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Controlling optoelectronic properties through protonation with π -extended triphenodioxazine diimides†

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We present a combined approach integrating chemical synthesis, computational methods and advanced ultrafast spectroscopy to explore the properties of triphenodioxazine diimides (TPDODI). The TPDODI derivative shows significant redshifted absorption ($\lambda_{\text{max}} = 556$ nm) and emission ($\lambda_{\text{max}} = 569$ nm) compared to other non-diimide TPDO derivatives. Protonation of the TPDODI leads to further redshifts in absorption ($\lambda_{\text{max}} = 638$ nm and 715 nm for mono- and diprotonated states, respectively), with the diprotonated form absorbing up to 800 nm. However, protonation also triggers competitive nonradiative decay processes, confirmed by transient absorption spectroscopy and linked to the Energy Gap Law via high-frequency molecular vibrations detected by two-dimensional electronic spectroscopy. Computational analysis supports these findings, particularly in highlighting the enhanced electron affinity of the monoprotonated species (LUMO = -3.61 eV vs. -4.98 eV for the neutral and monoprotonated forms, respectively). These results underscore the versatility of TPDODI for optoelectronic applications, providing key insights into the fine-tuning of n-type semiconductors, catalysts, and other advanced materials.

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Introduction

Over the past two decades, studies on organic molecules for optoelectronic applications, including organic photovoltaics (OPVs) and light-emitting diodes (OLEDs), have grown rapidly. The rising interest in these materials stems from their inherent advantages, including solution processability, flexibility and the potential to mould their photophysical properties by structural

fine-tuning.^{1,2} These characteristics make organic molecules highly attractive for use in a wide range of technologies from energy harvesting to light emission.

In this context, industrially producible triphenodioxazine (TPDO)³ derivatives have emerged as promising candidates due to their remarkable versatility. These derivatives have been successfully applied across a wide range of technologies, including air- and photo-stable pigments,^{4,5} OPVs,^{4–6} dye-sensitised solar cells,^{4,7} air-stable organic field effect transistors,^{8–10} liquid crystal luminescent devices,^{7,11} stimuli responsive materials,¹² and organic thin film transistors^{8,9,13} including those for toxin sensing.^{14,15} Of particular interest, TPDO derivatives display n-type semiconducting properties, which are relatively rare in organic molecules due to an inherent instability of organic n-type semiconductors which, among other methods, has been addressed by the incorporation of TPDO derivatives.^{4,8,9} This makes them attractive alternatives to traditional organic n-type semiconductors such as perylene diimides, naphthalene diimides, C₆₀ and fluorinated oligothiophenes.^{16–21}

Despite the promising optoelectronic properties of TPDO derivatives, there is still room to enhance their electron affinity and tuneability, which is crucial for optimizing their performance in semiconductors and other applications.²² One successful approach to achieving this is the introduction of

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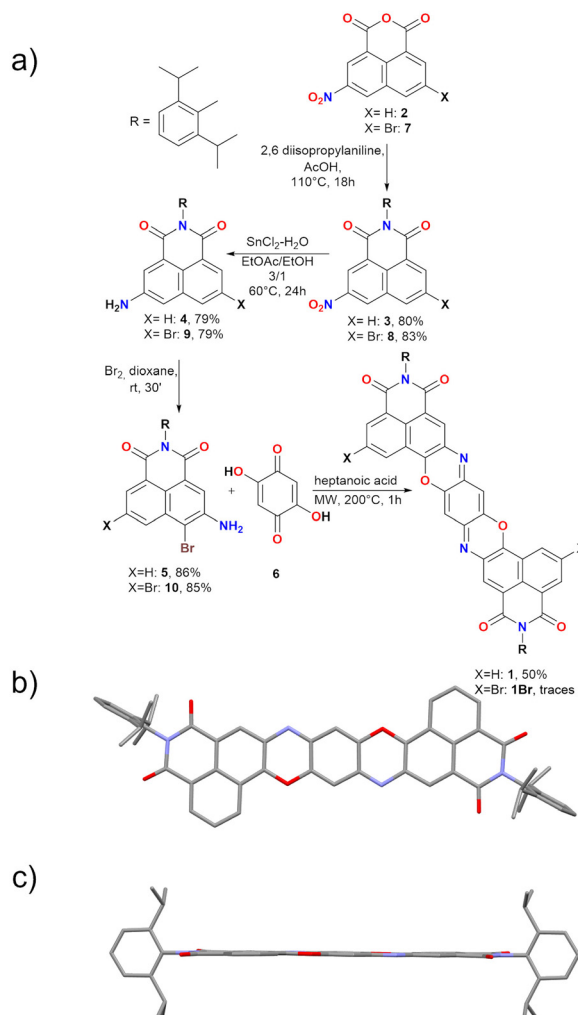
electron-withdrawing groups on the TPDO scaffold, which lowers the LUMO levels and enhances the electron affinity.^{8,23} Additionally, the conversion of TPDO to its diimide form has been shown to improve both optical and electronic properties, resulting in stronger visible absorption and higher emission quantum yields.²³ Recent studies on triphenodioxazine diimides (TPDODIs) have confirmed that the incorporation of the imide group enhances electron affinity, making these systems even more appealing for optoelectronic applications.²³ However, despite these advances, the effects of protonation on the optical and electronic properties of TPDOs and TPDODIs remain largely unexplored.

