18 with the permission of AIP Publishing, copyright 2002 and from ref. 20 with permission of Springer Nature, copyright 1999.

2.6  H-loss in the free PhOH molecule and its relation to ESHT

ESHT was established through the experimental and theoretical studies performed on solvated clusters. In monomers, the ESHT of clusters turns into an H loss channel from the excited state. H atom dissociation in photoexcited PhOH has been well

Fig. 3  Orbitals involved in the PhOH–NH₃ system. In the ground state, two electrons are localized in the $\pi$ orbital. In the first excited state, the $\pi\pi^*$ state, one electron is promoted into the $\pi^*$ orbital. In the second excited state, the $\pi\sigma^*$ state, one electron is localized in the very diffuse $\sigma^*$ orbital on the NH₃ moiety.

Fig. 4  Potential energy profiles of the lowest $^1\pi\pi^*$ states (squares), the lowest $^1\pi\sigma^*$ state (triangles) and the electronic ground state (circles) as a function of the OH stretch (PhOH) or NH stretch (indole, pyrrole) reaction coordinate. Geometries have been optimized in the excited electronic states at the CASSCF level; the PE profiles have been obtained with the CASPT2 method. Reproduced from ref. 23 with permission from the PCCP owner Societies, copyright 2002.
characterized by the measurement of the kinetic energy of the H atom. A bimodal distribution is observed. For the first component, the fast H atoms, the kinetic energy reflects the vibrational progression in the remaining PhO along the out of plane vibration. Here, out of plane vibrations are necessary in order to couple the \( \pi\pi^* \) and the \( \pi\sigma^* \) states because of the difference in \( C_s \) symmetry.\(^{29,32-34}\) Such a well-defined and structured kinetic energy distribution is characteristic of bond rupture on a dissociative potential. The H-loss channel is also evidenced by the formation of the PhO radical, as detected by VUV photoionization.\(^{35,36}\) The second component of the distribution, the slow H atoms, is assigned to a statistical fragmentation which occurs after internal conversion following the crossing of the \( \pi\sigma^* \) and the ground state. By measuring both the ps transients of parent and fragment and the total translational energy distribution of products as a function of the reaction time, it was evidenced that both the slow and fast components of the H fragment have the same tunneling origin and that the ground state is populated at the second \( \pi\sigma^*-\pi\sigma^* \) state and not through a direct \( \pi\pi^*-\pi\sigma^* \) internal conversion process.\(^{34,37}\)

The H-loss dynamics have also been studied in the time domain. The ps time-evolution of the \( S_1 \) excited state of PhOH shows that the lifetime decreases from 2 ns at the band origin \( \theta_0^* \) down to 600 ps at the band located 3500 cm\(^{-1}\) above the origin.\(^{38}\) These surveys show that the excitation of vibrational levels of A\(^*\) symmetry shortens the lifetime, mainly through \( S_1-\pi\pi^* \) vibronic coupling. At high energy, the direct excitation of the \( \pi\pi^* \) state leads to very fast dynamics of about 150 fs at 207 nm (excess energy of 12 000 cm\(^{-1}\)).\(^{39,40}\) The dynamics of the PhOH monomer has become a good example to test different theoretical methods for excited state dynamics, such as the quantum wave packet propagation method\(^{41}\) or the multistate semiclassical trajectory method. Xu\(^{42}\) et al. clearly confirm that hydrogen tunneling from the \( \pi\pi^* \) to the \( \pi\sigma^* \) potential surface is the main reaction mechanism in the PhOH H-loss photodissociation at the \( \theta_0^* \) band origin, and agree with experiments on the bimodal nature of the kinetic energy spectral distribution. They also show that the low kinetic energy release is due to both statistical dissociations to the ground state and direct dissociation to the first excited state of the phenoxyl radical. This interpretation is consistent with the translational energy distributions and branching ratios of the phenoxyl radicals produced in the different electronic states.\(^{38}\)

Besides \( \pi\pi^* \) to \( \pi\sigma^* \) internal conversion, another mechanism to explain H loss was proposed for the PhOH monomer.\(^{35}\) In this model, the first step was \( S_1 \) to \( S_0 \) internal conversion and the H loss was assumed to take place on the ground state surface. This model could not account for experimental observations such as the reaction in the ammonia cluster or the decrease of the reaction time upon the intermolecular OH–N vibration excitation and increase upon deuteration.\(^{35}\) This model was ultimately dismissed following the demonstration of the correlation of the excited state lifetime of PhOH and its derivatives with the \( \pi\pi^*/\pi\sigma^* \) gap.\(^{34,35,37}\) It should be stressed that the \( \text{NH}_4[\text{NH}_3]_n \) clusters can be seen as a very good messenger for characterizing the H loss on a \( \pi\sigma^* \) potential. Indeed, the H transfer requires a fairly direct process (the dynamics occur on a repulsive surface) since a process going through hot ground state molecules (H loss mediated by internal conversion to the ground state\(^{20}\)) would lead to a statistical process (Intramolecular Vibrational Redistribution: IVR) and the immediate evaporation of the whole cluster or some of the NH\(_3\) molecules. A strong driving force is imperative in order to attach the H atom and repel the \( \text{NH}_4[\text{NH}_3]_n \) cluster from the chromophore, and this cannot be achieved in a statistical process. One can also suppose that the kinetic energy of the \( \text{NH}_4[\text{NH}_3]_n \) clusters follows an impulsive type distribution, but the confirmatory experiment has yet to be performed. As we will show below, the variation of the dynamics with cluster size can be very well understood within this \( \pi\pi^*/\pi\sigma^* \) internal conversion framework.

### 2.7 Forgotten ESPT behind ESHT

Through extensive experimental and theoretical research, ESHT (or H loss in a monomer) has been characterized in terms of tunneling or internal conversion from the optically bright \( S_1 \) \( \pi\pi^* \) state to the dissociative \( \pi\sigma^* \) potential surface. In solvated clusters, the dissociative \( \pi\sigma^* \) potential will bring the H atom to the solvent moiety, the process terminating in H atom transfer. For monomers, the \( \pi\sigma^* \) state crosses the ground state potential surface, and internal conversion to the ground state is induced. This is the quenching process of photoexcited aromatic molecules, and is discussed as the built-in mechanism of aromatic biomolecules which have to remain photostable upon UV excitation. The first publications concerning ESHT described this reaction as the “Forgotten channel”\(^{16,17}\) but now it is a popularly invoked photochemical process, particularly in gas phase studies. On the other hand, ESPT has become a somewhat “forgotten” process now, and the relation between ESPT and ESHT has yet not been discussed, although both processes should be considered in the description of aromatic molecular systems. We should discuss their relations in aromatic molecular systems: which of ESHT or ESPT occurs, why so, where is the key on the potential surface, and how do ESHT and ESPT influence each other. PhOH–(NH\(_3\))\(_n\) clusters, at the origin of the ESHT issue, are one of the best systems with which to unravel these aspects, because the \( \pi\pi^* \) and \( \pi\sigma^* \) states are systematically controlled by the number of NH\(_3\) molecules in the cluster, and the reaction products \( \text{NH}_4[\text{NH}_3]_{n-1} \) are well characterized by both experimental spectroscopy and theory. In particular, their low ionization potentials\(^{42}\) and characteristic near infrared electronic absorption\(^{28}\) are important for experiments. Here, we will present the photodynamics for individual clusters of increasing size and discuss pertinent aspects of the ESHT/ESPT issue accordingly.

### 3. Methods

#### 3.1 Experimental

Although the experimental setups used to study PhOH–(NH\(_3\))\(_n\) clusters have been slightly different depending upon the experimental groups, the typical conditions are always the same. Clusters are produced by expansion of a gas mixture of NH\(_3\)-seeded He/Ne which has been flowed over a room temperature reservoir containing PhOH. The backing pressure...
is typically 2 bars and the nozzle diameter is around 500 μm. The cluster size is varied by changing the backing pressure and the partial pressure of NH₃ in the mixture, with the highest pressures giving rise to the largest clusters. The clusters are then laser excited and ionized in a second chamber, between the extraction plates of a TOF mass spectrometer.

