# **Chemical Science**

# EDGE ARTICLE



Cite this: Chem. Sci., 2024, 15, 11507

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 15th April 2024 Accepted 16th June 2024 DOI: 10.1039/d4sc02463j

rsc.li/chemical-science

## Introduction

Plasmonic nanocavities allow the localization and concentration of light into subnanoscale volumes and enable the generation of strong light–matter coupling.<sup>1-3</sup> This exceptional ability provides a novel platform for access to biosensing, quantum chemistry and materials science.<sup>4-7</sup> In particular, plasmonic nanocavities have been demonstrated to exhibit promising applications in mediating chemical reactions.<sup>8-10</sup> In theory, vibrational energy relaxation (VER) is fundamentally important for chemical reaction dynamics owing to its intrinsic connection with energy dissipation at reaction active sites and chromophores, as well as its ability to govern reaction coordinates.<sup>11</sup>–<sup>15</sup> Although plasmon-mediated chemical reactions (PMCRs) have achieved important milestones, $16,17$  the underlying effects of plasmons on the VER processes of interfacial molecules have largely not been explored.

Currently, several mechanisms for PMCRs have been proposed, including electromagnetic fields,<sup>18</sup> hot electrons

# Local electric field in nanocavities dictates the vibrational relaxation dynamics of interfacial molecules†

Xiaoxuan Zh[en](http://orcid.org/0000-0002-3286-5258)g[,](http://orcid.org/0009-0003-3257-3110) <sup>D</sup><sup>a</sup> Quanbing Pei,<sup>a</sup> Junjun Tan,\*<sup>b</sup> Shiyu Bai,<sup>a</sup> Yi Lu[o](http://orcid.org/0000-0003-0007-0394) D<sup>\*ab</sup> and Shuji Ye **D** \*ab

Plasmonic nanocavities enable the generation of strong light–matter coupling and exhibit great potential in plasmon-mediated chemical reactions (PMCRs). Although an electric field generated by nanocavities  $(E_n)$ has recently been reported, its effect on the vibrational energy relaxation (VER) of the molecules in the nanocavities has not been explored. In this study, we reveal the impact of an electric field sensed by molecules (para-substituted thiophenol derivatives) in a nanocavity ( $E_f$ ) on VER processes by employing advanced time-resolved femtosecond sum frequency generation vibrational spectroscopy (SFG-VS) supplemented by electrochemical measurements. The magnitude of  $E_n$  is almost identical (1.0  $\pm$  0.2 V nm<sup>-1</sup>) beyond the experimental deviation while  $E_f$  varies from 0.3 V nm<sup>-1</sup> to 1.7 V nm<sup>-1</sup> depending on the substituent. An exponential correlation between  $E_f$  and the complete recovery time of the ground vibrational C=C state ( $T_2$ ) of the phenyl ring is observed. Substances with a smaller  $T_2$  are strongly correlated with the reported macroscopic chemical reactivity. This finding may aid in enriching the current understanding of PMCRs and highlights the possibility of regulating vibrational energy flow into desired reaction coordinates by using a local electric field. EDGE ARTICLE<br>
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(holes) and heat generated by excited plasmons,<sup>19</sup>–<sup>21</sup> and static electric fields. $22-24$  In the last mechanism, the nanocavity can generate a direct current (DC) electric field  $(E_n)$  to mediate chemical reactions.<sup>25-29</sup> Here,  $E_n$  is an electrostatic field resulting from the photoexcitation of metal nanoparticles which is sustained over the full course of chemical turnover, rather than the instantaneous electric field resulting from the electromagnetic field concentration provided by plasmon resonance.<sup>26</sup> Although a chemical potential contributed by plasmonic excitation was recently shown,23,29,30 direct probing of the DC electric field and its influence on molecules in the nanocavity is rather challenging.<sup>25,26,30</sup> As a result, insight into the effect of plasmonic excitation on interfacial VER processes is poorly understood. In this study, we used self-assembled monolayers (SAMs) of para-substituted thiophenol derivatives on gold films and in plasmonic nanocavities as models and revealed the impact of the electric field sensed by the molecules in the nanocavity  $(E_f)$  on the VER processes. We determined the magnitudes of  $E_n$  and  $E_f$  by investigating the bandwidth of the C=C stretching mode ( $v_{\text{C}=C}$ ) of the phenyl ring via femtosecond sum frequency generation vibrational spectroscopy (SFG-VS) supplemented by electrochemical measurements. Here, the probed molecules are located in nanocavities based on a nanoparticle-on-mirror (NPoM) approach (Fig. 1a) (labeled as NPoM-SAMs).<sup>31,32</sup> The selected aromatic thiols are well known to form good SAMs on metal surfaces and are widely adopted in photocatalysis and biomedicine.<sup>33-37</sup> The feasibility and

