Femtosecond Wavepacket Dynamics Reveals the Molecular Structures in the Excited (S1) and Cationic (D0) States

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ABSTRACT: Molecular structures in the electronically excited (S1) and cationic (D0) states of 2-fluorothioanisole (2-FTA) have been precisely refined from the real-time dynamics of the femtosecond (fs) wavepacket prepared by the coherent excitation of the Franck–Condon active out-of-plane torsional modes in the S1 ← S0 transition at 285 nm. The simulation to reproduce the experiment in terms of the beating frequencies gives the nonplanar geometry of 2-FTA in S1, where the out-of-plane dihedral angle (φ) of the S-CH3 moiety is 51° with respect to the molecular plane. The behavior of the fs wavepacket in terms of the amplitudes and phases with the change of the probe (ionization) wavelength (λprobe = 300–330 nm) provides the otherwise veiled structure of the cationic D0 state. While the 2-FTA cation adopts the planar geometry (φ = 0°) at the global minimum, it is found to have a vertical minimum at φ ≈ 135° from the perspective of the D0 ← S1 vertical transition. Ab initio calculations support the experiment quite well although the simulation using the model potentials could improve the match with the experiment, giving the new interpretation for the previously disputed photoelectron spectroscopic results.

INTRODUCTION

Molecular structures dictate the fate of the chemical reactions. The conformational isomer, which could be converted to the other conformer quite easily by the nearly free internal rotation about the single bond axis, has been found to give a profound influence on the chemical reactivity.1,2 In ambient conditions though, it would be nontrivial to confine the molecule to a specific geometry as all possible structural isomers may interconvert freely with very low barriers among them. In this respect, the spectroscopy and dynamics on the supersonically cooled molecular beam have been extremely useful as the internal energy is minimized so that the molecule is frozen at the minimum-energy geometry with respect to all normal mode coordinates in the ideal case where the vibrational temperature is also extremely low. Once the molecule is frozen at the specific geometry, it could be further utilized for chemical reactions such as the photodissociation/unimolecular reaction or stereospecific bimolecular reactions. When two or more structural isomers are populated together in the molecular beam, one can select just one specific isomer at a time by either the spectroscopic3,4 or Stark deflection technique.5–7 As a matter of fact, although rare, there are notable examples that report extremely interesting conformational specificity in chemical reactions.8–10 In the photodissociation reactions, as the optical excitation is prompt, the phase space of the excited state is limited by the Franck–Condon window. If the excited state is repulsive along the particular bond extension coordinate, the initially chosen molecular geometry is then exactly reflected in the dynamic outputs. Namely, different geometrical isomer selection could lead to quite different outputs in terms of the rate, yield, and energy disposal dynamics, depending on the nature of the multidimensional excited-state potential energy surfaces.8,9

The more elaborate case however could be found when the excited state is bound or quasi-bound. In this case, as long as the Franck–Condon overlap is nonzero, one can explore many different excited-state nuclear configurations spanned by individual vibronic modes. Especially when the internal energy of the excited state is small (e.g., <1000 cm−1), the rate of intramolecular vibrational redistribution (IVR) is often slower than the rate of the excited-state reaction, giving the great opportunity to explore many different nuclear configurations of the excited state in terms of their chemical reactivities although the non-Franck–Condon region would be still blind.