Picosecond time-resolved IR/NIR (ps-TRIR or ps TRNIR) spectroscopy of PhOH–(NH₃)₃ was performed by utilizing time-resolved ultraviolet (UV)-IR/NIR-UV ion dip spectroscopy. The principle of the spectroscopy and details of the spectrometer are described elsewhere. The clusters obtained in the supersonic expansion have been studied by different excitation schemes which are illustrated in Fig. 5. Briefly, PhOH–(NH₃)₃ was excited to the ππ* state by a ps UV pulse, vexc, and the ESHT reaction was triggered. The reaction product, NH₄(NH₃)⁻⁻, was ionized by a ns UV laser pulse, vion, which was fired at (typically) 30 ns after vexc. The population of the reaction product was monitored at masses of NH₄⁺(NH₃)⁻⁻ using a time-of-flight mass spectrometer. A ps tunable NIR laser, vNIR, whose frequency was fixed to that of the 3p–3s Rydberg transition of NH₄⁺(NH₃)⁻⁻ in the NIR region, was fired for Δτ ps after vexc. The population of NH₄⁺(NH₃)⁻⁻ was depleted by excitation to the 3p Rydberg state. If vNIR is sufficiently strong, population of NH₄⁺(NH₃)⁻⁻ is generated until the vNIR irradiation (at Δτ) is completely depleted. Then, the ion signal of NH₄⁺(NH₃)⁻⁻ corresponds to the amount of NH₄⁺(NH₃)⁻⁻ produced after Δτ ps until 30 ns after from the vexc irradiation. Therefore, the ESHT dynamics can be monitored by measuring the intensity of NH₄⁺(NH₃)⁻⁻ as a function of Δτ.

This spectroscopy has several variations. If vNIR is changed to a pulse from a tunable ps IR laser, vIR, the time-resolved vibrational (IR) spectra can be measured. In the time-domain, we can measure the femtosecond dynamics by replacing vexc and vNIR by femtosecond UV and NIR laser pulses. The static IR and NIR spectra can be measured by using ns tunable lasers for all three laser pulses. It should be noted that the femtosecond UV/NIR lasers do not have high enough energy resolution to distinguish individual vibrational transitions. Thus, femtosecond spectroscopy is used only in the measurement of time-resolved electronic spectra in the NIR region.

3.2 Theoretical

The first calculations on the PhOH–NH₃ system showing evidence of the role of the πσ* and of the H transfer process were performed by Sobolewski et al. but these phenomena were inferred long ago by Evleth and collaborators on the basis of semi empirical molecular-orbital calculations. Using multi configuration methods (CASSCF/CASPT2) the calculations performed on PhOH–solvent, where solvent is NH₃ or H₂O, showed that the reaction of excited PhOH-solvent was exothermic for NH₃ clusters and endothermic for H₂O clusters. The role of the πσ* state could only be evidenced by the addition of diffuse functions in the basis set, necessary to fully describe the Rydberg character of this state. It was also shown that there is no barrier to the proton transfer in the ionic state (PhOH⁺–NH₃), adding one more piece of information dismissing the ESPT

![Fig. 5](Image)

Fig. 5 Different experimental schemes for studying the reaction dynamics in PhOH–(NH₃)₃ clusters: (a) simple pump–probe experiment: this allows the study of the decay of the parent ion, (b) ionization after a long delay between the pump and the probe. The excited state parent ions have decayed and only the reaction product can be ionized, (c) three color experiment. A NIR laser between the first and second UV lasers excites the NH₄(NH₃)⁻⁻ product immediately after the H transfer and leads to the depopulation of the reaction product. Reproduced from ref. 54 with permission from the PCCP Owner Societies, copyright 2020.
mechanism discussed in an earlier paper.\textsuperscript{31} It should be noted that earlier calculations were focused on the ESPT process and did not mention the $\pi\pi^*$ state.\textsuperscript{46-47} For clusters, the ground state geometries and the ground state vibrations have been mostly calculated at the MP2 level, and these calculations were used to assign the vibrational bands in the IR hole burning experiments and determine the cluster structure.\textsuperscript{31,44} For the excited state, two main methods were used: either (CASSCF/CASPT2) or Coupled Cluster (CC2). Both methods give good agreement with experiments. With CC2, the absorption of the excited state can be calculated and the nature of the excited state ($\pi\pi^*$ or $\pi\sigma^*$) compared with the experimental results.\textsuperscript{24} For large clusters, DFT/M06-2X/cc-pVTZ has also been used, with these calculations predicting a stable ground state proton transfer structure (zwitterionic structure) for PhOH$–$(NH$_3$)$_n$ where $n \geq 8$.\textsuperscript{48}

It is interesting to note that most of the theoretical work is published alongside experiments. The experimental and theoretical work has made progress together, nicely illustrating the interest of studying clusters, for which both approaches are directly comparable.

### 4. Experimental findings

The ESHT mechanism has been observed to vary as a function of cluster size. As we will describe below, the study of each cluster size of PhOH$–$(NH$_3$)$_n$ reveals some new information on the reaction dynamics, from the $n = 1$ complex in which the role of the intermolecular vibration is evidenced, to the $n = 5$ cluster in which the role of the proton transfer state in the ESHT is demonstrated.

#### 4.1 PhOH$–$(NH$_3$)$_1$: lifetime variation with intermolecular vibration

For the PhOH$–$(NH$_3$)$_1$ complex (hereafter “the $n = 1$ complex”), the time-evolution of the H transfer reaction can only be measured with the pump–probe scheme monitoring the decay of the parent ion. Indeed, since the lifetime of the $^1$NH$_4$ product is very short (13 ps), i.e. a lot shorter than the tunneling time from the $\pi\pi^*$ to the $\pi\sigma^*$ potential surface, the concentration of the product is negligible. One should mention that this method (Fig. 5a) measures the lifetime of the parent ion after the initial excitation pulse, which corresponds to the time for the tunneling, and subsequent dissociation, to occur. The latter (dissociation) corresponds to the time necessary for the $^1$NH$_4$ moiety to be far enough from the PhO. Examples of such an effect are demonstrated in the case of the Na$^+$ diatomic molecule or in Na$–$(NH$_3$)$_n$ clusters.\textsuperscript{50}

The excited state lifetime of the $n = 1$ complex has been measured to be 1.2 ns (ref. 51) which is shorter than the lifetime of the free PhOH molecule (2 ns (ref. 52) or 2.2 ns (ref. 24)). The excitation of the intermolecular vibration at +182 cm$^{-1}$ above the origin leads to a shortening of the lifetime down to 391 ± 100 ps. This vibration is the intermolecular OH···N stretching mode which is directly connected to the reaction path. Indeed, one can expect that when the OH···N distance decreases, the tunneling barrier decreases, as observed. This observation is very convincing evidence that the reaction proceeds via a tunneling mechanism. A similar observation was also made for the PhOH$–$(NH$_3$)$_2$ complex.\textsuperscript{51}

Another strong argument for a tunneling process in the reaction was the observation of the change in the reaction speed upon deuteration. For the $n = 1$ deuterated complex, the lifetime is 7 ns (compared to 1.2 ns for the hydrogenated species), and for the $n = 2$ cluster, it is 7 ns instead of 400 ps. Thus, the simplest explanation for this observation is the presence of a tunneling barrier.