Building on this, in this paper we report the first synthesis of a π -extended TPDODI presenting intense visible absorption and high emission quantum yield. We investigate the effect of mono- and double protonation on these derivatives, revealing significantly red-shifted absorption and emission associated with a dramatic decrease of the LUMO level of the molecule. An in-depth experimental and computational analysis reveals the optical and vibrational properties of the neutral and protonated states of the molecule, setting an important milestone for the future development of these materials.

Results and discussion

Synthesis and structural characterisation

In general, the most common route for the preparation of TPDO derivatives involves the reaction of 2,5-dihydroxy-1,4-benzoquinones with 1-aminophenols in acidic conditions, resulting in condensation reactions that lead to the formation of the heteroaromatic system. Aiming at the synthesis of TPDODI system **1**, the preparation of 5-amino-6-hydroxy naphthalene imide would be required, which is however a highly challenging multi-step procedure requiring noticeable synthetic effort. To avoid this, a similar reactivity could plausibly be obtained by using a 5-amino-6-bromo naphthalene monoamide (**5**, Scheme 1). In fact, in presence of 2,5-dihydroxy-1,4-benzoquinone and acidic conditions, the free amine group on **5** would be able to react with the carbonyl moiety in **6**, resulting in formation of an imine group. At this point, exploiting the electron-depleted nature of naphthalene imide systems and the high effective molarity arising from proximity, a nucleophilic aromatic substitution with release of HBr could take place, resulting in the formation of the final C–O bond. This would be a great advantage since it could lead to a facile synthesis of π -extended systems, avoiding the complexity related to the synthesis of amino hydroxy derivatives. Based on these assumptions, the synthesis of **5** was started by treating **2** with 2,6 diisopropyl aniline in the presence of acetic acid at 110 °C for 18 h, resulting in the formation of the corresponding naphthalene monoimide **3**. Subsequently, **3** was reduced in the presence of $\text{SnCl}_2 \cdot 2\text{H}_2\text{O}$ in EtOAc/EtOH at 60 °C to afford the free amine derivative **4** in a 79% yield. Treatment of **4** with bromine in dioxane smoothly afforded derivative **5** in an 86% yield. At this point, derivative **5** was suspended in heptanoic acid along with **6** followed by microwave (MW) irradiation at 200 °C for 1 h,



Scheme 1 (a) Synthesis of derivative **1**, R = diisopropylphenyl; (b) top view and (c) side view of crystal structure of **1**. Crystals obtained from slow diffusion of hexane in CHCl_3 . Space group: $P2_1/n$. Atom colours: O: red, N: blue, C: grey. Hydrogens omitted for clarity. CCDC 2383220.[†]

resulting in the formation of desired **1** as a dark red solid in a 50% yield (Scheme 1). Heptanoic acid was chosen as solvent since it ensures both the acid conditions for the imine formation and the high temperatures required for the nucleophilic aromatic substitution on the Br atom.

On the other hand, when **10** was reacted to form **1Br**, only traces of product could be obtained, suggesting reactivity also of the second bromine atom in the harsh reaction conditions, limiting characterization of **1Br** to scXRD (Table S4 and Fig. S45, ESI[†]).

Derivative **1** was completely characterised *via* NMR and high-resolution mass spectroscopy (HRMS), confirming the formation of the desired TPDODI system. Moreover, it was possible to grow crystals of sufficient quality for single crystal X-ray diffraction (scXRD, Scheme 1b and c). From the scXRD analysis, the formation of the desired structure was confirmed. The latter proved to be almost completely planar and presented a quinoidal structure in the central ring, highlighted by marked bond alternation, in line with similar non-diimide

derivatives.^{8,24} No significant π -stacking interactions were visible between derivatives, mostly due to the bulky nature of the diisopropyl-phenyl substituents which disrupt significantly any interaction. Also, no interactions were visible at the sp^2 nitrogen atoms which is plausibly related to the steric hindrance arising from the neighbouring phenyl ring. At this point, to understand if some nucleophilic character was present on the sp^2 nitrogen, an NMR experiment was performed on **1** to assess if this system could be protonated by acids resulting in the mono- and possibly doubly protonated form (Fig. 1), which are expected to present a very high electron affinity.

