

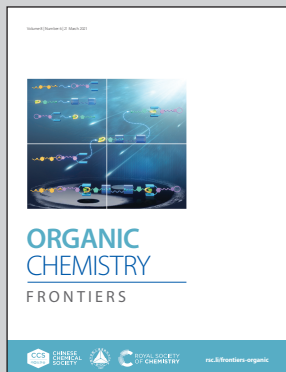
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Hole-mediated photoredox catalysis: tris(*p*-substituted) biarylaminium radical cations as tunable, precomplexing and potent photooxidants

Photocatalytic and electrochemical redox chemistries experiences fundamental limitations: i) the inabilities to harness the full photon redox energy and to engage molecules outside the solvent redox window as well as ii) thermodynamic redox potential-guided selectivity. Photoelectrochemistry circumvents such limitations for high-power redox processes. Triarylaminies are introduced as tunable, electroactivated photocatalysts for super-oxidations. Our discovery of dispersion precomplexation rationalizes i) the photochemistry of picosecond-lived photoexcited radical ion catalysts and ii) anti-Kasha photochemistry harnessing the full photon redox energy. Dispersion precomplexation presents a new selectivity handle, overturning redox potential-guided selectivity in photocatalysis.

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Hole-mediated photoredox catalysis: tris(*p*-substituted)biarylaminium radical cations as tunable, precomplexing and potent photooxidants†

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As a combination of visible light photoredox catalysis and synthetic organic electrochemistry, electrochemically-mediated photoredox catalysis emerged as a powerful synthetic technique in recent years, overcoming fundamental limitations of electrochemistry and photoredox catalysis in the single electron transfer activation of small organic molecules. Herein we report a tunable class of electroactivated photoredox catalyst, tri(*para*-substituted)biarylaminium cations, that become superoxidants in their photoexcited states even able to oxidize molecules beyond the solvent window limits of cyclic voltammetry (such as polyfluorobenzene and trifluorotoluene). Furthermore, we demonstrate that precomplexation not only permits the excited state photochemistry of tris(*para*-substituted)biarylaminium cations to overcome picosecond lifetime, but enables and rationalizes the surprising photochemistry of their *higher-order* doublet (D_n) excited states, unlocking extremely high oxidative potentials (up to a record-breaking $\sim +4.4$ V vs. SCE).

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Introduction

Synthetic Organic Electrochemistry (SOE)¹ and visible light PhotoRedox Catalysis (PRC),² which offer entries to single electron transfer (SET) chemistry and radical intermediates under mild conditions, have risen to the fore of contemporary organic synthesis. A key factor underpinning the success of PRC is the host of available photocatalyst structures with well-characterized photophysical and redox data, allowing chemists to ‘tune’ a given excited state to a desired process. Although PRC exhibits a selectivity benefit in transferring visible light photon energy to a colored transition metal-based or organic dye photocatalysts, its scope of applications are partially redox potential-limited by this photon energy (*ca.* 1.8–3.1 eV). Multiple-photon-accumulation strategies such as consecutive photoelectron transfer (conPET)³ and triplet-triplet annihila-

tion upconversion (TTA-UC)⁴ have proven elegant means to achieve powerful SET reductions, but their use in oxidations has eluded chemists. A limiting restriction in conPET is that both ground and radical ion states must be photoactive. Furthermore, net-oxidative/reductive PRC processes employ excess of a sacrificial oxidant/reductant which is necessary for photocatalyst turnover, but which may (or whose by-products may) (i) interfere with downstream chemistry and (ii) require separation.

In comparison to PRC processes, SOE can employ uncapped potentials to chemical redox reactions at the turn of a dial. However, electrode surfaces typically⁵ cannot discriminate between organic molecules aside from their innate order of thermodynamic redox potentials. Moreover, low electrical conductivity in organic solvents typically require applied potentials to be much higher than the target substrate’s redox potential.⁶ This encourages deleterious redox processes, such as those involving solvent especially if target SET processes lie near the electroactive window (typically *ca.* +3 to -3 V).^{6,7} In addition, mechanistic characterization of the heterogeneous electrolysis step (heterogeneous SET at the electrode surface)⁸ has remained a key challenge in SOE. Screening of electrode materials is inevitable, despite efforts to characterize materials by overpotential, resistivity, surface area, stability and cost.⁹