#### 4.2 PhOH$–$(NH$_3$)$_2$: biexponential lifetime and the role of the triplet state

The $S_1$ lifetime of PhOH$–$(NH$_3$)$_2$ was measured by two groups using a pump–probe ionization method.\textsuperscript{31,53} Both groups monitored the parent ion, PhOH$^–$(NH$_3$)$_2$, and measured the decay of $S_1$ by the loss of the parent ion (pump–probe method, see Fig. 5a). The reported lifetimes are significantly different: 80 ps (ref. 53) and 400 ps.\textsuperscript{51} The main reason for the difference is the simultaneous excitation of larger clusters. Even after careful preparation of the experimental conditions, it is very difficult to avoid the coexistence of clusters of various sizes. The transitions of the PhOH$–$(NH$_3$)$_n$ ($n = 2$–5) clusters appear in almost the same frequency range, thus, multiple species are invariably excited at the same time. Furthermore, larger clusters ($n > 2$) easily dissociate after ionization, and contaminate the PhOH$^–$(NH$_3$)$_2$ mass channel. The latter group subtracted the contributions from the larger clusters by measuring the time-evolution of the background signal, and obtained the longer lifetime. This lifetime is the convolution of the tunneling and the dissociation times. The pure H transfer lifetime (i.e. tunneling in this case) was measured by time-resolved UV-NIR-UV ion dip spectroscopy (Fig. 5c) which indicates the H transfer directly by NIR absorption. This method is insensitive to interference from larger clusters, thus the difference in the lifetimes corresponds to the dissociation lifetime.

All the time-evolutions of $^1$NH$_4$NH$_3$ formation detected by NIR absorption are biexponential.\textsuperscript{54} Here, the biexponential time evolution from the $S_1$ origin means that an electronic relaxation would be feasible to explain this behavior, while IVR makes no contribution (Fig. 6a). The single exponential fit to the early part of the time-evolution from the $S_1$ origin gives a lifetime of ~270 ps, thus the 130 ps difference from the lifetime measured by the conventional method\textsuperscript{51} is the contribution of $^1$NH$_4$NH$_3$ dissociation.

Two reaction paths and a contribution from the unknown state X in one of the paths are assumed to explain the biexponential time-evolution (see below).

$$\text{PhOH}^*–(\text{NH}_3)_2 \xrightarrow{k_1} \text{PhO}^* + \text{NH}_4\text{NH}_3$$

$X$
ESHT reaction

the common parameter of time scale from the (a) S1 origin (log scale on vertical axis) and (b) S1 vibronic levels. Here, we assumed that evolutions are \( \tau_1 = 268 \text{ ps} \) and biexponential linear scale on vertical axis) taken from ref. 54. A single exponential dynamics from all vibronic levels, and applied a global fitting with a biexponential growth, the observed time-evolution from the \( \tau_1 = 268 \text{ ps} \), \( \tau_2 = 440 \text{ ps} \), and \( \tau_3 = 440 \text{ ps} \) are shown by the solid straight line (a) and curve (b), respectively. Reproduced from ref. 54 with permission from the PCCP Owner Societies, copyright 2020.

The first reaction path with reaction rate \( k_1 \) is the simple ESHT reaction via the conical intersection from \( S_1 \pi \pi^* \) to \( \pi \sigma^* \). The second path including the state X also has an ESHT reaction for the decay of state X, but the formation of state X occurs via electronic relaxation. By solving the rate equations and fitting with a biexponential growth, the observed time-evolutions are fitted for the time-evolutions from several vibronic levels. Here, we assumed that \( k_2 \) is common for the dynamics from all vibronic levels, and applied a global fit. The example of the fitted results for the time-evolution from the overtone of the butterfly vibration in \( S_1 \) is shown in Fig. 6b along with the observed time-evolution on a linear vertical scale. The fitted values are \( \tau_1 = 1/k_1 = 268 \text{ ps} \) and \( \tau_3 = 1/k_3 = 440 \text{ ps} \), and the common parameter of \( \tau_2 = 1/k_2 \) converges at 125 ps. \( \tau_1 \) and \( \tau_3 \) vary according to the vibronic levels, but all of them are on the order of hundreds of ps.

What is state X? As mentioned above, IVR is not responsible for the biexponential behavior, thus a vibrational bath mode is not responsible. The high energy vibrational states of \( S_0 \) are also not a suitable candidate for X, because such states relax quickly by the fast IVR process, thus it is not possible to access the \( \pi \sigma^* \) state which promotes the ESHT reaction. A triplet state \( (3\pi \pi^*) \) of the PhOH–\( (NH_3)_2 \) cluster is also not reasonable, because \( k_2 \) must be the rate of the intersystem crossing, which would be much slower than ps.

The last candidate is a triplet \( \pi \sigma^* \) state \( (3\pi \sigma^*) \). The \( \sigma^* \) orbital is diffuse and located far away from the cluster (see Fig. 3). Thus, the \( 3\pi \sigma^* \) is expected to be energetically degenerate with the \( 1\pi \pi^* \). As a consequence, there is almost no energy gap in the intersystem crossing and thus the relaxation can be accelerated to the sub-ns order. The potential surface of \( 3\pi \sigma^* \) crosses that of \( 1\pi \pi^* \) at a longer OH distance, and a bound state can be produced because of the avoided crossing. Then, those molecules converted to \( 3\pi \sigma^* \) can be trapped in this bound state before dissociating to PhOH + \( \cdot NH_2NH_3 \) along the repulsive \( 3\pi \sigma^* \) potential surface (see Fig. 7).

This model was confirmed by the theoretical calculations of the potential curves of \( 1,3\pi \pi^* \) and \( 1,3\pi \sigma^* \) along the OH distance shown in Fig. 7. The potential curves were obtained by stretching the OH distance around the excited state equilibrium distance. The black curve represents the \( 1\pi \pi^* \) state and a local minimum at \( R_{OH} = 1.7 \text{ Å} \) corresponds to the proton transfer state. This PT state is not energetically accessible from the \( \pi \pi^* \) obtained by ground state excitation. The blue curve shows a triplet \( 3\pi \pi^* \) state that does not exhibit a local PT state minimum. The red and pink curves correspond to the singlet and triplet \( 1,3\pi \pi^* \) states, respectively. The potential curves of the singlet and triplet \( \pi \sigma^* \) states are almost degenerate in

The potential curves of the PhOH–\( (NH_3)_2 \) cluster along the OH bond length \( R_{OH} \) in different electronic states and spin multiplicities obtained at the RI-CC2/aug-cc-pVDZ level. \( C_2 \) symmetry has been imposed for the calculations, thus avoided crossing should be considered between \( 1,3\pi \pi^* \) and \( 1,3\pi \sigma^* \) states. Reproduced from ref. 54 with permission from the PCCP Owner Societies, copyright 2020.
energy. The dissociative potential curves of $^1\pi\pi^*$ cross those of both $^1\pi\pi^*$ and $^3\pi\pi^*$, thus confirming the mechanism detailed above: the biexponential time-evolutions are due to the triplet $\pi\sigma^*$ formation and successive trapping in the triplet manifold.

4.3 PhOH–(NH$_3$)$_3$: isomerization within the reaction products

The PhOH–(NH$_3$)$_3$ cluster discussed above has no isomers, and the reaction product of ESHT is a single species, 'NH$_4$(NH$_3$)$_3$, rendering the study of this species relatively simple. For clusters containing more than two ammonia molecules, the ESHT reaction becomes complicated in two ways. The first is the coexistence of several conformers of PhOH–(NH$_3$)$_n$. Excitation of different conformers leads to a difference in the initial geometry of the cluster undergoing the ESHT reaction. This complication can be minimized if the electronic transition of each conformer is well separated in energy and can be selectively excited. The second point is the conformational variety of the reaction product 'NH$_4$(NH$_3$)$_{n-1}$ ($n \geq 3$). Isomerization between these conformers is also possible, according to the excess energy available after ESHT. This second point is essential and cannot be avoided experimentally.