<sup>&</sup>quot;Hefei National Research Center for Physical Sciences at the Microscale, Department of Chemical Physics, University of Science and Technology of China, Hefei, Anhui, 230026, China. E-mail: yiluo@ustc.edu.cn; shujiye@ustc.edu.cn

<sup>&</sup>lt;sup>b</sup>Hefei National Laboratory, University of Science and Technology of China, Hefei, Anhui, 230088, China. E-mail: jjtan@ustc.edu.cn

<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4sc02463j>



Fig. 1 (a) Picture depicting the SAMs assembled in NPoMs,  $R = NO<sub>2</sub>/$ CN/CF<sub>3</sub>/COOH/NHCOCH<sub>3</sub>/OCH<sub>3</sub>/NH<sub>2</sub>/N(CH<sub>3</sub>)<sub>2</sub>. The abbreviations for the corresponding substances are 4-NTP, MBN, TFTP, MBA, 4- AATP, 4-MTP, PATP and 4-DMATP. (b) The SE-SFG spectra of 4-NTP, MBN, TFTP and MBA in the frequency range of 1500–1700 cm $^{-1}$ . (c) The SE-SFG spectra of 4-AATP, 4-MTP, PATP and 4-DMATP in the frequency range of 1500–1700 cm $^{\rm -1}$ .

superiority of using the  $C=C$  stretching mode of the phenyl ring as a probe have been confirmed by surface-enhanced Raman spectroscopy (SERS).<sup>38,39</sup>  $E_n$  is approximately 1.0  $\pm$  0.2 V nm $^{-1}$ , and  $E_{\rm f}$  varies from 0.3 V nm $^{-1}$  to 1.7 V nm $^{-1}$ . An exponential correlation between  $E_f$  and the complete recovery time of the ground vibrational C=C state  $(T_2)$  was observed. A substance with a smaller  $T_2$  shows a strong correlation with the macroscopic chemical reactivity and plasmon-molecule interactions reported for PMCRs.40,41 This work highlights the possibility of regulating vibrational energy flow into desired reaction coordinates using the local electric field in which the molecules are sensed in nanocavities. In addition, it is well known that local electric field strengths on the order of 1 V nm<sup>-1</sup> are not only responsible for the vibrational Stark effect in both single-molecule systems<sup>42</sup> and condensed phases,<sup>43</sup> but can also selectively promote, drive, and control chemical reactions<sup>44</sup> such as proton transfer<sup>45</sup> and catalytic processes,  $46-49$  for example, the catalytic synthesis of ammonia from nitrogen and water microdroplets.<sup>50,51</sup> Therefore, the findings in this work may aid in enriching the current understanding of these processes including vibrational dynamics and catalytic processes.

### Results and discussion

#### The ultrafast vibrational dynamics of the  $C=C$  stretching mode of NPoM-SAMs

We first measured the spectra and ultrafast vibrational dynamics of the  $v_{C=C}$  mode of NPoM-SAMs using femtosecond time-resolved IR pump-SFG probe measurements. Fig. 1b and c



Fig. 2 (a)  $\chi^{(2)}$  decay of the  $\nu_{C=C}$  mode of the phenyl rings of 4-NTP, MBN, TFTP and MBA. (b)  $\chi^{(2)}$  decay of the  $\nu_{C=C}$  mode of the phenyl rings of 4-AATP, 4-MTP, PATP and 4-DMATP.