From these experiments it is possible to see that the diagnostic H_a proton associated with the quinoidal system undergoes a marked downfield shift upon addition of acids. Particularly, addition of TFA in $CDCl_3$ (0.2 M) results in an immediate colour change from bright red to deep blue, associated with a downfield shift of 0.83 ppm. Recording the same spectrum in pure TFA does not significantly change the chemical shift of the peaks, suggesting quantitative protonation in both cases. The symmetric nature of the obtained spectrum however does not allow to discern between a monoprotonation with a fast interchange between the two nitrogen atoms or the doubly protonated form. Diffraction of crystals obtained in presence of TFA also did not provide satisfactory answers to this question (see ESI[†]). As such, to make this point clear, a further experiment was performed by adding 1 drop of D_2SO_4 to a TFA solution of **1**. The resulting NMR spectrum presents an even more marked downfield shift for H_a (0.54 ppm), suggesting that a second protonation is occurring in this case. To support this, the absorption spectrum of **1** was measured in solvent systems of different acidity and is shown in Fig. 2a and Fig. S33 (ESI[†]). First, in $CHCl_3$ **1** presents a structured absorption with a $\lambda_{max} = 556$ nm ($\varepsilon = 6.5 \times 10^4 M^{-1} cm^{-1}$) associated with an emission ($\lambda_{em,max} = 569$ nm, Fig. S28, ESI[†]) with a small Stokes

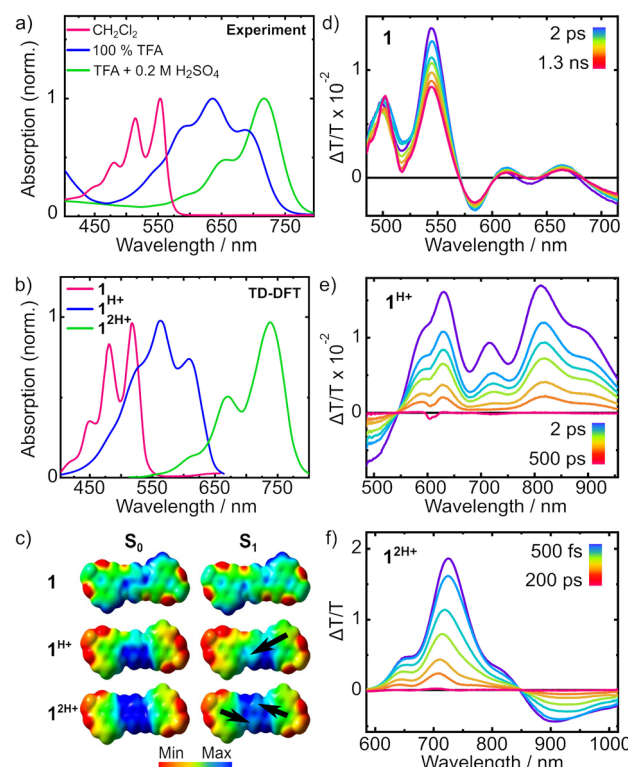


Fig. 2 (a) Normalized absorption of **1** in CH_2Cl_2 , pure TFA (1^{H^+}) and TFA with 0.2 M H_2SO_4 (1^{2H^+}). (b) Calculated absorption spectra of **1**, 1^{H^+} and 1^{2H^+} from TD-DFT. (c) Calculated electrostatic potentials for neutral **1**, 1^{H^+} and 1^{2H^+} . The colour scale indicates the full range of electrostatic potential, with the range being $-0.02e$ to $0.02e$ for **1**, $0.00e$ to $0.15e$ for 1^{H^+} and $0.10e$ to $0.26e$ for 1^{2H^+} . As expected, the scale is shifted to increasingly positive values with more protonation. (d)–(f) Transient absorption spectra for **1** (d), **1** in TFA (1^{H^+} , (e)) and **1** in H_2SO_4 (1^{2H^+} , (f)) at selected timepoints within the range given by the respective colour bar for each spectrum.