For PhOH–(NH$_3$)$_3$, hole burning spectroscopy indicates that this cluster exists in only a single geometry in the ground state.$^{26}$ Its cyclic hydrogen-bonded structure, shown in the lower left of Fig. 8, was determined by interpretation of the IR spectrum with the help of theoretical calculations.$^{21}$

Fig. 9a shows the NIR transition probing the electronic structure obtained by scanning the $\nu_{NIR}$ frequency in the UV-IR-UV pump–probe spectroscopy of PhOH–(NH$_3$)$_3$. The NIR absorptions at 7800 cm$^{-1}$ and 9000 cm$^{-1}$ are assigned to 3p–3s transitions of NH$_3$–NH$_4$–NH$_3$ while those at 11 000 cm$^{-1}$ and 12 000 cm$^{-1}$ originate from that of 'NH$_4$–NH$_3$–NH$_3$. The NIR absorption of 'NH$_4$–NH$_3$–NH$_3$ (Fig. 9c) rises faster than that of NH$_3$–NH$_4$–NH$_3$. Another important point is that both species exist even after a long delay time. This suggests an equilibrium between the two species.

The fast formation of 'NH$_4$–NH$_3$–NH$_3$ demonstrates the memory effect of the initial geometry in the parent PhOH–(NH$_3$)$_3$ cluster (see Fig. 8). After photoexcitation to $S_1$, O–H bond radical dissociation in the cyclic PhOH–(NH$_3$)$_3$ induces H atom transfer to the terminal NH$_3$. This reaction generates the initial product 'NH$_4$–NH$_3$–NH$_3$ exclusively. The 'NH$_4$–NH$_3$–NH$_3$ species is less stable and thus isomerizes to the more stable NH$_3$–NH$_4$–NH$_3$ (the barrier height is 3.1 kcal mol$^{-1}$ (ref. 43 and 63)). The back reaction to the initial product under an isolated condition, leads to an equilibrium between the two species at long delay times.
The three-state model is confirmed by the application of rate equations, using the rate constants \( k_1 \) (ESHT), \( k_2 \) (isomerization), and \( k_{-2} \) (back-reaction) shown in Fig. 8. The details of the fitting procedures are reported in the reference, thus we will show only the fitted results here. Fig. 10 shows the time-evolution of (a) the meta-stable product \(^1\text{NH}_4\text{NH}_2\text{NH}_3\) and (b) the final product \(\text{NH}_3\text{NH}_4\text{NH}_3\). The experimental time evolutions were obtained by the averaging of the time evolutions of both the NIR and IR spectra. The fitted curves obtained by the global fit of the rate equation analysis are shown by solid curves as well as the residual errors between the observations and the fit functions. The fits reasonably reproduce the time-evolution of the meta-stable and the final products with 24 ps, 6 ps, and 9 ps for the inverse of rate constants \(1/k_1\), \(1/k_2\), and \(1/k_{-2}\), respectively. Thus, we conclude that the initial formation of the metastable product is due to the memory effect, followed by isomerization to the most stable species, and final equilibrium is attained by the back reaction.

### 4.4 PhOH–(NH₃)_4: an even more complex system

While time-resolved spectroscopy has been applied to ESHT reactions in PhOH–(NH₃)₂,₃,₅ clusters, no time-resolved spectroscopy has yet been reported for the PhOH–(NH₃)₄ system. To complete the systematic study of PhOH–(NH₃)₄, the ps TRIR and TRNIR spectroscopy of PhOH–(NH₃)₄ has been performed for this review. Before presenting the time resolved data, we would like to summarize the previous nanosecond spectroscopy of PhOH–(NH₃)₄ associated with theoretical calculations.¹⁻³⁵

The UV action spectra of PhOH–(NH₃)₄ obtained by the monitoring by delayed ionization detection of the product with (a) ps and (b) ns lasers are shown in Fig. 11 (see Fig. 5b for the spectroscopic method). Four color hole burning spectroscopy shows that PhOH–(NH₃)₄ has three isomers, whose bands are indicated in the figure.³⁶ Due to the lower spectral resolution of the ps laser (15 cm⁻¹), a low frequency progression of isomer C cannot be resolved in Fig. 11a. Also, sharp and weak bands of isomers A and B cannot be seen in Fig. 11a. As a result, the ESHT time-evolution can be measured only for isomer C.

Theoretical calculations at the MP2/6-31++G(d, p) level show that a cyclic hydrogen bonded structure is the most stable for this cluster. In the early stages of the ESHT studies in PhOH–(NH₃)₄, the coexistence of isomers was not known. For this reason, the IR spectrum obtained by labeling the band at 35 348 cm⁻¹ (isomer A) was assigned to the cyclic hydrogen-bonded structure. However, this assignment should be reconsidered because the most stable species should appear more strongly than other isomers. In this case, isomer A is significantly weaker than isomer C. Also, no IR spectra of isomers B and C have been measured. Thus, we proceed in the analysis with a tentative assignment of the isomer C as the cyclic hydrogen-bonded structure.

The cyclic hydrogen-bonded structure shown in Fig. 12 is similar to that of PhOH–(NH₃)₃. Fig. 12 also shows possible conformations of the reaction products \(^1\text{NH}_4\text{NH}_3\text{NH}_3\) with their relative energies. If ESHT takes place in the parent cluster of the cyclic structure, the H atom will transfer to the terminal NH₃ in the ammonia moiety. In this case, only the meta-stable product \(^1\text{NH}_4\text{NH}_3\text{NH}_3\text{NH}_4\) (4c in Fig. 12) is produced. The NIR spectrum of reaction products from PhOH–(NH₃)₄ measured by ns laser systems is also shown in the figure. Theoretically predicted 3p–3s Rydberg transitions for the products 4a to 4d are also indicated in the spectrum. The spectrum clearly shows the coexistence of all four reaction products 4a to 4d. Therefore, it is reasonable to consider the isomerization of the meta-stable product 4c to the other products. However, it is too complicated to trace all the possible isomerization paths and equilibria after ESHT because of the multiple isomers of \(^1\text{NH}_4\text{NH}_3\text{NH}_3\). Thus, we neglected the contributions of species 4b and 4c, and applied the three-state model to the photoexcited PhOH–(NH₃)₄, the meta-stable product 4c and the most stable product for this cluster, 4d.

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Fig. 10 The time evolutions of the products (a) \(^1\text{NH}_4\text{NH}_2\text{NH}_3\) monitored at 12 000 cm⁻¹, (b) the final product \(\text{NH}_3\text{NH}_4\text{NH}_3\) monitored at 6000 cm⁻¹ for ESHT in the PhOH–(NH₃)₃ cluster. The fitted results are shown as solid curves along with the residual. Reproduced from ref. 43 with permission of AIP Publishing, copyright 2007.

![Image](image_url)
Time-evolutions of the meta-stable product 4c and the most stable product 4d were monitored by following the NIR absorption at 11 000 cm\(^{-1}\) and 7000 cm\(^{-1}\), respectively. The observed time-evolutions and the best fit of the three-state model are shown in Fig. 13. The absorption of the meta-stable product 4c rises quickly after the photoexcitation to S\(_1\) and decays gradually after 30 ps. On the other hand, the absorption of the most stable product 4d increases slowly. This is essentially the same mechanism as that observed in PhOH–(NH\(_3\))\(_3\): selective generation of the metastable 4c product due to the memory effect, and successive isomerization towards the most stable product, 4d. Good reproduction of the observed time-evolutions by the rate equations confirms the equivalent ESHT mechanism to PhOH–(NH\(_3\))\(_3\). The inverse of the rate constants 1/\(k_1\), 1/\(k_2\) and 1/\(k_3\) are determined to be 15 ps, 25 ps and 38 ps, respectively.