As a result of these limitations, synthetic photoelectrochemistry is emerging as a state-of-the-art in SET-mediated

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chemistry.^{10–12} Different categories for the merger of photo- and electrochemistry have been reported, including interfacial photoelectrochemistry (iPEC) involving photoelectrodes¹³ and decoupled photoelectrochemistry (dPEC) where photo- and electrochemical steps serve distinct mechanistic roles.¹⁴ A third category involves an intimate and synergistic relationship of photo- and electrochemical steps within the same catalytic cycle. Xu,^{15a,b} Lambert,^{15c} Lin^{15d} and Ackermann^{15e} reported elegant examples using electrochemistry to turnover spent photoredox catalysts thus obviating the need for sacrificial oxidants/reductants. This category can also electrogenerate photocatalysts to achieve super-oxidations or super-reductions beyond those accessible *via* photocatalysis alone.¹⁶ A variety of nomenclature has been coined for this sub-category: “electro-photocatalysis”^{16e} “photoelectrocatalysis”^{16f} and “electron-primed photoredox catalysis”.^{16g} We coined nomenclature “electrochemically-mediated PhotoRedox Catalysis (e-PRC)” as a blanket term to cover both net-oxidative/net-reductive variants¹⁰ and to avoid any misunderstanding with iPEC (Fig. 1A). In the net-oxidative direction, Moutet and Reverdy achieved the first e-PRC bimolecular cyclization of 1,1-diphenylethylene using phenothiazine (PTZ) as catalyst (Fig. 1C and E), but utilized UV light and only a few catalyst turnovers were reported.

Recently, Lambert reported an exotic trisaminocyclopro-phenium cation (TAC) as an e-PRCat for Nicewicz-type oxidative coupling. Chloro-, dichloro- and fluorobenzene, typically beyond reach of photoredox catalysis alone, could be engaged (Fig. 1C and E).^{16e} In context of these seminal reports, a class of e-PRC catalyst that offers tunability, to provide more powerful potentials or new handles for selectivity, is highly attractive.

Tri(*p*-substituted)arylamines, discovered by Walter and co-workers¹⁷ are renowned for their photophysical properties as hole-transport materials in OLEDs and in photovoltaics,¹⁸ as oxidative mediators in SOE^{19,20} and as oxidants in radical chain reactions.²¹ They exhibit fully reversible 1e[–] oxidations by cyclic voltammetry. Generally colorless in their neutral state, their oxidized tris(*para*-substituted)aminium radical cation forms are intensely colored. While the commercial tri (*4*-bromophenyl)aminium radical cation presents issues in synthesis,²² better behaved radical cations derive from tri (*para*-substituted)biaryl amines (TPAs) that are easily be prepared in a single/few steps from Pd-catalyzed cross-coupling reactions of tri-*p*-bromophenylamine with appropriate partners or after appropriate activation. Barham reported the use of tri (*para*-substituted)arylaminium radical cations bearing *p*-Me or *p*-Ph substituents that exhibited good solution stability and excellent stability as isolated solids, that mediated highly selective *N*-CH₃ oxidative functionalizations.²³ Given their favorable properties and facile synthetic accessibility, we foresaw TPAs as a suitable class of customizable e-PRC catalyst that would open new avenues to customize redox and photophysical parameters of e-PRCats. Moreover, the excellent stability of isolated TPA⁺s could be leveraged in photophysical and mechanistic characterization. Herein, we disclose TPAs as powerful and effective e-PRCats (up to E^{p}_{ox} (${}^*\text{Ar}_3\text{N}^+/\text{Ar}_3\text{N}$) ~ +4.3 V vs. SCE) that engage unactivated or electron-poor aromatic systems (E^{p}_{ox} = +2.1 V to >3.0 V) in SET oxidation (Fig. 1D and E). Of key importance is the first discovery of *precomplexation* in the ground state (Fig. 1B), which (i) circumvents ultrashort lifetimes of excited states, (ii) allows engagement of higher order excited states and (iii) enables contra-thermodynamic redox selectivity. Photoexcited triarylaminium radical cations are demonstrated as novel superoxidants in organic synthesis.