show the ppp spectra of the NPoM-SAMs in the frequency range of 1500–1700 cm−<sup>1</sup> . All the peaks indicate a positive phase. The spectra in the frequency range of 1200–1800  $cm^{-1}$  are displayed in Fig. S1.† All the samples had a  $v_{C=}$  frequency of 1585  $(\pm 15)$  cm<sup>-1</sup>. Compared to those of the samples on the gold film without NPoMs (labeled as Au-SAMs) (Fig. S2†), the resonance signals of the molecules on the NPoM-SAMs are amplified with an enhancement factor  $>10^3$ . Fig. 2a and b depict the decay in the effective SFG peak strength  $(\chi^{(2)})$  of the  $v_{C=C}$  mode of thiophenol derivatives. In theory, SFG probes the surface population density of a vibrational state.<sup>52,53</sup> The excitation of  $v_{C=C}$ from the ground state  $(v_0)$  to its first vibrational state  $(v_1)$  can reduce the SFG intensity. In general, the efficiency of vibrational excitation at the interface is very low. Currently, vibrational dynamics of only a few groups, such as OH and amide modes, have been reported.<sup>52,54-56</sup> In the absence of nanocavities, the bleaching value is too small to be detected (Fig. S3†). However, in the presence of nanocavities, clear bleaching following excitation at 1580 cm<sup>-1</sup> at  $t = 0$  was observed (Fig. 2a and b) due to plasmon-enhanced vibrational excitation.<sup>57,58</sup> Following this bleaching, the  $v_1$  state relaxes to an intermediate state, and the SFG intensity gradually recovers. The vibrational dynamics of the  $v_{C=C}$  mode exhibit biexponential decays with a fast component  $(T_1)$  and a slow component  $(T_2)$ . These two components were attributed to the fast population transfer between different C=C modes  $(T_1)$  and the slow complete recovery of the ground vibrational state  $(T_2)$ , respectively.  $T_2$  is dominated by a combination of intramolecular vibrational relaxation (IVR) and nonadiabatic relaxation which contributes to the excitation of electron–hole pairs (EHPs) in gold substrates and gold nanoparticles.<sup>59,60</sup> EHPs have been proven to be the primary energy transfer channel on metal surfaces.<sup>61,62</sup> The  $T_1$ values of the different thiophenol derivatives are essentially identical at approximately 1 ps, which is consistent with the time required for rapid energy equilibrium among the delocalized  $v_{C=C}$  modes.<sup>63</sup> In contrast, significant differences in  $T_2$ values were observed across the samples (Table  $S1\dagger$ ). Specifically, the  $T_2$  values are 21.4 ( $\pm$ 1.3), 20.0 ( $\pm$ 0.4), 17.3 ( $\pm$ 0.5), 15.2  $(\pm 0.3)$ , 13.8  $(\pm 0.4)$ , 12.7  $(\pm 0.5)$ , 9.7  $(\pm 0.7)$  and 10.4  $(\pm 1.4)$  ps for

4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4-DMATP, respectively. The difference in  $T_2$  may arise from various factors, such as intermolecular interactions, molecular orientation, and the electric field.<sup>64-66</sup> The studied SAMs exhibit very similar orientations with respect to the metal surface, thus ruling out orientation as the main reason for the difference in  $T_2$ .<sup>64</sup> The effect of intermolecular interactions on  $T_2$  can also be ruled out. Earlier reports suggested that the value of  $T_2$  is strongly dependent on the  $\pi$ – $\pi$  interactions between molecules, with stronger intermolecular interactions promoting faster molecular energy dissipation.<sup>64</sup> However, the observed results contradict this hypothesis. For instance, 4-NTP, which was expected to have a faster  $T_2$  due to its strong  $\pi-\pi$  interaction, actually exhibited a longer  $T_2$  (21.4 ( $\pm$ 1.3) ps). Similarly, the presence of a –CF<sub>3</sub> group theoretically weakens  $\pi$ – $\pi$  interactions and may result in an increase in the  $T_2$  of the phenyl ring. However, these speculations in terms of intermolecular interactions contrast with our findings. Therefore, the electric field is most likely a significant factor in regulating vibrational relaxation dynamics. To verify this conjecture, we measured the magnitudes of  $E_n$  and  $E_f$  based on the DC Stark effect on the bandwidth of the  $v_{C=C}$  mode of the phenyl ring by combining SFG-VS and electrochemical measurements. Edge Article Chemical Computer Chemical C