As compared to PhOH–(NH\(_3\))\(_n\), 1/\(k_1\), the time constant for the tunneling is two times faster (15 ps instead of 30 ps) whereas the isomerization constants (1/\(k_2\) and 1/\(k_3\)) are longer (25/38 ps instead of 6/9 ps for PhOH–(NH\(_3\))\(_3\)). The shorter tunneling time implies that the tunneling barrier for PhOH–(NH\(_3\))\(_n\) is smaller than that in PhOH–(NH\(_3\))\(_3\), in agreement with the interpretation given in Section 4.7 “General model for ESHT and ESPT”. Concerning the isomerization, the barriers are nearly the same for ‘NH\(_4\)(NH\(_3\))\(_2\) and ‘NH\(_4\)(NH\(_3\))\(_3\) so the slower reaction rate can be accounted for by the higher density of states in ‘NH\(_4\)(NH\(_3\))\(_3\) compared to NH\(_4\)(NH\(_3\))\(_2\).

### 4.5 PhOH–(NH\(_3\))\(_n\): the return of proton transfer

S\(_1\)–S\(_0\) electronic spectra of PhOH–(NH\(_3\))\(_n\) (n \(\leq\) 4) exhibit well-resolved structures. This is consistent with the picosecond order of the ESHT reaction rate. In contrast, PhOH–(NH\(_3\))\(_3\) gives rise to a broad absorption, which indicates a significant change of the reaction mechanism for this cluster. Here, we would like to summarize the structural information obtained from nanosecond “static” spectroscopy. The structure of the PhOH–(NH\(_3\))\(_5\) cluster in S\(_0\) has been assigned to a bicyclic hydrogen-bonded

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**Fig. 12** Reaction scheme for the PhOH–(NH\(_3\))\(_4\) cluster. – The initial cluster (upper left) reacts at the rate \(k_1\) to give the PhO\(^{+}\)–NH\(_4\)(NH\(_3\))\(_3\) species. The initial ESHT product starting from ‘NH\(_4\)(NH\(_3\))\(_3\) (4c) evolves to structures 4d and 4b and then isomerizes to the most stable C\(_{3v}\) isomer 4a (upper right). – Lower trace: electronic spectrum of the ‘NH\(_4\)(NH\(_3\))\(_3\) species, which includes the contributions of the four isomers. The colored sticks are the calculated excited states of the different isomers. Reproduced from ref. 18 and 63 with the permission of AIP Publishing. copyright 2002 and 2003, respectively.

**Fig. 13** Picosecond time-evolutions of reaction products of PhOH–(NH\(_3\))\(_5\) measured by the transitions in the IR (NH vibrations at 3180 cm\(^{-1}\) (red curve)) and NIR (electronic excitation). Blue curve – excitation at 11 000 cm\(^{-1}\) – probes the ‘NH\(_4\)(NH\(_3\))\(_3\) in the chain structure (4c). Green curve – excitation at 7000 cm\(^{-1}\) – probes the most stable C\(_{3v}\) structure (4a): this structure is obtained at longer delay times.
structure (see Fig. 14).\textsuperscript{21} Photoexcitation of the cluster triggers O–H cleaving by ESHT, with the H atom transferred to the nearest ammonia in the ammonia moiety. From this geometry, the expected nascent product is the structure where the fifth ammonia molecule is hydrogen bonded to one of the NH\textsubscript{3} in \textsuperscript{14}NH\textsubscript{4}(NH\textsubscript{3})\textsubscript{5} in C\textsubscript{3v} symmetry, which is shown in Fig. 14 as C\textsubscript{3v}+1. However, the NIR spectrum obtained by ns UV-NIR-UV pump-probe spectroscopy (Fig. 2 and lower right panel in Fig. 14) shows a strong and broad band at \( \sim 6000 \text{ cm}^{-1} \) which is assigned to \textsuperscript{14}NH\textsubscript{4}(NH\textsubscript{3})\textsubscript{5} in the most stable tetrahedral structure \( T_d \), not C\textsubscript{3v}+1. Therefore, several intermediate species are expected to play a role in the ESHT reaction of PhOH–(NH\textsubscript{3})\textsubscript{5}.

The time-evolutions of the NIR signals at 6000 cm\(^{-1}\), 7000 cm\(^{-1}\) and 8000 cm\(^{-1}\) are shown in Fig. 14.\textsuperscript{37,58} The signal at 6000 cm\(^{-1}\), which corresponds to the most stable \( T_d \) product, grows gradually and becomes constant at around 100 ps. The signal at 8000 cm\(^{-1}\) rises rapidly and decays slowly. The ultrafast rise of the signal is consistent with the broad spectral feature of the UV absorption transition of PhOH–(NH\textsubscript{3})\textsubscript{5}. The time-evolution at 7000 cm\(^{-1}\) shows a sharp rise and the signal is constant after the rise. This is explained by an overlapping of the decreasing signal at 8000 cm\(^{-1}\) and the increasing signal of the \( T_d \) species at 6000 cm\(^{-1}\). Quantum chemical calculations suggest two possible transient species in this region. One is the nascent ESHT product of C\textsubscript{3v}+1 symmetry, of which the \( 3p_{x,y,z} \rightarrow 3s \) transitions are shown by blue bars in Fig. 14a. Another is the 4p Rydberg-\( \pi^* \) transition of the CT complex originating from the occupation of an electron in the \( \pi^* \) orbital, of which the transition energy is 0.85 eV (6900 cm\(^{-1}\)). These two species are considered potential candidates for the 8000 cm\(^{-1}\) absorption. In both cases, this signal is a signature of electron transfer from the aromatic \( \pi^* \) to the \( \sigma^* \) orbitals.

The NIR transitions are a good indicator of electron transfer. On the other hand, the real N–H bond formation dynamics can be observed independently by IR spectroscopy in the 3 \( \mu \text{m} \) region. Fig. 15 shows the ps-TRIR spectra of PhOH–(NH\textsubscript{3})\textsubscript{5}. The theoretical IR spectra of both the C\textsubscript{3v}+1 and \( T_d \) products are also shown in Fig. 15b. Here, the \( T_d \) product will exhibit intense vibrational transitions at 2965 cm\(^{-1} \) (H-bonded NH stretching of \textsuperscript{14}NH\textsubscript{4}) and 3260 cm\(^{-1} \) (free NH stretching of NH\textsubscript{3}), while the C\textsubscript{3v}+1 species is calculated to give rise to IR absorption at 3217 cm\(^{-1} \) (free NH stretching of NH\textsubscript{3}). The ps-TRIR spectra show a strong transition at \( \sim 3200 \text{ cm}^{-1} \) and a broad absorption at \( \sim 3000 \text{ cm}^{-1} \). Time-evolutions at 2983 cm\(^{-1} \) (A in Fig. 15a), 3165 cm\(^{-1} \) (B) and 3248 cm\(^{-1} \) (C) are shown in Fig. 15c. Both signals at A and C gradually grow in intensity with increasing time.

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Fig. 14  H transfer dynamics of PhOH–(NH\textsubscript{3})\textsubscript{5}: (a) absorption spectrum of the \( T_d \) \textsuperscript{14}NH\textsubscript{4}(NH\textsubscript{3})\textsubscript{4} species, the most stable structure, (b) time evolutions of the electronic structures: at 8000 cm\(^{-1} \) excitation of the charge transfer state within PhOH–(NH\textsubscript{3})\textsubscript{5}, at 6000 cm\(^{-1} \) the final \( T_d \) structure, and at 7000 cm\(^{-1} \) the overlap of both contributions. The right panel shows a scheme of the reaction events and time evolution of the electronic structures. (c) shows the population evolution deduced from the fit. The C\textsubscript{3v}+1 structure is barely populated. Reproduced from ref. 58 with the permission of Wiley-VCH Verlag GmbH & Co, copyright 2018.
delay time, and no sharp rise was found. The signal at B should be sensitive to the presence of the nascent \( C_{3v}^{+1} \) product. However, the time-evolution does not show an ultrafast rise, and can be fitted by single exponential functions with lifetimes of \( \sim 20 \) ps. This suggests that the ps-TRIR spectra reflect mainly the vibrational transitions of the \( T_d \) product.