The mode-selective reaction dynamics have long been investigated for many interesting chemical systems, which involve predissociation or tunneling process. The mode dependence of the dynamic outputs such as reaction rates and/or product yields has been extremely useful in disentangling the complicated coupling dynamics among...
numerous vibronic modes associated with the reaction channels. In this aspect, the dynamics around conical intersections have been studied in impressive detail in recent studies on thioanisoles,11–17 thiophenols,18–25 or phenols.26–37 For instance, the nonadiabatic transition probability has been found to be strongly mode-dependent in the S-CH3 bond predissociation of the electronically excited (S1) thioanisole.11,13,14,16,17 The nonadiabatic transition probability is strongly enhanced when (the nuclear configuration of the reactive flux given by) the specific vibronic mode accesses the conical intersection region. From the exploration of the wide range of the phase space, the otherwise formidable task of capturing the conical intersection in terms of the structure and its dynamic role becomes a reality now. The vibronic mode dependence has also been observed in the tunneling reactions. Namely, the tunneling in the H-atom detachment reactions of phenols and thiophenols turns out to be mediated by the upper-lying surface crossings in the proximity of the conical intersections along the multidimensional seam coordinates.38–40

As the direct optical transition has the intrinsic limit in terms of the Franck–Condon window, however, the expansion of the exploring region of the excited-state phase space is quite desirable for the investigation of multiple facets of the conical intersection, for example. One may employ the double resonance excitation technique such as the IR + UV scheme.40,41 Here, the IR laser pulse prepares the specific rovibrational mode in S0 whereas the subsequent UV excitation would then modify the accessible Franck–Condon region. Depending on the extent of IVR in S0 prior to the UV excitation, the characteristic of accessible quantum states could also be modified. Interestingly, the other way to expand the exploring region of the excited state is the chemical substitution. A recent study of the photodissociation dynamics of 2-fluorothioanisole (2-FTA) is quite notable as it has been found that, unlike the case of thioanisole, it undergoes the substantial structural change upon the S1 ← S0 optical transition.42 Namely, whereas thioanisole maintains its molecular planarity in the S1 ← S0 optical transition,11,12,13 the planar 2-FTA in the ground state (S0) undergoes the significant structural change to adopt the nonplanar geometry in S1 in terms of the dihedral angle (φ). As the S-CH3 dihedral angle is one of the most strongly coupled coordinates in the branching plane of the conical intersection,12,13 the nonplanarity of the S1 state should open the chance of exploring the S1/S0 conical intersection region from the completely different perspective; the otherwise veiled nonadiabatic path via the pseudo-conical intersection seam could be opened.42 This approach is similar to the exploration of the other side of the moon. Even though the structural change upon the S1 ← S0 optical transition could be inferred from the spectroscopic evidence, it is not straightforward to grasp the exact shape of the excited-state potential energy surface, especially because the ab initio calculation of the excited state is quite challenging for polyatomic molecules at the present time.

Herein, we have found that the femtosecond coherent wavepacket interferometry is extremely useful in characterizing the excited-state geometry of 2-FTA. The molecular geometries of thioanisole and its derivatives have long been controversial in terms of the dihedral angle of the S-CH3 moiety with respect to the molecular plane.35–36 Although the presence of the geometrical isomer of φ = 90° has been conjectured in many previous studies, thioanisole prepared in the supersonic jet has been found to adopt the planar geometry (φ = 0°) in the ground state as the global minimum structure.43–46 As mentioned above, although thioanisole remains planar in S1, 2-FTA becomes nonplanar upon the S1 ← S0 transition. The planar-to-nonplanar structural change of 2-FTA then allows the coherent excitation of the dihedral torsional modes within the coherence width of the femtosecond (fs) pump laser pulse.47–54 The consequent fs wavepacket in S1 propagates along the dihedral torsional angle (φ) coordinate, giving the clear-cut quantum beats in the parent ion transient obtained by monitoring the ionization cross section as a function of the delay time between the pump (S1 ← S0) and probe (D0 ← S1) fs laser pulses. Through the simulation of the experiment, the exact shape of the S1 potential energy surface along the dihedral angle could be extracted. The D0 structure could also be inferred from the apparent phase inversion of the quantum beat as the ionization wavelength is varied. The current fs wavepacket interferometry of 2-FTA unravels the detailed molecular structure of the S1 state to give the most critical information regarding the conical intersection structure, allowing the exploration of the nonadiabatic dynamics from a completely different perspective.