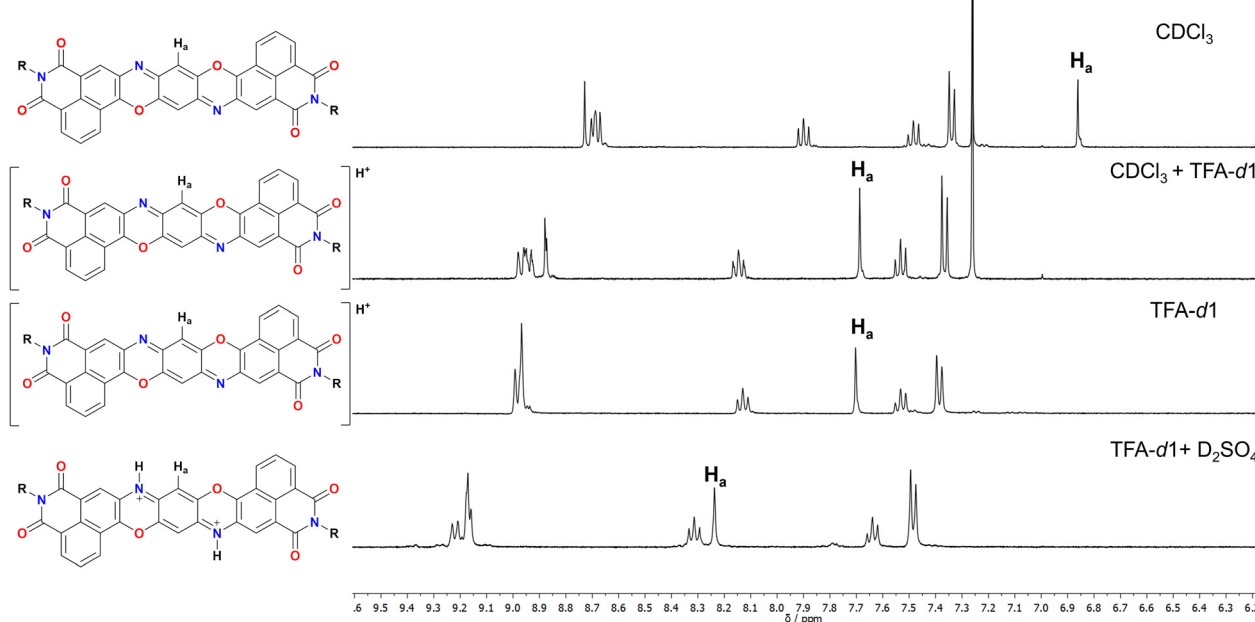


Fig. 1 Detail of the aromatic part of **1** in different solvent systems.

shift consistent with its highly rigid structure. Interestingly, this emission occurs with a high quantum yield (QY) of 70%. Upon addition of increasing amounts of TFA to solutions of **1** in CH_2Cl_2 a new set of redshifted absorption bands appears ($\lambda_{\text{max}} = 638$ nm, Fig. S29, ESI[†]), suggesting the formation of the monoprotonated form with complete conversion at *ca.* 0.2 M concentration of TFA. A similar titration performed with H_2SO_4 in THF afforded a similar result (Fig. S30 and S33, ESI[†]). Moreover, the absorption spectrum in pure TFA ($\lambda_{\text{max}} = 629$ nm) is very similar to those in $\text{THF} + \text{H}_2\text{SO}_4$ ($\lambda_{\text{max}} = 620$ nm), $\text{CH}_2\text{Cl}_2 + \text{TFA}$ ($\lambda_{\text{max}} = 638$ nm) and $\text{CHCl}_3 + \text{TFA}$ ($\lambda_{\text{max}} = 638$ nm), thus ruling out important solvatochromic effects (Fig. S33, ESI[†]). Interestingly, the spectra of the neutral and monoprotonated derivatives are very similar to non-imide derivatives²⁴ but are redshifted by *ca.* 36 nm corresponding to lowering of 0.16 eV in the transition energy, plausibly due to the enhanced conjugation arising from the extended diimide system. Also, compared to the non-diimide derivatives, in the case of **1**, a 10-fold higher concentration of TFA is required to form quantitatively **1**^{H+}, suggesting that the electron depleted diimide systems are indeed decreasing the basic nature of the pyridinic nitrogen.²⁴ Interestingly, a weak (QY < 1%) red emission ($\lambda_{\text{em,max}} = 738$ nm) was associated with **1**^{H+} suggesting a great enhancement of the nonradiative decay deactivation pathways upon protonation (Fig. S31, ESI[†]). In line with the previously discussed NMR experiments, also in this case, dissolving **1** in a TFA solution containing 0.2 M H_2SO_4 resulted in the formation of a new set of redshifted absorption bands ($\lambda_{\text{max}} = 715$ nm) consistent with the formation of **1**^{2H+}.

To estimate the HOMO and LUMO levels for derivatives **1** and **1**^{H+}, cyclic voltammetry experiments were carried out in CH_2Cl_2 (Fig. S36 and S37, ESI[†]). In the case of **1**, an irreversible oxidation was visible at +0.95 V *vs.* Fc/Fc^+ , while two reversible reductions were detected at -1.19 V and -1.62 V *vs.* Fc/Fc^+ respectively. These values are associated with a HOMO level of -5.75 eV and a LUMO of -3.61 eV *vs.* Fc/Fc^+ which are consistent with TPDO derivatives bearing electron-withdrawing groups such as CF_3 ,⁸ and differently positioned diimides.²³ Interestingly, upon addition of TFA (0.2 M) the two reversible reduction peaks appear at much more positive potentials (Fig. S37, +0.18 V and -0.23 V *vs.* Fc/Fc^+ , ESI[†]) which are consistent with a dramatic decrease of the LUMO level (-4.98 eV *vs.* Fc/Fc^+). This suggests that protonation results in great enhancement in the electron affinity for TPDO diimides. This result is consistent with the trend observed in calculated HOMO and LUMO levels (Fig. S38 and Table S1, ESI[†]), the computational details of which are discussed in detail below. The theoretical results also reveal decreasing HOMO and LUMO levels and a narrowing HOMO-LUMO gap with increasing protonation.

