Real-Time Observation of Fermi Resonances in the $S_1$ State of Phenol

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Supporting Information

ABSTRACT: Fermi resonances in the first electronically excited ($S_1$) state of phenol have been observed in real time. Quantum beats associated with coherent superposition of Fermi resonant eigenstates are manifested as temporal oscillations of the ionization cross sections of which the amplitudes are strongly dependent on the total ionization energy. This indicates that coherently excited eigenstates are effectively decomposed into their zeroth-order states, providing the unique opportunity for the investigation of nonstationary state dynamics in real-time. Energy gaps ($\Delta \nu$) of eigenstates within the laser coherence width have been most precisely determined up to date, giving $\Delta \nu \sim 3.302 \pm 0.001$ or $1.655 \pm 0.001$ cm$^{-1}$ for the $1^{1}/41\text{b}_{1}$ or $12^{2}/8\text{a}_{1}$ Fermi doublet, respectively. Dephasing rate suddenly increases as the $S_1$ internal energy becomes above $\sim1500$ cm$^{-1}$, revealing the important role of energy randomization dynamics during the H atom tunneling process of phenol in $S_1$.

Fermi resonances are misleadingly considered to be quite common in polyatomic molecular systems as there are supposedly many chances for a couple of quantum states belonging to the same irreducible symmetry species to coincide in their vibrational energies. In the ground-state infrared or Raman spectroscopy, it is often that overtone or combination modes otherwise optically dark obtain significant oscillator strengths because of strong couplings with nearly isoenergetic fundamental modes. Although it is somewhat straightforward to identify Fermi doublets in small molecular systems, it is nontrivial to identify those doublets in vibrational spectra of relatively large polyatomic molecules with many vibrational degrees of freedom. Real-time probing of such resonances provides more direct evidence where eigenstates comprising the superposition state are coherently excited within the coherence width of the pump laser pulse. Then quantum interferences of two or more distinct time-dependent wave functions embedded in the superposition state are manifested as oscillatory time-evolution features in various spectroscopic probing schemes. These include laser-induced dispersed fluorescence, time-resolved photoion, or photoelectron spectroscopy. Although many examples of quantum beats among coherently excited states were reported, real-time probing of strongly coupled Fermi resonances in polyatomic molecular systems has been quite rare. Photochemistry of phenol ($C_6H_5OH$) has been very much-studied because it is one of the prototypical systems undergoing the quantum interferences of two or more distinct time-dependent wave functions embedded in the superposition state. As a result, H atom tunneling takes place through the adiabatic reaction barrier under the $S_1/\text{S}_2$ conical intersection seam, and its height is predicted to be quite high assuring the prevalence of the tunneling process even at $\sim3500$ cm$^{-1}$ above the $S_1$ origin. Recent picosecond time-resolved spectroscopic studies have revealed that the H atom tunneling rate strongly depends on the low-frequency vibrational modes seemingly orthogonal to the reaction coordinate, demonstrating that the tunneling dynamics is indeed multidimensional in nature. The lifetime of $S_1$, which mainly reflects the tunneling rate, has been measured to be $\sim2$ ns at the $S_1$ origin and it decreases to $\sim0.8$ ns at $\sim3600$ cm$^{-1}$ above $S_1$. Naturally, in these circumstances, it is an important question to what extent the energy randomization of intramolecular vibrational redistribution (IVR) plays a role during the concomitant tunneling process in terms of washing out the mode effect and thus averaging out the whole reaction dynamics governing reaction rates and energy disposals. In this work, we report the real-time observation of Fermi resonances and the energy flow dynamics among vibrational manifolds of the $S_1$ phenol undergoing the H atom tunneling predissociation.