#### Determination of the magnitudes of the electric field  $E_n$  and  $E_f$

In principle,  $E_f$  in nanocavities consists of two components, the molecular internal electric field  $(E_i)$  and  $E_n$  (eqn (1)). In the absence of nanocavities,  $E_f = E_i$ . It has been reported that  $E_i$  is linearly related to the Hammett constant  $(\sigma)$  of the parasubstituted group on the chemical moiety.<sup>67-69</sup> For simplicity of analysis, the magnitude of  $E_i$  in this work is the relative magnitude compared to that at  $\sigma = 0$ . Earlier studies indicated that the electric field can generate a vibrational Stark effect, yielding a linear correlation between the electric field and the frequency  $(\omega)^{70-73}$  or peak area (intensity)<sup>74</sup> of a vibrational mode. In addition to  $\omega$  and peak area, the bandwidth also displays a linear dependence on the electric field and provides another molecular-level descriptor for evaluating the electric field (see eqn  $(2)-(4)$ ).<sup>67,75</sup> To determine the magnitude of  $E_n$ , we



Fig. 3 (a) Schematic diagram for obtaining the magnitude of  $E_n$  based on the application of an external electric potential (V) to Au-SAMs. (b)  $\Gamma_{Au}^{\sigma}$  of Au-SAMs plotted against  $\sigma$ , and (c)  $\Gamma_{NPoM}^{\sigma}$  of NPoM-SAMs plotted against  $\sigma$ 

utilized an external electric potential (V) to increase the molecular bandwidth of the films in the absence of a nanocavity (Fig. 3a). When this bandwidth broadening of the "Au-SAMs + applied potential  $(V)$ " case equals the bandwidth broadening caused by the nanocavity, the value of the external electric potential can be used to determine the magnitude of the corresponding  $E_n$  using eqn (1)–(4). Specifically, by probing the  $v_{C}$  $C<sub>C</sub>$  bandwidth change of each Au-SAM at different applied potentials  $(\Delta\Gamma_p)$  using SFG-VS supplemented by electrochemical measurements, the dependence of  $\Delta\Gamma_p$  on V can be determined by using eqn (2). The  $E_i$  and  $E_n$  of each sample can be deduced using eqn (3) and (4), respectively.

$$
E_{\rm f} = E_{\rm n} + E_{\rm i} \tag{1}
$$

$$
k^{\sigma} = \Delta \Gamma_p^{\ \sigma} / V \tag{2}
$$

$$
E_{\rm i} = \frac{\left(\Gamma_{\rm Au}^{\sigma} - \Gamma_{\rm Au}^{\sigma=0}\right)}{k^{\sigma=0}} \bigg/ d \tag{3}
$$

$$
E_{\rm n} = \frac{(F_{\rm NPoM}^{\sigma} - F_{\rm Au}^{\sigma})}{k^{\sigma}} / d \tag{4}
$$

where  $k^{\sigma}$  is defined as the Stark bandwidth broadening rate,  $\Gamma_{\text{Au}}^{\sigma}$  is the bandwidth of the  $v_{\text{C=C}}$  mode of thiophenol derivatives in the absence of nanocavities at zero applied electric potential,  $\Gamma_{\text{NPoM}}^{\sigma}$  is the corresponding bandwidth of the samples in the presence of nanocavities, and  $d$  is the spacer thickness of SAMs in the nanocavity. The subscript  $p$  represents the change induced by V, and the superscript  $\sigma$  corresponds to the Hammett constant. According to eqn (1)–(4), determining the values of  $E_n$  and  $E_f$  requires knowledge of  $\Gamma_{Au}^{\ \ \sigma}$ ,  $\Gamma_{\rm NPoM}^{\ \sigma}$ , and  $k^{\sigma}$ .