Electronic structure characterisation

To gain insights into the effect of protonation on the electronic structure of **1**, computational studies were performed by means of density functional theory (DFT) for the ground state (GS), and time-dependent density functional theory (TD-DFT) for the first excited state employing long-range corrected CAM-B3LYP²⁵ hybrid functional together with cc-pVQZ²⁶ basis set as implemented in

the Gaussian16²⁷ software package. Solvent effects were treated implicitly through the polarisable continuum model (PCM).²⁸ GS optimisation of the three systems shows that the structure is not affected upon different degrees of protonation retaining the aromatic core planar structure, in line with crystal structures obtained in presence of TFA (Fig. S44, ESI[†]). Calculation of vertical excitation energies reveals that the S_1 bright state is characterised by the HOMO-LUMO transition (Fig. S38 and Table S2, ESI[†]) and, in agreement with experimental trend, we observe a progressive redshift of the absorption peak with increasing protonation: in particular, $\lambda_{\text{abs}} = 499$ nm for the neutral system, $\lambda_{\text{abs}} = 577$ nm for the monoprotonated system and $\lambda_{\text{abs}} = 767$ nm for the diprotonated system. Computed vibronic spectra of the three systems are shown in Fig. 2b. To reproduce the vibronic fine structure, we made use of the iSpectron²⁹ code providing the GS equilibrium geometry together with its normal modes and energy gradient of the S_1 excited state at the Franck-Condon point. We assume that potential energy surfaces (PESSs) of the electronic states are quadratic in the coordinates with different equilibrium geometries. Moreover, different electronic states are described by the same normal modes and frequencies which are calculated only for the GS. This procedure is referred to as the vertical gradient (VG)³⁰ approach in the literature. The simulated spectra show good agreement with experiments, although it was necessary to scale the gradients to reproduce the exact line shape (computed vibronic spectra without scaling factors are shown in Fig. S39, ESI[†]). We note that in the three systems, the vibronic structure is given by the activation of vibrational modes between 1300–1500 cm^{-1} assigned to stretching deformations of the aromatic core. In the neutral and diprotonated systems, the dominant mode is characterised by a symmetric stretching while in the monoprotonated system an anti-symmetric stretch is the dominant mode (Fig. S40, ESI[†]).

Electrostatic potentials for **1**, **1**^{H+} and **1**^{2H+}, on both S_0 and S_1 are shown in Fig. 2c. The neutral molecule **1** exhibits weak electron density deficiency around the central ring of the molecule in the GS. Protonation leads to delocalization of a positive charge over the central ring, asymmetrically in the case of **1**^{H+} and symmetrically in the case of **1**^{2H+}. Due to the positive charge residing on the central ring, a donor-acceptor-donor motif arises that is increasingly prevalent with increasing protonation. In all three molecules, excitation to S_1 results in a partial charge displacement towards the centre (indicated by black arrows in Fig. 2c). Both **1** and **1**^{2H+} exhibit a symmetric shift on excitation due to the symmetry in their structure. Conversely, the asymmetry in **1**^{H+} extends to asymmetric electrostatic potentials on both S_0 and S_1 , giving rise to a non-zero dipole moment in S_1 .

Characterisation of excited state dynamics

To characterise the optical properties of the three systems, the photoinduced dynamics following excitation to S_1 were investigated by transient absorption (TA) spectroscopy. The transient differential transmission ($\Delta T/T$) spectra at selected probe delays are shown in Fig. 2 for **1** (Fig. 2d), **1**^{H+} (Fig. 2e) and **1**^{2H+} (Fig. 2f). For all three systems we observe large ground state



bleach (GSB) features at wavelengths corresponding to the linear absorption spectra in Fig. 2a. Due to the probe spectral range (490–710 nm), we were able to detect only the two lowest-lying vibronic peaks in the absorption spectrum for **1** (Fig. 2a), giving GSB features at 500 nm and 550 nm (Fig. 2d). At longer probe wavelengths, stimulated emission (SE) bands can be observed at 610 nm and 665 nm, though their amplitude is depleted by a superimposed negative photoinduced absorption (PA) band. The transient spectrum features isosbestic points at 570 nm and 600 nm, which lie on the $\Delta T/T = 0$ baseline, indicating that the spectrum observed at early times evolves directly to zero. This reveals that the S_1 state relaxes directly to the GS and therefore the timescale of its decay denotes the timescale of $S_1 \rightarrow S_0$ relaxation. All spectral features decay slowly and persist beyond the 1.3 ns pump–probe delay available in the measurement.

For $\mathbf{1}^{1\text{H}^+}$ (Fig. 2e), we observe GSB features at 590 nm and 630 nm. A positive band at 710 nm comprises a superposition of GSB and SE contributions, while the remaining SE peaks appear at 805 nm and 910 nm. We identify an isosbestic point on the baseline for $\mathbf{1}^{1\text{H}^+}$ at 550 nm, confirming that the S_1 state of the monoprotonated molecule also relaxes directly to the ground state without branching photophysical processes. In this case, however, the ground state is entirely repopulated before 500 ps, as observed by the recovery of the signal to zero. For $\mathbf{1}^{2\text{H}^+}$ (Fig. 2f), the highest-lying vibronic peak lies outside of our probe window, so we observe GSB bands at 650 nm and 725 nm. The spectrum also reveals a strongly quenched emission manifesting as only a shoulder of SE at 830 nm, otherwise masked by a PA that spans from 850 nm to longer wavelengths. Again, the isosbestic point at 850 nm demonstrates that the S_1 state of $\mathbf{1}^{2\text{H}^+}$ relaxes directly back to the ground state.