EXPERIMENTAL SECTION

Experimental details have been described elsewhere.16,37 Briefly, the 2-FTA sample was mixed with the neon carrier gas and expanded into vacuum through a nozzle orifice (φ = 100 μm) of the Even-Lavie pulsed valve at the repetition rate of 200 Hz with the backing pressure of ~2 bar. The molecular beam was collimated by a skimmer (φ = 2 mm) and crossed by femtosecond ultraviolet laser pulses. The ions were accelerated in the time-of-flight tube toward a Chevron-type microchannel plate (MCP) detector coupled with the phosphor screen to be detected by the photomultiplier tube (PMT). The femtosecond pump (285 nm) and probe (300–330 nm) pulses (cross-correlation full width at half-maximum (FWHM) of 150 fs) were generated by two independent optical parametric amplifier (OPA) units (TOPAS-prime, Light Conversion). The OPA units were pumped by the 25:75 split of the fundamental output of 1 kHz femtosecond Ti:Sapphire regenerative amplifier system (Legend Elite-USP, Coherent, 35 fs) seeded by an 80 MHz femtosecond Ti:Sapphire oscillator (Vitara-T-HP, Coherent). The probe pulse was linearly polarized, which is parallel to the detection plane, and the polarization of the pump pulse was kept at the magic angle (54.7°). The pump–probe delay was manipulated by a 300 mm long linear translational stage (M-531.PD1,Physik Instrumente) combined with a retroreflector. The geometry optimization and excited-state calculations were carried out using the state-averaged complete active space self-consistent field (SA-CASSCF) method with the 6-311++G(2df,2pd) basis set using the Molpro program package.75 For the lowest four electronic states of cationic 2-FTA, the (11,11) active space was generated using three pairs of π/π* orbitals on the benzene ring, two pairs of π/π* orbitals on the S-CH3 bond, and a nonbonding p-type orbital of the sulfur atom. Potential energy curves along the S-CH3 dihedral angle (φ) coordinate were obtained either by the geometry optimization of the D0 state at each step of the φ increment (relaxed scan) or by freezing all of the geometric parameters other than φ (rigid scan). The single-state second-order perturbation theory (SS-
CASPT2) correction was implemented with an imaginary level shift of 0.3 au.

## RESULTS AND DISCUSSION

In our previous report, the resonance-enhanced two-photon ionization (R2PI) spectrum of 2-FTA obtained by the nanosecond laser pulse exhibits the prominent low-frequency S1 vibronic progressions of 32 and 82 cm$^{-1}$ corresponding to one ($\tau^1$) and two quanta ($\tau^2$) of SCH3 torsion mode. The equilibrium geometry and wavefunctions of the S1 state of 2-FTA have been constructed by solving a Hamiltonian matrix in the discrete variable representation (DVR) with 1440 grid points of the SCH3 torsion angle from $-180$ to $180^\circ$ (Figure 1a). The ground-state potential is modeled by a harmonic (quadratic) function, whereas the double-well potential energy function of $V(x) = \frac{1}{2}kx^2 + Ae^{-\beta x}$ has been employed for the depiction of $S_1$. Parameters of the model potentials are optimized to fit the experiment using the genetic algorithm (Supporting Information (SI)). The resultant analytic S1 potential energy curve suggests that the (most probable) dihedral angle of the S-CH3 moiety should be $51^\circ$ at the zero-point energy level. In other words, two minima of the potential energy curve are located at $\varphi = \pm 57^\circ$ with a small barrier of 40 cm$^{-1}$. The out-of-plane dihedral angle in $S_1$ is found to be actually much nearer to the perpendicular geometry than the previously reported value of $\sim 12.5^\circ$ predicted based on the unscaled ab initio calculations. The Franck–Condon simulation based on the calculated wavefunctions of $S_0$ and $S_1$ then gives the band intensity ratio of $\tau^3:\tau^1:\tau^2 = 1.0:1.8:0.4$. This is in excellent agreement with the one-color R2PI experiment. The large negative anharmonicity (which is larger than unity for the steeper potential energy curve than predicted by the harmonic approximation) of $\gamma = (\langle \tau^3 - \tau^2 \rangle - \langle \tau^1 - \tau^3 \rangle) = 1.58$ also strongly supports the largely distorted geometry of the S1 2-FTA. This is quite consistent with the previous reports on other double-minima systems of 2-fluorophenol ($\gamma = 1.28$, $\varphi \approx 37^\circ$), 2-hydroxyphenyl ($\gamma = 1.23$, $\varphi \approx 24^\circ$), or 2-methoxythiophenol ($\gamma = 1.13$, $\varphi \approx 20^\circ$) in terms of the correlation between $\gamma$ and $\varphi$.