To determine the values of  $\Gamma_{Au}^{\ \ \sigma}$  and  $\Gamma_{NPoM}^{\ \ \sigma}$ , we fit the ppp spectra of the Au-SAMs and NPoM-SAMs (Fig. 1b, c and S2†) using a standard procedure (eqn  $(S1)$ †). The fitting parameters are shown in Tables S2 and S3.† The average  $\Gamma_{Au}^{\sigma}$  values were determined to be 4.1 ( $\pm$ 0.4), 4.6 ( $\pm$ 0.1), 5.1 ( $\pm$ 0.4), 5.3 ( $\pm$ 0.4), 6.2  $(\pm 0.5)$ , 6.3  $(\pm 0.7)$ , 7.2  $(\pm 0.3)$  and 7.7  $(\pm 0.6)$  cm<sup>-1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4-DMATP, respectively. The average  $\Gamma_{\text{NPoM}}^{\sigma}$  values were determined to be 4.6 ( $\pm$ 0.2), 5.2 ( $\pm$ 0.1), 5.8 ( $\pm$ 0.2), 6.6 ( $\pm$ 0.1), 8.2 ( $\pm$ 0.6), 8.6  $(\pm 0.2)$ , 11.2 ( $\pm 0.3$ ), and 12.1 ( $\pm 0.5$ ) cm<sup>-1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4-DMATP, respectively. The error bars represent more than 5 repeated experimental results (Tables S4 and S5†). The Hammett constants of the substituent groups in 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4- DMATP are 0.78, 0.66, 0.54, 0.45, 0, −0.27, −0.66, and −0.83, respectively.<sup>76</sup> A good linear correlation between the bandwidth and Hammett constant is observed  $(\Gamma_{Au}^{\sigma} = -2.0 \times \sigma + 5.9 \text{ and})$  $\Gamma_{\text{NPoM}}^{\sigma}$  = -4.3  $\times$   $\sigma$  + 8.1) (Fig. 3b and c), indicating that the bandwidth can provide an effective descriptor for evaluating the electric field. Indeed, recent investigations on the carbonyl group of acetophenones have also demonstrated the linear dependence of the bandwidth on the Hammett constant<sup>77</sup> and potential.<sup>67</sup> Compared to those of  $\Gamma_{Au}^{\sigma}$ , the plot of  $\Gamma_{NPOM}^{\sigma}$ against the Hammett constant has a greater slope and intercept (Fig. 3b and c), revealing that the introduction of nanocavities leads to peak broadening. Accordingly, the changes in bandwidth  $(\Delta \Gamma_{\rm n} = \Gamma_{\rm NPOM}^{\sigma} - \Gamma_{\rm Au}^{\sigma})$  in the absence and presence of nanocavities are 0.5 (±0.3), 0.6 (±0.1), 0.7 (±0.3), 1.3 (±0.4), 2.0  $(\pm 0.3)$ , 2.3  $(\pm 0.7)$ , 4.0  $(\pm 0.1)$ , and 4.4  $(\pm 0.3)$  cm<sup>-1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4-DMATP, respectively. Theoretically, peak broadening can be caused by several factors, e.g., pressure,<sup>78</sup> temperature,<sup>79</sup> and the electric field.<sup>24,75,80,81</sup> The effects of pressure and temperature can be excluded because the contributions caused by pressure and temperature are expected to be similar for different substituted groups with the same fluences of input laser pulses, which is opposite to what we have observed.

The value of  $k^{\sigma}$  can be determined by measuring the dependence of the bandwidth of the Au-SAMs (eqn (2)) on the potential via electrochemical testing experiments. Fig. 4a displays the potential-dependent SFG spectra of  $v_{C=C}$  of MBN. The data for other Au-SAMs are provided in Fig. S4a.† Fig. 4b and S4b† depict the dependence of  $\Delta\Gamma_p$  on V. As the potential increases,  $\Delta\Gamma_p$  for MBN, 4-NTP, TFTP, MBA, 4-AATP, and 4-MTP linearly decreases (Fig. 4b and S4b†), while those for PATP and 4-DMATP linearly increase (Fig. S4b†). The values of  $k^{\sigma}$  are determined to be  $-0.5$  (±0.1),  $-0.6$  (±0.1),  $-0.9$  (±0.1),  $-1.4$  $(\pm 0.1)$ , −2.0 ( $\pm 0.1$ ), −1.9 ( $\pm 0.2$ ), 4.0 ( $\pm 0.4$ ), and 4.4 ( $\pm 0.9$ ) cm<sup>-1</sup> V−<sup>1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4- DMATP, respectively (Table S6†). Based on the linear relationship between  $k^{\sigma}$  and  $\sigma$  shown in Fig. S5,†  $k^{\sigma=0}$  is deduced to be 2.2 cm<sup>-1</sup> V<sup>-1</sup>. With the assumption of  $d \sim 1$  nm,<sup>32,82,83</sup>  $E_i$  is determined to be  $-0.7, -0.6, -0.5, -0.4, 0, 0.2, 0.6,$  and 0.7 V nm<sup>-1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP, and 4-DMATP, respectively. Note that an electron withdrawing group (EWG) is equivalent to applying a negative electric field, while an electron donating group (EDG) represents the application of a positive electric field. This distinction arises from the fact that an upward electric field is defined as positive. When the substituent is an EWG, it induces an upward movement of the charge of the phenyl ring, equivalent to imposing a downward (negative) electric field.<sup>68</sup> In contrast, an EDG corresponds to Chemical Science<br>
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Fig. 4 (a) Potential-dependent SFG spectra of MBN between 0.6 V and −0.6 V vs. Ag/AgCl.  $\Delta\Gamma_p$  (b) and  $\Delta\omega_p$  (c) of  $v_{C=C}$  of MBN as a function of the electric potential. (d) The  $\Delta\Gamma_{\text{n}}$  values of thiophenol derivatives plotted against  $E_f$ .