To examine the lifetime of the S_1 state in each system, a global fit was performed for each time-resolved spectrum. The kinetic traces that show the decay of the S_1 state, together with monoexponential fits to their evolution, are presented in Fig. 3. From the monoexponential fits, a lifetime that describes the relaxation from S_1 to S_0 was extracted and is also reported in Fig. 3. The fit for the neutral molecule **1** revealed a time constant 1.2 ± 0.13 ns. In $\mathbf{1}^{1\text{H}^+}$, the $S_1 \rightarrow S_0$ relaxation is accelerated to 92 ± 0.7 ps, more than an order of magnitude faster. Meanwhile, the diprotonated form $\mathbf{1}^{2\text{H}^+}$ relaxes on a timescale of 27 ± 0.2 ps, accelerated a further threefold.

The coupling of electronic transitions to high-frequency ($1000\text{--}1600\text{ cm}^{-1}$) molecular vibrations, commonly carbon–carbon stretching modes, is usually responsible for accelerating the nonradiative depopulation of the fluorescent state in organic molecules.³¹ This generally accepted principle prompted us to investigate whether molecular vibrations that are coupled to the $S_1 \rightarrow S_0$ electronic transition play a role in its altered dynamics. The ground state and excited state vibrational spectra of each molecule were therefore calculated using the same level of theory previously described. The ground state spectra can be found in the ESI† (Fig. S41) and the S_1 Huang–Rhys spectra are shown in Fig. 4. All calculated spectra are generally in good agreement with one another and are also consistent with experimental

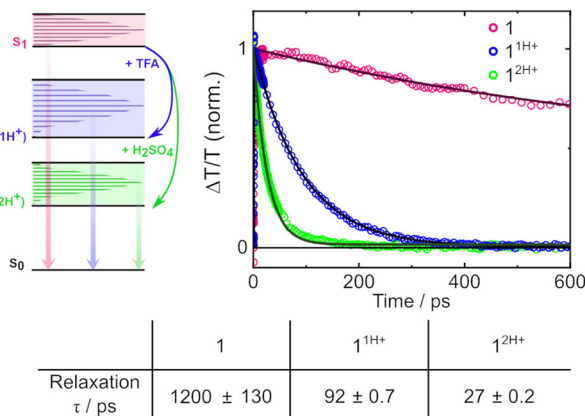


Fig. 3 Energy level diagram for **1** under neutral conditions, in TFA and in TFA + 0.2 M H_2SO_4 where vertical arrows depict the probed transition of relaxation from S_1 to S_0 , alongside the dynamics of this process in each molecule and the associated lifetimes from global fitting to TA spectra in Fig. 2.

ground state spectra obtained by Raman spectroscopy (Fig. S42, ESI†). All three systems show a dense manifold of high frequency vibrations in the region $1300\text{--}1500\text{ cm}^{-1}$ on both the ground and excited state, which correspond to the same stretching modes which give rise to vibronic spectra, and the frequency of the highest intensity peak is consistent for all three molecules between $1430\text{--}1450\text{ cm}^{-1}$.

To experimentally verify the observations from theoretical vibrational analysis, two-dimensional electronic spectroscopy (2DES) was also carried out. Such experiments employ sub-15-fs broadband pulses, granting a sufficiently high time resolution to probe high-frequency vibrations up to 1500 cm^{-1} .³² Indeed, ultrashort pulses impulsively excite vibrational wavepackets, the temporal evolution of which induces an oscillating signal amplitude superimposed on the photoinduced dynamics.³³ The period of the oscillation corresponds to the pump-induced nuclear motion and can be projected onto the frequency domain using the Fourier transform to resolve the vibrational spectrum of a given electronic state. In typical TA measurements, the use of broadband excitation results in a signal integrated across the entire pump wavelength range. The outcome of this experimental condition is that dynamics (especially minor oscillations in the signal due to molecular vibrations) arising from photoexcitation at different pump wavelengths may cancel to an amplitude that is within the signal-to-noise limit of the measurement. 2DES instead allows to resolve the transient $\Delta T/T$ spectra as a function of pump wavelength as well as probe wavelength, still preserving high temporal resolution, thus enabling the sensitive detection of high frequency oscillations. The resulting 2D maps can be found in the ESI† (Fig. S43).

For each molecule, the temporal evolution of a negative signal in the 2D map was extracted by plotting the signal intensity at a specific pump and probe wavelength as a function of time. As the negative signals correspond to PA transitions following resonant excitation to S_1 , these signals arise due to excited state absorption from S_1 to a higher-lying electronic