A couple of well-separated peaks observed at $S_1$ internal energies ($E_{nm}$) of 935 and 938 cm$^{-1}$ in the $S_1$--$S_0$ nanosecond laser resonant two-photon ionization (R2PI) spectrum had been assigned as the $1^{1}/41\text{b}_{1}$ or $12^{2}/8\text{a}_{1}$ Fermi doublet, respectively. Similarly, Bist et al. had reported the room-temperature gas-phase absorption spectrum showing closely spaced states in the same frequency region separated by $\sim3.37$ cm$^{-1}$. In the R2PI spectrum taken...
here using the picosecond laser pulse with the temporal duration of 1.2 ps and the coherence width of \(\sim 25 \text{ cm}^{-1}\), these \(1^1/4^1\text{0b}^1\) bands with a separation of \(\sim 3 \text{ cm}^{-1}\) are coherently excited, giving the necessary condition for the observation of quantum interferences thereafter. It should be noted here that the clear-cut characteristic of the Fermi-resonance is that the optically dark states get significant oscillatory strength via the coupling to the optically bright state. We have carried out Franck–Condon simulations for the \(S_1\)–\(S_0\) transition of phenol (see the Supporting Information). It is evident that the Franck–Condon factor of the \(4^1\text{0b}^1\) combination band is predicted to be 4–6 orders of magnitude smaller than that of the \(1^1\) fundamental mode. However, in the experiment, those coupled \(4^1\text{0b}^1\) and \(1^1\) modes show comparable intensities, strongly indicating that these modes belong to the Fermi doublet. Herein, including the \(1^1/4^1\text{0b}^1\) doublet, we have found several outstanding Fermi resonances in the \(S_1\) phenol. Interferences from coherently excited eigenstates have been clearly observed in our picosecond time-resolved parent ion transients, giving the most precise energy gaps to date between strongly coupled eigenstates. Most surprisingly, it is found that the oscillatory amplitude of the quantum beat here is unusually large (\textit{vide infra}) if the ionization energy is tuned just above the ionization threshold, strongly indicating that Fermi resonance eigenstates are almost completely decoupled into nonstationary zeroth-order quantum states in real time.

Prominent oscillatory features on the \([\text{C}_6\text{H}_5\text{OH}]^+\) transients are observed following the coherent population of \(1^1/4^1\text{0b}^1\) eigenstates of the \(S_1\) phenol, giving the beating period of \(\sim 10\) ps (Figure 1). The transient has been analyzed either by subtracting the exponential decaying function from the experiment or by fitting the transient with a damped-cosine-implemented formula (see eq S1 in the Supporting Information), giving consistent values as listed in Table 1. The frequency gap \(\Delta \nu\) of the coherently excited \(1^1/4^1\text{0b}^1\) eigenstates is estimated to be \(3.302 \pm 0.001 \text{ cm}^{-1}\). This is the most precise value to date while it is widely in accord with the previously reported spectroscopic value of 3.37 or 3.3 \(\pm 0.1 \text{ cm}^{-1}\). When the laser pulse populates another \(S_1\) Fermi doublet located at 1564 and 1566 \(\text{ cm}^{-1}\), which had been assigned as \(8^2\) and \(8a^1\) modes, respectively, the clear beating pattern has also been found, this time with the oscillation period of \(\sim 20\) ps, Figure 2b. Transient analyses give the beating frequency of 1.655 \(\pm 0.001 \text{ cm}^{-1}\). Notably, this is quite different from the previously reported literature value of 1.78 \(\pm 0.001 \text{ cm}^{-1}\), demonstrating that time-resolved measurement here gives the much-improved coupling constants in terms of both accuracy and precision, Table 1. Intriguingly, different types of \(1^1/4^1\text{0b}^1\) Fermi resonances are also found in combination with the \(6a^1\) or \(12^1\) mode to give the \(1^16a^1/4^1\text{0b}^16a^1\) (1408/1412 \(\text{ cm}^{-1}\)) or \(1^112^1/4^1\text{0b}^112^1\) Fermi doublet (1716/1719 \(\text{ cm}^{-1}\)), respectively (see Figure S2 in the Supporting Information). The beating frequency analysis for the \(1^16a^1/4^1\text{0b}^16a^1\) and \(1^112^1/4^1\text{0b}^112^1\) doublet gives \(\Delta \nu = 3.472 \pm 0.004\) or 3.201 \(\pm 0.013 \text{ cm}^{-1}\), respectively.