The parent transients of 2-FTA obtained by the pump (285 nm) and probe (330–300 nm) laser pulses are shown in Figure 1b. Time-resolved quantum beats added on the slow decaying background ($\tau \approx 500$ ps) are clearly observed. Fast Fourier transform (FFT) of the oscillatory part gives the beating frequencies of 31.6 and 49.7 cm$^{-1}$, and these exactly match with the frequency differences of coherently excited torsional modes of ($\tau^3 - \tau^2$) and ($\tau^1 - \tau^3$), respectively. The fs wavepacket interferogram thus confirms and refines the 2-FTA ($S_1$) structure of the nonplanar geometry with the precise dihedral angle of $\sim 51^\circ$. While the beating frequencies remain constant, the amplitudes and phases of the oscillatory part are found to be strongly modified by changing the probe laser wavelength. For instance, the phase in the transient obtained at $\lambda_{\text{probe}} = 330$ nm is completely inverted by the factor of $\pi$ in the transient at $\lambda_{\text{probe}} = 300$ nm. Meanwhile, the oscillating amplitude almost vanishes in the transient at $\lambda_{\text{probe}} = 315$ nm. As a matter of fact, the decrease of the quantum beat amplitude could be attributed to the fact that the D0 $\leftrightarrow$ S1 cross section becomes insensitive to the molecular planarity as the wider phase space of the cationic state (with respect to $\varphi$) is sampled with increasing the probe energy from 330 to 315 nm. And yet, it is quite remarkable that the beating amplitude gradually increases again as $\lambda_{\text{probe}}$ decreases further with the inverted phase. This experimental finding indicates that there should be two distinct ionization processes having the totally different phases in the probing wavelength range of 330–300 nm. As the D0 minimum-energy structure is planar, the quantum beat is the consequence from the swing of the S1 wavepacket along the out-of-plane dihedral angle. Namely, the planarity of the $S_1$ molecular structure is monitored by the D0 $\leftrightarrow$ S1 cross section, which is either maximized or minimized as the S1 wavepacket is located at the planar or nonplanar geometry, respectively. Since the temporal oscillation is solely due to the out-of-plane motion of the S1 wavepacket, the ionization cross section is influenced only by the planarity of the cationic state.
Therefore, the experimental fact that the phase gets inverted as $\lambda_{probe}$ changes from 330 to 300 nm indicates that the ionization cross section at $\lambda_{probe} = 300$ nm becomes maximized when the $S_1$ wavepacket is instantly located at the nonplanar geometry.

The adiabatic potential energy curves along $\phi$ have been calculated for the three lowest electronic states of the 2-FTA cation (Figure 2). The minimum-energy structure of $D_0$ optimized at the SA4-CASSCF(11,11)/6-311++G(2df,2pd) level, has been employed for the rigid or relaxed scan. All geometrical parameters except $\phi$ were frozen or optimized in the rigid (filled symbols connected with the solid line) or relaxed scan (opened symbols connected with the dotted line), respectively. The gray-shaded area represents the local minimum in the vertical transition (see the text). The minimum energy of the $D_1$ state optimized by itself at the same level of theory ($\phi \approx 90^\circ$) is denoted as a red star.