Fig. 5 (a)  $E_f$  plotted against the  $T_2$  of thiophenol derivatives. (b) The ppp  $\chi^{(2)}$  decay of the  $\nu_{C=C}$  of 4-AATP at different potentials.

a positive electric field. Using eqn  $(4)$ ,  $E_n$  is deduced to be 1.0, 1.0, 0.8, 0.9, 1.0, 1.2, 1.0, and 1.0 V nm−<sup>1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP, and 4-DMATP, respectively. The magnitude of  $E_n$  for all the studied samples is almost identical  $(1.0 \pm 0.2 \text{ V nm}^{-1})$  beyond the experimental deviation. The magnitude of  $E_n$  can be further confirmed by the results determined using Stark shift (shown in the  $ESI<sup>†</sup>$ ), which is an acknowledged tool for quantifying electric fields.<sup>73,84</sup> As a result, the  $E_f$  values of the different substituents were deduced to be 0.3, 0.4, 0.5, 0.6, 1.0, 1.2, 1.6 and 1.7 V nm<sup>-1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4-DMATP, respectively (Fig. 4d). The consistency of the electric field obtained through the peak shift and peak bandwidth provides robust support for the effectiveness of employing the bandwidth as a metric for electric field measurements.

#### The correlation between  $E_f$  and relaxation time  $T_2$

We can now establish a correlation between  $E_f$  and  $T_2$  (Fig. 5a). Evidently,  $T_2$  exhibits an exponential dependence on  $E_f$ . A larger  $E_f$  corresponds to faster vibrational relaxation dynamics, while a smaller  $E_f$  corresponds to longer vibrational relaxation lifetimes. This correlation was further supported by the results obtained by applying a potential directly to 4-AATP (Fig. 5b).  $T_2$ is 10.8 ps and 15.6 ps for  $E_{\rm f}$   $=$  1.5 and 0.5 V nm $^{-1}$ , respectively.  $E_{\rm f}$ is related to the molecule–metal coupling. A larger  $E_f$  facilitates more efficient metal-molecule electron transfer and leads to faster transfer of molecular vibrational energy to the metal substrate, resulting in a shorter  $T_2$ . Accordingly, a substance with a smaller  $T_2$  is expected to exhibit stronger reactivity because of its enhanced coupling to the metal. Indeed, this has been supported by some recent reports associated with PMCRs. For example, the  $H_2$  production rate of 4-MTP in plasmonmediated hydrogenolysis reactions was approximately 60 times greater than that of 4-NTP.<sup>40</sup> The plasmon-mediated methyl dehydrogenation occurrence rate of 4-methylbenzenethiol (4-MBT) exhibited a more than twofold increase at +0.5 V compared to that at  $-0.5$  V.<sup>41</sup>

### Conclusion

In this study, we utilized femtosecond broadband SFG to investigate the electric field sensed by the molecules in a nanocavity  $(E_f)$  and its impact on the ultrafast dynamics of para-substituted thiophenol molecules confined within nanocavities. We demonstrate that the magnitude of the electric field generated in the nanocavity  $(E_n)$  and  $E_f$  can be determined by

investigating the bandwidth of the  $C=C$  stretching mode of the phenyl ring by combining SFG-VS and electrochemical measurements.  $E_{\rm n}$  is approximately 1.0  $\pm$  0.2 V nm $^{-1}$ .  $E_{\rm f}$  is determined to be 0.3, 0.4, 0.5, 0.6, 1.0, 1.2, 1.6 and 1.7 V nm<sup>−1</sup> for 4-NTP, MBN, TFTP, MBA, 4-AATP, 4-MTP, PATP and 4- DMATP, respectively. An exponential correlation between  $E_f$  and the complete recovery time of the ground vibrational  $C=C$  state  $(T<sub>2</sub>)$  is observed. By extending this correlation to the reported macroscopic chemical reactivity  $(k_{CR})$ , a profound connection between a higher  $k_{CR}$  and a smaller  $T_2$  is found. This finding highlights the possibility of regulating vibrational energy flow into a desired reaction coordinate by introducing rationally designed molecular functionalities within spatially confined nanoscale environments. A molecular-level perspective on the relationship between  $k_{CR}$ ,  $E_f$  and  $T_2$  may aid in enriching the current understanding of PMCRs. Edge Article Common access Article Common Access Article is limited on 2022. The state of the Southern State of