The close proximity of two quantum levels belonging to the same irreducible symmetry species does not always guarantee the existence of a Fermi resonance. Moreover, it is not straightforward to observe quantum interferences experimentally as the total ionization cross section, for instance, would average out constructive and destructive quantum effects. In this regard, our clear-cut observation of quantum beat from the ion transients is quite notable. The oscillation amplitude is found to be strongly dependent on the probing wavelength \(\lambda_{pr}\) as its modulation depth exceeds more than 50% of the ion transients at \(\lambda_{pr} = 309 \text{ nm}\) for the \(1^1/4^1\text{0b}^1\) Fermi levels. The oscillation amplitude decreases sharply with the decrease of the probing wavelength, showing that it falls off less than 15% of the total ion signal at \(\lambda_{pr} = 301 \text{ nm}\) (Figure 2). Considering the propensity rule of \(\Delta \nu = 0\) in the \(D_0\)–\(S_1\) transition, this indicates that the vibrational adiabatic ionization leads to the large fluctuations of the ionization cross section as the nearly degenerate zeroth-order \(S_1\) states split remote in the \(D_0\) state. For instance, theoretical calculations predict that the vibrational frequency of the \((4^1\text{0b}^1)^+\) combination mode is much higher than that of the \(1^1\) mode in \(D_0\) (see Table S1 in the Supporting Information). Therefore, the ionization cross section at \(\lambda_{pr} = 309 \text{ nm}\) (32 362.5 \(\text{ cm}^{-1}\)) should mostly represent the Franck–Condon (FC) overlap between \(1^1\) and \(1^1\) in \(S_1\) and \(D_0\) respectively. This acute FC overlap fades away in terms of its sensitivity on the probing spectral window as many more different vibrational modes of \(D_0\) contribute to the total ionization cross-section as \(\lambda_{pr}\) is decreased to 301 nm (33 222.6 \(\text{ cm}^{-1}\)), giving the significant reduction of the oscillation amplitude.
Further variation of $\lambda_{pr}$ into the shorter wavelength results in the entire absence of the oscillating feature (see Figures S2 and S3 in the Supporting Information). This phenomenon could also be seen for 12$^2$/8a and all other Fermi levels, giving the interference pattern where the oscillating depth is largely dependent on $\lambda_{pr}$.

Decoupling of Fermi resonances into the zeroth-order quantum states in real time could be rationalized by the following simple model. Namely, we could represent Fermi resonant eigenstates as follows:

$$|i\rangle = \frac{1}{\sqrt{2}}(|\phi_1\rangle + |\phi_2\rangle) \quad |j\rangle = \frac{1}{\sqrt{2}}(|\phi_1\rangle - |\phi_2\rangle)$$

Here $|i\rangle$ and $|j\rangle$ are molecular eigenstates whereas $|\phi_1\rangle$ and $|\phi_2\rangle$ are zeroth-order states where magnitudes of coupling coefficients are assumed to be equivalent. The coherently excited state by the picosecond laser pulse is then the superposition of $|i\rangle$ and $|j\rangle$, giving the time-dependent wave function, $\Psi(t)$.

$$\Psi(t) = \frac{1}{\sqrt{2}}(e^{-i(E_1/h)t} + e^{-i(E_2/h)t})$$

The ionization cross-section near the ionization threshold will reflect the projection of $\langle \Psi(t) | \langle \phi_1^+ | \phi_1 \rangle = e^{-i(E_1/h)t} + e^{-i(E_2/h)t} \rangle$.

$$\langle \phi_1^+ | \Psi(t) \rangle = \frac{1}{2} \langle \phi_1^+ | \phi_1 \rangle (e^{-i(E_1/h)t} + e^{-i(E_2/h)t})$$

Here, according to the propensity rule of $\Delta \nu = 0$ in the ionization, $\langle \phi_1 | \phi_1 \rangle \equiv 0$ should be a good approximation. The ionization cross-section near the threshold then will be proportional to $l(\phi_1^+ | \Psi(t) \rangle)^2$ as follows where $\langle \phi_1^+ | \phi_1 \rangle \equiv c_1$.

$$l(\phi_1^+ | \Psi(t) \rangle)^2 = \frac{1}{2} c_1 (e^{-i(E_1/h)t} + e^{-i(E_2/h)t})$$

This value spans from zero to $c_1^2$ as a function of time, demonstrating that the coherently excited Fermi doublet could be completely decoupled into the zero-order state according to the oscillatory quantum beat in real time.