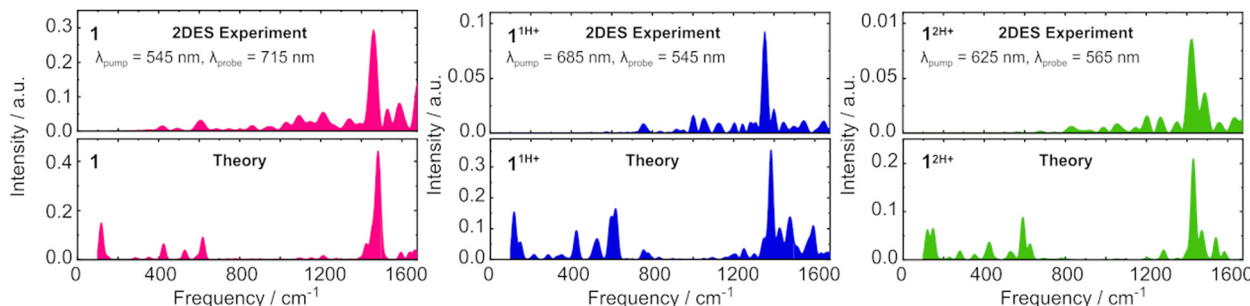


Fig. 4 Experimental impulsively excited vibrational spectra from two-dimensional electronic spectroscopy (top) by taking the Fourier transform of kinetic traces pertaining to transitions on the S_1 state and calculated Huang–Rhys spectra of each molecule on the S_1 electronic excited state (bottom).

state (S_n). Therefore, by ensuring that the negative time-dependent signal is extracted from a point on the 2D map that is sufficiently far from GSB and SE features to rule out overlap with vibrational wavepackets on the ground electronic state, the oscillatory signal for a given molecule describes pump-excited vibrational wavepackets on the S_1 state.³⁴ In each case, the Fourier transform of the time-dependent signal provides a frequency-dependent spectrum that reveals the experimental vibrational spectrum of the S_1 state. In Fig. 4, these S_1 vibrational spectra are plotted alongside the calculated Huang–Rhys spectra, $HR_k = d_k^2 \omega_k / 2\hbar$, where d_k is the horizontal displacement of the S_1 PES with respect to S_0 along the k th normal mode with frequency ω_k , obtained with iSpectron in the aforementioned VG approximation.

We note an excellent agreement between theory and experiment around $1420\text{--}1450\text{ cm}^{-1}$, arising due to the different equilibrium geometries of S_1 and S_0 along the high frequency stretching mode also responsible for the vibrational fine structure in the GS absorption spectra (Fig. S40, ESI[†]). In detail, the dominant vibrational modes from the 2DES measurement and from the theoretical approach, respectively, are $1458 \pm 10\text{ cm}^{-1}$ and 1482 cm^{-1} for **1**, $1354 \pm 10\text{ cm}^{-1}$ and 1384 cm^{-1} for 1^{1H^+} and $1426 \pm 10\text{ cm}^{-1}$ and 1428 cm^{-1} for 1^{2H^+} demonstrating a close agreement between theory and experiment. The discrepancies between experimental and theoretical results at lower frequencies are signatures of the different PES topology of the S_n state giving rise to the PA at the wavelength chosen for the analysis with respect to S_1 (Fig. S43, ESI[†]) and are thus less relevant for the present discussion. Overall, the similarity between the vibrational spectra for all three molecules confirms that the protonation does not have a dramatic effect on the vibrational energy levels on the S_1 state, nor on the ground state.

The results in Fig. 4 reveal that the acceleration of non-radiative decay with increasing protonation is a consequence only of the decreasing $S_0\text{--}S_1$ energy gap, as the frequency of the dominant vibrational mode is largely unaffected by the protonation. We confirm this by invoking the well-accepted Energy Gap Law for calculating the rate (k_{nr}) of radiationless transitions, widely used for evaluating the competing channels of luminescence and other loss pathways that contribute to S_1 decay, which states that k_{nr} accelerates as the $S_0\text{--}S_1$ energy gap decreases.³⁵ Computing the ratio of nonradiative decay rates by varying only the $S_0\text{--}S_1$ energy gap, ΔE , and using a constant

value of vibrational frequency for all three molecules, we can accurately predict the trend of accelerating nonradiative decay thus further confirming that this effect comes from the decreasing energy gap alone. In the following discussion we therefore employ an effective frequency value ω by taking an average of the three molecules' dominant vibrational modes (Fig. S40 and Table S3, ESI[†]), which is 1431 cm^{-1} according to the peaks from calculated Haung–Rhys spectra, and 1413 cm^{-1} according to the results from 2DES experiments. In order to calculate the expected acceleration of nonradiative decay with narrowing ΔE , we compare approximated decay rate ratios, calculated using ΔE and ω values according to the Energy Gap Law, to the experimentally determined nonradiative decay rates. Nonradiative decay rates are calculated from the experimental S_1 decay lifetimes in Fig. 3 and the fluorescence QY values (Table S3, ESI[†]). Because 1^{1H^+} and 1^{2H^+} both have negligible QY and are thus dominated by non-radiative decay, the k_{nr} is equal to the experimentally measured S_1 decay rate, while the high (0.7) QY of **1** means that k_{nr} is three times faster than the measured S_1 decay lifetime given in Fig. 3, according to $k = k_r + k_{\text{nr}}$ and $\text{QY} = k_r / (k_r + k_{\text{nr}})$.