$D_0$ has been employed for the rigid or relaxed scan. All geometrical parameters except $\phi$ were frozen or optimized in the rigid scan, whereas those are continuously optimized in the relaxed scan. The PT2 correction has been made to obtain smoothly varying potential energy curves for all electronic states of $D_0$, $D_1$, and $D_2$. As mentioned above, the 2-FTA cation is planar at the global minimum of $D_0$, whereas it adopts the perpendicular geometry ($\phi = 90^\circ$) in both $D_1$ and $D_2$. The phase inversion in the quantum beat would have been nicely explained if the $D_1 \rightarrow S_1$ ionization could be invoked together with the $D_0 \rightarrow S_1$ transition as the former would show the completely inverted phase from the latter. However, from previous and present experimental and theoretical results, the $D_1$ state is most likely located at $\sim 1$ eV above the $D_0$ state, whereas the phase inversion has been found to occur at $\sim 3000$ cm$^{-1}$ (0.37 eV) above the adiabatic ionization threshold (8.07 eV)\(^{12}\) (Figure 1). Thus, it seems to be not appropriate to invoke the $D_1 \rightarrow S_1$ transition for the phase inversion observed in the current experiment. In this regard, it is quite notable that the $D_0$ state has a vertical minimum well at $\phi \approx 135^\circ$, while its global minimum adopts the planar geometry of $\phi = 0^\circ$ along the potential energy curve calculated by the rigid scan. The nonplanar vertical minimum (which is the minimum along $\phi$ from the perspective of the $D_0 \rightarrow S_1$ vertical transition) is calculated to be $\sim 0.4$ eV above the planar global minimum. Although the much-lowered local minimum could be found at $\phi \approx 180^\circ$ in the relaxed scan, it should be emphasized that the $S_1$ wavepacket is subject to the prompt vertical transition in the ionization process. The vertical transition from $S_1$ then should experience the vertical minimum of $D_0$ at $\phi \approx 135^\circ$. Namely, the nonstationary $S_1$ wavepacket (pumped by 285 nm) is only allowed to be swung along the out-of-plane dihedral angle of $\phi$, while all other geometrical parameters are frozen (see Figure S4 in the SI).

Subsequent ionization by the fs probe laser pulse is strictly confined to the instant $S_1$ geometry given by the wavepacket location on the $\phi$ coordinate according to the Franck-Condon principle, validating that the rigid scanned potential energy curve should be employed for the explanation of the experiment. Based on the ab initio potential energy curves, the experiment has been simulated by calculating the vibrational wavefunction overlap between the nonstationary time-dependent $S_1$ wavepacket and the stationary $D_0$ state as a function of pump-probe time delay, Figure 3. The rigid scanned potential energy surface has been used to reflect stationary wavefunctions at the global (planar) and vertical ($\phi \approx 135^\circ$) minima. Most interestingly, the phase inversion could be perfectly reproduced by the simulation. And yet, the probe energy of $\sim 6000$ cm$^{-1}$ above the adiabatic ionization threshold should be required for the observation of the clear phase inversion when $ab$ initio potential energy curve is used without modification. This suggests that the $D_0$ vertical minimum energy might have been overestimated in the calculations. For the better match with the experiment, the model potential energy curve composed of two harmonic functions is employed with the variable energy gap ($\Delta E$) between their distinct minima located at $\phi = 0^\circ$ and $\Delta \phi$. Two parameters are then varied until the simulation fits to the experiment to give $\Delta E = 2000$ cm$^{-1}$ and $\Delta \phi = 135^\circ$ (see Figure S5 in the SI).