The right panel in Fig. 14 summarizes the ESHT reaction mechanism and spectroscopic signatures for the possible species involved. The time-evolutions are analyzed by rate equations with the rate constants \( k_{S_1\text{-CT}} \), \( k_{\text{CT-}C_{3v}^{+1}} \), and \( k_{\text{isomerization}} \). The global fit reproduced the observed time-evolutions, as shown by black solid curves in Fig. 14 and 15, with \( 1/k_{S_1\text{-CT}} \ll 3 \) ps, \( 1/k_{\text{CT-}C_{3v}^{+1}} = 20.2 \) ps, and \( 1/k_{\text{isomerization}} = 0.4 \) ps. The population changes of CT, \( C_{3v}^{+1} \) and \( T_d \) species extracted from the fit are also illustrated in Fig. 14c. The fit shows that the electron transfer from PhOH to the ammonia moiety is very fast (\( \ll 3 \) ps). However, the proton transfer is slower (20 ps) and is decoupled from the electron transfer. It is clearly a different mechanism compared to that observed in the smaller clusters (\( n \leq 4 \)). It should be noted that the time resolved experiment in the femtosecond regime later determined the time constant of the initial electron transfer as \( 370 \) fs.

The electron-proton decoupling for \( n = 5 \) is rationalized by the strong distortion of the potential surfaces of the \( S_1 \) and \( S_2 \) states. Fig. 16 shows the calculated potential energy curves obtained by stretching the OH distance, \( R_{\text{OH}} \). The other coordinates are fixed at the geometry in \( S_n \), of which the symmetry is \( C_1 \). Due to the \( C_1 \) symmetry, \( \pi/\sigma \) symmetry does not exist, but we distinguish the major character of the excited states from the shape of their orbitals. If the orbital is mainly localized on the aromatic ring, like the valence orbital, we call it \( \pi \) or \( \pi^* \) and it is denoted by blue squares. In the case where the orbital is diffuse like the Rydberg orbital, it is denoted as \( \sigma^* \) and is indicated by red circles. Where \( R_{\text{OH}} \) is smaller than 1.2 \( \AA \), \( S_1 \) and \( S_2 \) have \( \pi/\pi^* \) and \( \pi/\sigma^* \) characters, respectively. This character is reversed beyond the conical intersection (CI) at \( \sim 1.2 \) \( \AA \) and the \( S_1 \) becomes \( \pi/\sigma^* \) in character. From the Franck–Condon (FC) position (\( \sim 1.0 \) \( \AA \)), the reaction takes place without any barrier in \( S_1 \). This means that the proton moves fast along the reaction coordinate. When the proton crosses over the CI, the character of \( S_1 \) changes to \( \pi/\pi^* \). This means that the \( \pi^* \) electron is transferred to the Rydberg-like \( \sigma^* \) orbital, and the CT complex is generated. The CT complex gives rise to the NIR transitions at \( \sim 8000 \) cm\(^{-1} \) which corresponds to the fast rise of \( 370 \) fs. At that time, the proton – which goes over the CI with significant speed – proceeds into the \( S_2 \) state rather than following the \( S_1 \) potential curve. Here, \( S_2 \) has \( \pi/\pi^* \) character beyond the CI, thus this corresponds to the classical ESPT reaction. The proton can be trapped in \( S_2 \) for a while, and due to the Coulomb attraction this cluster cannot dissociate. The \( S_2 \) state relaxes by internal conversion to \( S_1 \) (of \( \pi/\pi^* \) character), which immediately causes dissociation and the successive formation of the final \( T_d \) product. In this scenario, the observed \( \sim 20 \) ps lifetime for the formation of the \( T_d \) product corresponds to the rate of the IC, \( i.e. \) the lifetime of \( S_2 \). Further detailed discussion allowing a systematic understanding of the ESHT mechanism will be given in the next section.

### 4.6 Larger clusters: proton transfer in the ground state

ESHT has not been detected in PhOH–(NH\(_3\))\(_n\) when \( n = 6 \) or larger, because \( \text{NH}_4(\text{NH}_3)_{n-1} \) (\( n \geq 6 \)) are not observed in the mass-spectrum after two-photon ionization of PhOH–(NH\(_3\))\(_n\) via
the $S_1$ state (see Fig. 1, for $n = 6, 7, \ldots$). The absence of NH$_4^+$(NH$_3$)$_{n-1}$ suggests two possibilities: ESPT or proton transfer in the neutral ground state (GSPT). However, theoretical calculations show that $\pi\pi^*$ states which promote ESPT are higher in energy than the $\pi\sigma^*$ state for clusters of $n \geq 6$, as shown in Fig. 17. Thus, ESPT cannot be responsible for the absence of the reaction products NH$_4^+(NH_3)_{n-1}$.

GSPT is reasonable to explain the observed results. At a certain size, the proton affinity of the ammonia moiety in the cluster will be large enough to induce proton transfer from PhOH in the ground state, leading to the formation of a zwitterion. When the zwitterionic cluster PhO$^-$-NH$_4^+$(NH$_3$)$_{n-1}$ is excited, there is no driving force to dissociate the cluster, since the most stable NH$_4^+$ structure is already formed and this system relaxes in the excited state by evaporation of NH$_3$ fragments.$^{13,15}$ This is consistent with the absence of the fragments NH$_4^+(NH_3)_{n-1}$.

The obvious question, just like in the case of ESPT, is: what is the threshold size for GSPT? The results of the mass-spectra naturally suggest the threshold size $n = 6$. This is supported by the measurement of the ionization potential (IP) of PhOH-(NH$_3$)$_n$ (ref. 61) through a one photon experiment using synchrotron light to avoid the excited state reaction in a two photon process through $S_1$. The measured IP drops for $n = 6$ and larger clusters. This can be explained by the formation of the PT structure in the ground state. It is clear from calculations$^{11,13,44}$ and experiments that the most stable form in the ionic state is the PT structure (PhO-NH$_4^+$) by around 0.8 eV compared to the PhOH-NH$_3$ form. Thus, a PT structure in $S_0$ has a good FC factor to access the adiabatic ionic minimum (i.e. the PT structure), whereas the non-PT structure of ground state PhOH-NH$_3$ cannot access the adiabatic IP (PT structure) in the ion and this leads to a higher IP.

However, the size-selected IR spectra of PhOH-(NH$_3$)$_n$ ($n = 0-11$) measured by IR-UV dip spectroscopy raised a doubt about the threshold size of $n = 6$. The IR spectra showed the signature of PhOH vibrations (such as C-O stretching) even for clusters of $n = 6$ and the expected marker band (C=O stretching) of PhO$^-$ was not found. The disappearance of the C-O stretching band was not clear, even in larger sized clusters, whereas the C-O-H bending vibration, another vibrational signature of PhOH, disappeared at $n \geq 9$. From these results, it was suggested that $n = 6$ may be the beginning of GSPT, but that both PT and non-PT structures co-exist for $n = 6-8$, and all the co-existing species become PT structures at $n = 9$.

A recent theoretical paper, at the M06-2X/cc-pVTZ level of theory, has revealed that clusters of $n \geq 9$ exhibit zwitterionic structures. In this paper, a rather impressive number of cluster structures were investigated and the energy of zwitterionic structures becomes comparable to the non-PT structures only for clusters of $n \geq 9$. There is obviously a discrepancy compared to previous studies and a few possibilities could be responsible.
for this. One is that the evaporation processes following either ionization or excited state reaction are very important and have been overly neglected in the experimental interpretations. In this hypothesis, the ions observed at \( n = 6 \) come from larger ones: this may be tested experimentally but has not been done yet. Another possibility might be that the DFT method is not appropriate for describing such a system and the energy of some structures can be tested with other methods (MP2, CCSD(T), …), although this requires huge computational power at the present time. These possibilities have not been investigated, leaving work to be done by the next generation.

4.7 General model for ESHT and ESPT

We have determined the following three points on the ESHT process mainly from experiments: (1) the reaction speed increases as the cluster size increases, (2) the reaction mechanism changes at \( n = 5 \) and (3) the H transfer proceeds through a tunneling mechanism for small clusters \((n < 5)\).