In fact, coherently populating Fermi resonance levels could be regarded as one of the prototypical examples of the restricted IVR as it represents the typical energy flow between the optically bright zeroth-order state and a nearly isoenergetic yet optically dark quantum state if the terminology of IVR is used in the strict way. As long as further IVR into dark quantum manifolds is not activated, interferences due to coherent excitation of Fermi eigenstates are expected to persist during the whole $S_1$ lifetime. However, if the “so-called” dissipative IVR takes in action, then such an interference would dephase accordingly. Namely, whereas the $S_1$ lifetime of phenol is largely determined by the H atom tunneling rate, the oscillation amplitude could decrease with time as the interference of Fermi levels dephases as IVR into dark manifolds intrudes the otherwise two-state coupling of Fermi resonance. From the fits to the experiment using eq S1, the dephasing time constants ($\tau_{deph}$) of infinity or ~400 ps are obtained for the 1$^1$/4$^1$10b$^1$ (Figure 1) and 12$^2$/8a$^1$ Fermi levels, respectively (Figure S1). This is quite reasonable as the
S$_1$ internal energy of $1^{1}/4^{1}10^{b_{1}}$ is centered at $\sim$937 cm$^{-1}$ while that of $12^{2}/8a_{1}$ is located at $\sim$1565 cm$^{-1}$. As such, the dissipative IVR is not activated yet within the $S_{1}$ lifetime of $\sim$1.0 ns at $\sim$937 cm$^{-1}$. Meanwhile, the dissipative IVR beyond Fermi-type coupling occurs at $E_{\text{int}} \sim 1565$ cm$^{-1}$ with a rate faster than tunneling. In retrospect, this behavior seems to be quite consistent with recently measured tunneling rate constants of the $S_{1}$ phenol which show large fluctuations at $E_{\text{int}}$ lower than $\sim$1300 cm$^{-1}$ and become monotonic at the higher internal energy. It is notable that the dephasing rate gets increased sharply with increasing the internal energy, giving $\tau_{\text{dep}}$ of $\sim$75 ps at $E_{\text{int}} \sim 1718$ cm$^{-1}$ associated with the $1^{12}/4^{1}10^{b_{1}}12^{1}$ Fermi doublet (Figure S2b). In addition, the fast decaying component with the ultrashort lifetime less than tens of picoseconds has been found in the initial part of the $1^{12}/4^{1}10^{b_{1}}12^{1}$ transient. This ultrafast decaying component of the transient should be associated with IVR in $S_{1}$ as the ionization cross-section becomes rapidly dispersive with the accompanying mode-randomization.

It is notable again that the Fermi-type interference survives during the entire $S_{1}$ lifetime of phenol even at the internal energy of $\sim$937 cm$^{-1}$, giving the full support for the observation of the strong mode-dependent tunneling rates. This also promises the mode-selective chemistry of the excited polyatomic systems as the mode randomization is found to be rather slow compared with the reaction rate. Although low-frequency modes are hardly associated with the reaction coordinate of the specific bond cleavage, for instance, their excitations are closely related to the subtle structural changes that are quite often very critical in nonadiabatic transitions on the multidimensional conical intersection seam or in shaping the adiabatic transition states diversely defined along the orthogonal conformational coordinates. In this regard, exploration of mode-effect on excited-state dynamics and its relation to the energy flow among close-lying quantum states as observed here will shed new light on the structure—function relationship in a number of molecular systems.

## EXPERIMENTAL METHODS

Experimental methods of frequency- and time-resolved pump–

probe ion yield spectroscopy have been recently given elsewhere.$^{31}$ Briefly, tunable laser pulses in the ultraviolet wavelength range are generated from two optical parametric amplifiers (TOPAS-800 ps, light conversion) pumped by the 50:50 split fundamental outputs from the 1 kHz picosecond Ti:sapphire regenerative amplifier (UBBR2.5-1UV, Newport) mounted on a computer-controlled 220 mm optical delay line stage (DDS220, Thorlabs), where the delay step sizes of 0.8 ps are used for the range of monitoring oscillatory features.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcl.9b03393.

Details in fitting procedure of the transients with oscillations; transients in the full range of time delay along with the beating frequency analyses; transients for the additional Fermi levels of combined $1^{1}/4^{1}10^{b_{1}}$ with the modes 6a and 12; comparison of transients for $S_{1}$ vibronic levels near $1^{1}/4^{1}10^{b_{1}}$ with various probe wavelengths; results of the harmonic frequency calculations; additional notes on the $S_{0}$ → $S_{1}$ Franck–Condon simulation; time-dependent Fermi doublet in the experiment (PDF)

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### Notes

The authors declare no competing financial interest.

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