Actually, this double-well $D_0$ potential energy curve may be helpful to resolve the previous disputed issue of the photoelecton spectroscopic results on thioanisole and 2-FTA regarding the possible existence of the conformational

![Figure 2](image_url)  
**Figure 2.** Calculated potential energy curves of 2-FTA in the ground and excited cationic states of $D_0$ (black), $D_1$ (red), or $D_2$ (blue) obtained at the level of SA4-CASSCF(11,11) with the single-state PT2 correction. All geometrical parameters except $\phi$ were frozen or optimized in the rigid (filled symbols connected with the solid line) or relaxed scan (opened symbols connected with the dotted line), respectively. The gray-shaded area represents the local minimum in the vertical transition (see the text). The minimum energy of the $D_1$ state optimized by itself at the same level of theory ($\phi \approx 90^\circ$) is denoted as a red star.

![Figure 3](image_url)  
**Figure 3.** Simulations (solid lines) are overlaid with the experimental residuals after subtracting the exponential decay function of $\tau \approx 500$ ps (dots). Simulations are based on the rigid scanned potential energy curve of $D_0$ calculated at the SS-CASPT2 level. The probe laser wavelength used in the experiment is denoted for each transient, whereas the $D_0 \rightarrow S_1$ ionization energy (above the adiabatic ionization threshold) used in each simulation ($E_{simul}$) is shown within parentheses (see the text).
isomer of the perpendicular geometry in S₀. In the previously reported photoelectron spectrum of 2-FTA,⁵ the broad D₀ ← S₀ band is accompanied with the significant shoulder in the high-binding-energy region. This shoulder could be now ascribed to the vertical minimum of D₀, which is accessed by the vertical transition. The growth of the shoulder intensity with increasing temperature had been previously reported in the photoelectron spectra of thioanisole.⁴⁵,⁴⁸ This was attributed to the increase of the relative population of the perpendicular conformational isomer of thioanisole in S₀. And yet, it might be interpreted now to be due to the more excitation of the S₀ torsional modes at the higher temperature yet, it might be interpreted now to be due to the more excitation of the S₀ torsional modes at the higher temperature. The beating frequencies are successfully reproduced by the fs pump (285 nm) laser pulse prepares the vertical transition. The growth of the shoulder intensity with increasing temperature had been previously reported in the photoelectron spectroscopic results in the literature.

Conclusions

In this work, the excited-state (S₁) molecular structure, in terms of the molecular planarity, has been precisely determined from the analysis of the fs wavepacket dynamics. Coherent excitation of the out-of-plane torsional progression bands (S₁) by the fs pump (285 nm) laser pulse prepares the nonstationary wavepacket, giving rise to the clear-cut quantum beat in the parent ion transient obtained by the pump–probe scheme. The beating frequencies are successfully reproduced by the S₁ double-well model potential, giving the nonplanar geometry of 2-FTA with the out-of-plane dihedral angle (φ) of 51° at the zero-point level. This nearly perpendicular molecular structure in S₁ should open the great opportunity to explore the otherwise blind unexplored aspect of the conical intersection dynamics. Intriguingly, the amplitudes and phases of the fs wavepacket interferograms are strongly modified as the ionization (probe) energy is changed. With the aid of ab initio calculations of the cationic states, it has been found that the 2-FTA cation (D₀) adopts the planar geometry at the global minimum, whereas it has a vertical minimum at φ ≈ 135° as the vertical ionization cross section is monitored along the dihedral angle and all of the other geometrical parameters except φ are frozen. The experiment is quite well reproduced by the simulation, especially when the double-well model potential curve is energetically modified (toward the lower location of the vertical minimum with respect to the global minimum in D₀) from ab initio values. The present experimental finding also shed new light on the long-disputed interpretation of the photoelectron spectroscopic results in the literature.

Associated Content

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpca.1c04976.

Simulation details, full time-resolved ion yield data of 2-fluorothioanisole, spectra at different pump wavelengths, visual aid graphics for the theoretical interpretation, and quantum beat simulation using two harmonic potential functions (PDF)

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Notes

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References