Full treatment of the Energy Gap Law expression for a transition from S_1 to S_0 includes terms that consider the energy difference ΔE between the involved transition, the electronic coupling between the S_1 and S_0 states, and the relaxation energy related to the frequency of molecular vibrations. According to TD-DFT calculations, there is an $S_0\text{--}S_1$ conical intersection only for the deprotonated molecule that is, however, far too high in energy (*ca.* 2 eV from the S_1 minimum) to be accessible. We therefore deduce that all three systems are in the regime of weak electronic coupling between S_0 and S_1 and approximate that there is no major difference in electronic coupling between the three studied molecules. This approximation allows us to derive an expression for the ratio of the rates for any given two compared molecules for which the ΔE is the only variable that is unique to a given molecule. The ratio between the rate of the $S_1 \rightarrow S_0$ transition in, for example, 1^{1H^+} and 1^{2H^+} according to our approximation of the Energy Gap Law is therefore:

$$\frac{k_{2\text{H}^+}}{k_{1\text{H}^+}} = \sqrt{\frac{\Delta E_{1\text{H}^+}}{\Delta E_{2\text{H}^+}}} \cdot \frac{\exp\left(-\frac{\Delta E_{2\text{H}^+}}{\omega}\right)}{\exp\left(-\frac{\Delta E_{1\text{H}^+}}{\omega}\right)} \quad (1)$$



where k_{nH^+} is the rate of S_1 decay for $\mathbf{1}^{nH^+}$, ΔE_{nH^+} is the S_0 - S_1 energy gap for $\mathbf{1}^{nH^+}$ and ω is the effective frequency value taken as the average of the three dominant vibrational modes. The expression can also be applied to compare to the rate of nonradiative decay in the neutral form of $\mathbf{1}$, for which we denote the rate k_1 . Due to the approximation and the omission of several terms, we do not expect to accurately predict the rate ratios but to qualitatively predict a trend that is consistent with the experimental trend of nonradiative decay acceleration. We extract the ΔE values from the linear absorption spectra in Fig. 2a and obtain $\Delta E = 2.24$, 1.79 and 1.73 eV for $\mathbf{1}$, $\mathbf{1}^{1H^+}$ and $\mathbf{1}^{2H^+}$ respectively. Using an ω value of 1431 cm^{-1} from the calculated Huang–Rhys spectra, we estimate ratios for k_{2H^+}/k_{1H^+} and k_{1H^+}/k_1 of 1.4 and 14.0 , respectively. Instead, taking an ω value of 1413 cm^{-1} from the Fourier transform spectra of the 2DES measurements, we arrive at similar ratios for k_{2H^+}/k_{1H^+} and k_{1H^+}/k_1 of 1.4 and 14.5 , respectively. Finally, we compare these values with the ones that can be directly extracted from the experimental k_{nr} values (Table S3, ESI†). In this experimental case, the ratios k_{2H^+}/k_{1H^+} and k_{1H^+}/k_1 were determined to be 3.4 and 43 , respectively; our approximation thus consistently underpredicts the ratio by a factor of three. The consistency of the trend in the predicted rate ratios with the trend obtained by experimental results suggests that this is a reasonable qualitative approximation to demonstrate that the acceleration of S_1 relaxation follows the Energy Gap Law and is dominantly caused by the narrowing S_0 - S_1 energy gap. Furthermore, we approximate an acceleration of over an order of magnitude on monoprotonation and a lesser acceleration on diprotonation, consistent with experimental results. The excellent agreement between this approach when considering the theoretically determined vibrational frequencies compared to those from 2DES measurements further validates the application of the Energy Gap Law model here. The kinetic and vibrational analyses therefore characterise the novel molecule to describe its photophysical properties and justify the behaviour in terms of a well-understood physical model.

Conclusions

In this work, we have synthesized a novel π -extended TPDO diimide derivative that exhibits intense visible absorption and high fluorescence QY. In particular, the absorption was red-shifted compared to other electron depleted TPDO derivatives, plausibly due to the extended conjugation. Interestingly, this molecule demonstrates remarkable sensitivity to pH: protonation of the molecular core further shifts the absorption spectrum significantly to the near infrared (up to 800 nm for $\mathbf{1}^{2H^+}$) and lowers the LUMO levels (*ca.* -4.98 V *vs.* Fc/Fc^+ for $\mathbf{1}^{1H^+}$). Using transient absorption spectroscopy and two-dimensional electronic spectroscopy with high temporal resolution, supported by DFT-based simulations, we have shown that protonation accelerates nonradiative decay by an order of magnitude. This behaviour is consistent with the Energy Gap Law, where the narrowing of the S_0 - S_1 energy gap, rather than changes in the

vibrational spectra, accounts for the faster decay rates. The optical properties of this new organic semiconductor molecule can therefore be finely tuned *via* pH, resulting in an unprecedented level of control over its absorption characteristics. This tuneability particularly in achieving low-lying LUMO levels and visible to NIR absorption has significant potential for diverse applications of organic semiconductors, especially in the photocatalysis field where red to near-infrared light absorption is highly desirable.

Data availability

The data supporting this article have been included as part of the ESI.† Crystallographic data for $\mathbf{1Br}$ and $\mathbf{1}$ have been deposited at the CCDC under 2383219 and 2383220-2383224 deposition codes.†

Conflicts of interest

There are no conflicts to declare.

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