Let us discuss the cluster size dependence of the ESHT lifetime, which is summarized in Table 1. The ESHT lifetimes for \( n = 2, 3, 4, \) and 5 (electron transfer ET and proton transfer PT) are measured by the rise of the reaction products \( ^{-}\text{NH}_4(\text{NH}_3)_{n-1} \), while that of \( n = 1 \) is extracted from the pump–probe experiment in which the size selection of the cluster is not perfect because of the evaporation of NH\(_3\) after ionization. Thus, the lifetime of \( n = 1 \) may not be “clean” in comparison to other values, however, the order of the lifetime can be trusted. Thus, the ESHT lifetime exhibits a tendency to drop significantly upon increasing the number of solvent molecules in the cluster: nanosecond at \( n = 1 \), sub-nanosecond at \( n = 2 \), tens of ps at \( n = 3 \) and 4, and finally femtosecond for \( n = 5 \).

At first, this change was attributed to the efficient stabilization of \( \pi\sigma^* \) by the higher solvation. It sounds reasonable because the diffuse \( \sigma^* \) electron can easily be stabilized by solvents. Nevertheless, such a sharp stabilization was not found in the theoretical calculations\(^{60,63}\) the absolute energy is lowered by the solvation but the energy gaps between \( \pi\pi^* \) and \( \pi\sigma^* \) are not changed much at the vertical excitation.

This is shown in Fig. 18. Here, the clusters are maintained in \( C_s \) symmetry and the initial structures are optimized in the ground state.\(^{60}\) The clusters are then optimized in the \( \pi\pi^*(A') \) state and the potentials are obtained by stretching the OH distance. For the PhOH–(NH\(_3\))\(_n\), there is no barrier, so that the FC \( \pi\pi^* \) region optimization requires the constraint of the OH distance at 1 Å. It should be noted that the potential curves for \( n = 5 \) (1–5 in the figure) are essentially the same as those in Fig. 16 but are calculated with simpler assumptions, such as \( C_s \).

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| Size (n) | \( \tau_1 \) | \( \tau_2 \) | Reference |
|---------|-------------|-------------|-----------|
| 1       | 1.23 ns (HT)| —           | 51        |
| 2       | 267 ps (HT) | —           | 54        |
| 3       | 23 ps (HT)  | 7 ps (isomerization) | 43        |
| 4       | 15 ps (HT)  | 25 ps (isomerization) | This paper |
| 5       | 370 fs (ET) | 20 ps (PT)  | 59        |

Fig. 18 Potential curve along the OH coordinate in clusters with \( C_s \) symmetry as a function of the cluster size. In red, the \( \pi\pi^* \) potential, and in black, the \( \pi\pi^* \) state. Along this \( \pi\pi^* \) state there are two minima for small sizes: the FC region corresponding to the excited PhOH\(^*\)=(NH\(_3\))\(_n\) species, and at longer OH distance the PhO\(^*\)=(NH\(_4\))\(^+\)(NH\(_3\))\(_{n-1}\) proton transferred (PT) minimum. As the cluster size increases the barrier diminishes and is no longer present for \( n = 5 \). The \( \pi\sigma^* \) potential does not change significantly with the cluster size and in particular the energy gap between the \( \pi\pi^* \) in the FC region and the \( \pi\sigma^* \) stays nearly constant.
symmetry. As one can see, the πσ* potential curves at long O–H distances have a minimum which corresponds to the attachment of the H to N (i.e. formation of ‘NH₄⁺’ but along the O–N coordinate, this ππ* state is dissociative and this is responsible for the separation of the ‘NH₄⁺(NH₃)ₙ₋₁’ cluster away from PhO⁻; the one dimension calculations does not take into account this dissociation of ‘NH₄⁺(NH₃)ₙ₋₁’ from the clusters. Although these curves cannot be compared directly with the experiments, an essential picture emerges from these calculations:

(a) At the equilibrium of the ππ* geometry (around the FC region), the ππ*/πσ* gap does not change as the cluster size increases, being 0.5 eV for the free molecule, 0.4 for n = 2, 3, 4 and rising back to 0.5 eV for n = 5.

(b) The PT structures (ππ* at long OH distance) gradually decrease in energy, being of comparable energy to the ππ* state for n = 3 and being lower in energy for n ≥ 4.

(c) The minimum of the PT structure (ππ*) is obtained at 1.75 Å for n = 1 (1–1 in Fig. 18) and is shifted to 1.5 Å for n = 5, indicating a shrinking of the O–N distance as the cluster size increases in the PT structure.

(d) Since the reaction is controlled by tunneling, both (b) and (c) favor a lowering of the barrier and thus the HT reaction becomes faster.

In Fig. 18, it can be seen that the shape of the πσ* remains approximately the same as the cluster size changes, although the energies are shifted by the cluster size. This is opposite to the pattern observed for the πσ* state. The πσ* potential curves each have two minima, except for n = 5: one at around the FC region and another for the PT (PhO⁻–NH⁻₄(NH₃)ₙ₋₁) structures at greater OH distance. In contrast to the πσ* case, along the O–N coordinate this is a bound state because ionic character prevents fragmentation. For n = 1–3, the minima in the FC region are the global minima but the relative energies of the PT minima gradually decrease and become comparable at n = 3. The PT minimum becomes the global minimum at n = 4 and further stabilization of the PT structure makes the ππ* potential barrier-less at n = 5.

The H transfer tunneling barrier along the OH coordinate is generated by the potential crossing (conical intersection) between the ππ* and the πσ* potential curves. Therefore, the height of the barrier strongly depends on the steepness of the ππ* state as the OH is stretched. Another barrier in this system is that to the PT structure in the ππ* potential curves. For small sizes (n = 1), the PT structure is higher in energy than the FC region and the barrier to the PT structure on the ππ* is also quite high. Thus, the PT reaction cannot take place after optical excitation to S₁. The HT reaction is possible but the barrier is quite high because of the steep ππ* potential. This is a reasonable explanation for the slowest HT reaction in the series. As the cluster size increases, the PT structure and the barrier on the ππ* potential decrease, which also suppresses the barrier height of the H transfer. This tendency terminates in the absence of any barrier for n ≥ 5. Gradual suppression of the barrier can be used to rationalize the size dependence of the ESHT rate as shown in Table 1. This means that the ESHT dynamics vary according to the ππ* potential curve, which in turn is controlled by the stability of the PT structure.

This systematic understanding of the ESHT reaction rate provides a comprehensive understanding of the relation between ESHT and ESPT. It is interesting to note that the initial idea of ESPT which started this research in the 1980s was dismissed at the beginning of the 21st century. However, this review has revealed “The return of ESPT” in a quite different form as being a key mechanism in ESHT.