### Methods

### Materials and sample preparation

4-(Triuoromethyl)-thiophenol (TFTP, with a purity >98%), 4 acetamidothiophenol (4-AATP, with a purity >95%) and 4 methoxythiophenol (4-MTP, with a purity >98%) were purchased from Shanghai Macklin Biochemical Technology Co., Ltd. 4-Nitrothiophenol (4-NTP, with a purity >95%) and paminothiophenol (PATP, with a purity >98%) were obtained from Aladdin. 4-Mercaptobenzonitrile (MBN, with a purity >97%) was obtained from Shanghai Haohong Scientific Co., Ltd. P-mercaptobenzoic acid (MBA, with a purity >90%) was purchased from Shanghai Acmec Biochemical Co., Ltd. 4- Dimethylamino-thiophenol (4-DMATP, with a purity >98%) was purchased from Shanghai Canspec S&T Co., Ltd. Ethanol and nhexane were purchased from Sinopharm Chemical Reagent Co., Ltd. Ultrapure deionized (DI) water was purified by using a Milli-Q reference system. All the chemicals were used as received.

The substituents in 4-NTP, MBN, TFTP and MBA are EWGs – NO2, –CN, –CF3 and –COOH, respectively, while 4-AATP, 4-MTP, PATP and 4-DMATP are substituted with the EDGs –NHCOCH<sub>3</sub>, –OCH<sub>3</sub>, –NH<sub>2</sub> and –N(CH<sub>3</sub>)<sub>2</sub>, respectively. The specific sample preparation method is described in the ESI.†

### SFG-VS experiments

All SFG experiments were carried out using a femtosecond timeresolved SFG-VS system. Detailed information about the instrument parameters was provided in our previous studies.52,54 For a IR pump- SFG probe time-resolved SFG instrument, the pulse energies of the pump IR and probe IR pulses were  $\sim$ 12 µJ and 11 µJ at  $\sim$ 1580 cm<sup>−1</sup>, respectively. The pulse energy of vis was  $0.4 \mu$ J. The specific spectral fitting procedures and data analysis have been presented in our publications<sup>56,85</sup> and are included in the ESI<sup>†</sup> for reference.

#### Electrochemical measurements

Electrochemical measurements were performed using a CHI660e workstation. The working electrode was a Au-SAM

sample with a wire in contact with the Au film, the reference electrode was Ag/AgCl (in 3 M KCl), and the counter electrode was a Pt wire electrode. The electrolyte was a 0.3 M KCl solution. All potentials in the paper were relative to Ag/AgCl/3 M KCl, and a schematic diagram of the device is shown in Fig. S6.† In situ SFG measurements of Au-SAMs were performed over a potential range of −0.6 to 0.6 V to prevent desorption of thiol groups with a potential interval of 0.1 V. Fig. S7† illustrates the cyclic voltammetry curve with MBN as a typical example.

# Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

## Author contributions

X. X. Zheng: conceptualization, investigation, formal analysis, visualization, writing - original draft preparation, writing reviewing and editing; Q. B. Pei: investigation, writing – reviewing and editing; J. J. Tan: software, funding acquisition, project administration, data curation, writing – reviewing and editing; S. Y. Bai: writing – reviewing and editing; Y. Luo: conceptualization, funding acquisition, manuscript drafting; S. J. Ye: conceptualization, funding acquisition, supervision, data analysis, manuscript drafting, writing - reviewing and editing.

# Conflicts of interest

The authors declare no competing financial interest.

## Acknowledgements

This work was supported by the National Natural Science Foundation of China (92250306, 21925302, and 22273094), the Innovation Program for Quantum Science and Technology (2021ZD0303303), the Strategic Priority Research Program of the Chinese Academy of Sciences (XDB0450202), and the Anhui Initiative in Quantum Information Technologies (AHY090000).

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