4.8 Other examples

4.8.1 Naphthol. Despite the extensive study of ESPT (see 1. Introduction), no ESHT has been reported in solvated naphthol clusters. Although naphthol and PhOH are both aromatic alcohols, their excited state dynamics are quite different. The main difference in electronic structure between naphthol and PhOH is that the ππ* is lower in energy (3.8 eV) in naphthol than in PhOH (4.5 eV). From our calculations, it appears that the ππ*/πσ* gap at the ππ* geometry is significantly higher in 1-naphthol (0.95 eV at the aug-cc-pVDZ/CC2 level, compared to 0.52 eV in PhOH). At the ground state geometry, the πσ* in naphthol is around 4.9 eV compared to 5.4 eV in PhOH. This can be understood with very simple arguments. The πσ* state potential is a dissociative curve along the OH coordinate. There is no reason for the shape of the curve to change when the aromatic ring is changed, thus the energy of the πσ* state at the OH equilibrium geometry (1.0 Å) is only dependent on the OH dissociation energy. In this crude approximation, the energy of the πσ* is a constant. In contrast, the energy of the ππ* lowers as the size of the aromatic (particle in a bigger box) – and thus the ππ*/πσ* energy gap – increases, which gives rise to a higher barrier in naphthol than that in PhOH. This gives a simple but comprehensive explanation as to why ESPT, rather than ESHT, takes place in naphthol. This simple scheme has also been used to explain the variation of the excited state lifetime of protonated aromatic amino acids, in which the πσ* state is related to the dissociation of the NH bond.⁴⁴

4.8.2 Substituted PhOH. The H transfer mechanism has been studied in several PhOH derivatives.⁴⁵ Most of the molecules have been studied in their complexes with a single solvent molecule such as ammonia and amines. The first goal was to demonstrate that ESHT occurs in various aromatics, and the mechanisms were discussed in terms of tunneling. It was shown that the excited state lifetimes of fluorophenols, methylyphenols, and their complexes with ammonia were remarkably well correlated with the calculated ππ*/πσ* energy gaps, the lifetime being shorter when the gap is smaller, i.e. the barrier height smaller. In addition to the ππ*/πσ* gap, the coupling of the ππ* and the πσ* is an important parameter to control the reaction rate. In Cₛ symmetry, the ππ* and the πσ* states belong to different symmetry representations A’ and A”, respectively, thus can be coupled only through out of plane vibrations and in particular the out of plane torsion.⁴⁸ In most of the complexes studied, the Cₛ symmetry of the system is conserved and thus the relationship between the lifetime and
the barrier height is relatively regular. In contrast, the excited state lifetime of the cis-ortho-fluorophenol–NH₃ complex⁶⁴,⁶⁵ is very short due to the breaking of the C₆ symmetry in the structure of this complex.

4.8.3 Pyrrole–NH₃ complex. In pyrrole (PyH), the lowest energy excited state is the πσ* state which leads to a direct dissociation⁶⁷ within 80 fs.⁶⁸,⁶⁹ In the PyH–ammonia cluster,⁷⁰ the experimental results reveal that the excited state hydrogen transfer reaction cannot be considered as a simple N–H bond rupture along a repulsive potential energy curve, as was at first suggested by comparison with the free molecule. The results of the lifetime measurements as well as the competition between the H-atom transfer reaction and the evaporation imply that the reaction proceeds via a fairly long lived (10–30 ps), relatively deep intermediate state. The excited state potential energy surface calculated at the CASSCF level agrees with these conclusions: two minima corresponding to very different structures are apparent and the reaction path after the initial excitation can be described in four steps. The first step is a contraction of the N–N distance accompanied by a vibrational motion of the pyrrolic N–H bond. Then, at the appropriate N–N distance, an electron transfer from PyH to the solvent occurs, immediately followed by proton transfer from PyH to ammonia (concerted electron/proton), which produces the radical pair Py’–NH₄ with a very diffuse Rydberg electron cloud around the ammonium. Then the cluster breaks into the two separate radicals.

4.8.4 Indole–NH₃ complex. In indole, as for PhOH, the H loss in the free molecule⁷¹ and the hydrogen transfer to the NH₃ cluster have been demonstrated.⁷²–⁷⁷ In the free molecule, the H loss through the πσ* state⁷⁷ has been observed but is also contaminated by H atoms generated by one or more (unintended but unavoidable) multiphoton processes. UV excitation of indole–(NH₃)ₙ clusters leads to a H atom transfer reaction and the formation of ‘NH₄(NH₃)ₙ radicals. With an excess of a few tenths of an electron volt, the reaction proceeds very quickly in the 100 ps regime. As for PhOH, the H transfer process explains why the excited-state lifetime is short (sub-nanosecond) in indole-ammonia clusters as compared to all other indole clusters (with water, methanol, etc.) where the lifetime is much longer.

The lifetime of the first excited state in the 3-methylindole–NH₃ complex is strongly dependent on the intermolecular vibrations. As experimentally observed,⁷³ the excitation of the NH…N stretching coordinate is expected to favor the H transfer reaction. High-level ab initio optimization of the equilibrium geometry in the S₁ state and calculated vibrational frequencies compared with the experimental observations seem to indicate that the in-plane bending vibration also favors the H transfer from the aromatic chromophore (3-methylindole) to the solvent (ammonia), the reaction being 530 ps on the 0–0, 145 ps for 1 quantum in the N–H…NH₃ stretch and 205 ps for 1 quantum in the N–H…NH₃ bend.

4.8.5 Hydrogen transfer and protonated molecules. Unexpectedly, the competition between the PT and the H transfer mechanisms has been found in the excited state dynamics of protonated aromatic amino acids. In the ground state, if the proton is localized on the amino group, electronic excitation triggers the proton transfer onto the aromatic ring and this lead to Cₓ–Cᵧ bond rupture. In an intermediate energy range, the hydrogen transfer between the protonated amino group and the carboxylic group leads to CO₂ and H₂O loss, while H loss was experimentally evidenced upon excitation at high energy.⁷⁸

5. Concluding remarks

PhOH–(NH₃)ₙ is a very rich system in which many subtleties of the reaction dynamics of the hydrogen mediated reaction exist. The excited state dynamics is systematically changed by the size of the ammonia moiety, of which the basicity increases with the number of molecules. Although this idea, initiated in Leutwyler’s group⁷⁹ was used for the study of ESPT in clusters, it also explains systematic variation in the ESHT mechanism. This is because the change in the basicity distorts the potential surface of the optically active πσ* state due to stabilization of the PT local minimum. The potential distortion gives rise to a significant change in the tunneling barrier to H atom release, which is produced by the crossing to the πσ* potential surface.

The lowering in energy of the PT structure is strongly size dependent. For n = 1, the PT structure at O–H = 1.8 Å is higher in energy by 0.15 eV than the FC structure (PhOH–(NH₃)ₙ), and its energy decreases as the cluster size increases. At n = 4, this structure seems to be more stable than that of the FC, although a barrier to PT transfer still exists, and at n = 5 the energy of the PT structure is so low (–0.22 eV) that the PT reaction becomes barrierless. In contrast to the πσ* state, the πσ* state is relatively insensitive to the cluster size, primarily because the potential shape of this repulsive state is essentially independent of the molecular system. This insensitive character is given by the energy of its asymptote at long distance. It corresponds to the bond dissociation energy of the OH (NH) which is more or less the same for many molecules (OH bond energy 4.8 eV, or 3.2 for NH). Thus, the systematic change of the πσ* potential surface directly gives the clear size dependence of the ESHT mechanism, as well as the ESPT (rather than ESHT) reaction in naphthol.

Quite unexpectedly, the role of the hypervalent dissociative Rydberg state – evidenced experimentally and theoretically for the PhOH–(NH₃)ₙ system – has been found to be key to processes in the excited states of many protonated ions such as the aromatic amino acids. In these cases, the excess proton on the amino group (NH₃⁺) can easily accept an electron after optical excitation to a hypervalent dissociative state which leads to H loss or H transfer depending on the nearby atom. This first step controls both the ensuing long term dynamics, and the selectivity of the fragmentation processes.⁷⁹ This work is a benchmark, demonstrating the role of the excited state reaction in leading to the formation of radical pairs through the H loss, which was too often neglected before. The H loss from aromatic molecule is now quite a standard paradigm. Extension of the ESHT reaction to describe other processes is now occurring, notably in the description of excited state H abstraction by aromatic molecules, which is the reverse process of ESHT and is demonstrated in photoexcited pyridine–H₂O system.⁸⁰–⁸¹ These
reverse process studies are still in their infancy but might grow rapidly to resolve the world energy crisis. We hope that the up-to-date mechanism of the ESHT reaction and re-definition of the role of ESPT will thus provide a firm base for the understanding and design of this new direction of excited state dynamics.

Author contributions
C. J., M. M. and M. F. conceived and designed the project. M. M. measured and analysed the spectra. C. J. performed and analysed the quantum chemical calculations. C. J. and M. F. co-wrote the paper.

Conflicts of interest
There are no conflicts to declare.

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