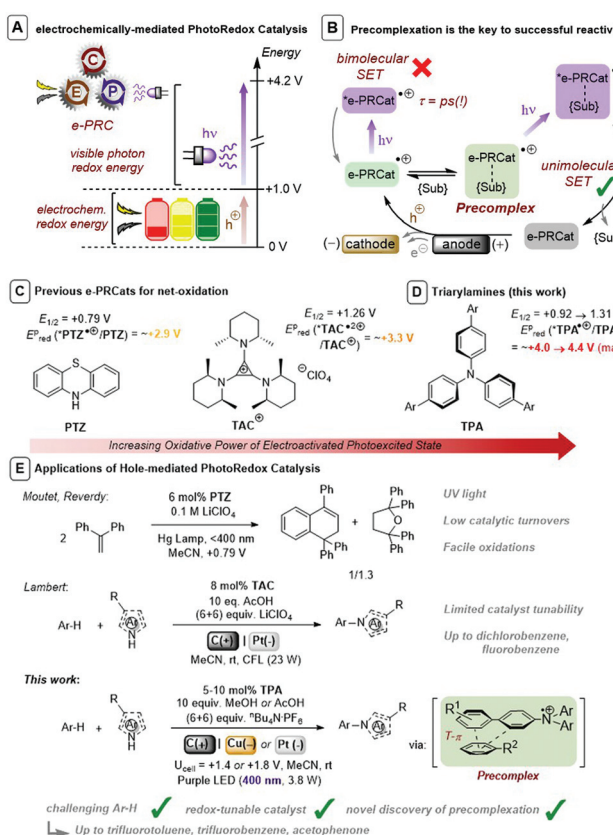


Fig. 1 (A) Concept of electrochemically-mediated photoredox catalysis. (B) Precomplexation rationalizes successful reactivity of picosecond lifetime photoexcited radical ions. (C) Previous e-PRCats for net oxidation. (D) Triarylamines as e-PRCats. (E) Applications of hole-mediated photoredox catalysis and contextualization of this work.

Results and discussion

Synthetic results

We began by screening different TPA e-PRCats (Generation 1, Fig. 2) with different half-wave oxidation potentials in the Nicewicz model reaction;²⁴ the oxidative C–H amination of mesitylene ($E^{\text{p}}_{\text{ox}} = +2.1$ V vs. SCE)²⁴ with pyrazole (Table 1). After setting an anodic constant potential of $U_{\text{cell}} = +1.4$ V (Fig. 3), color developed in the anodic compartments for each TPA e-PRCat, consistent with the expected formation of their TPA⁺s. Gratifyingly, irradiation under the specified conditions afforded C–H activation product **3aa**. Control reactions in absence of light, potential or e-PRCat confirmed the operation



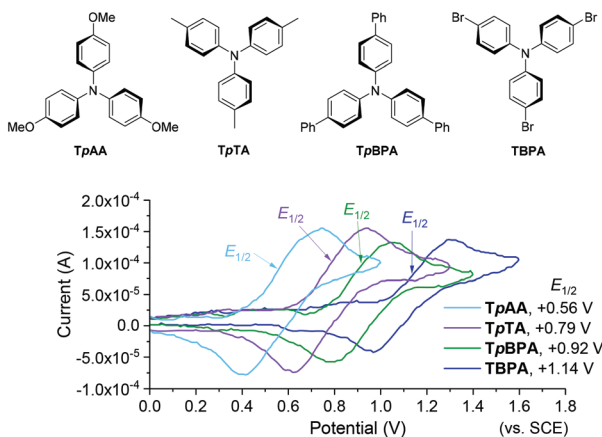
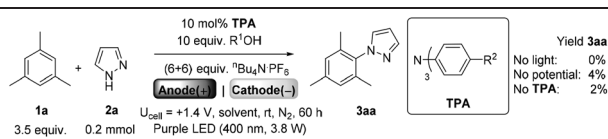


Fig. 2 First generation TPA e-PRCats (top). Cyclic voltammograms (bottom).

Table 1 Optimization of e-PRC C–H heteroamination using first generation TPAs



Entry	R ¹	R ²	Anode/cathode	Yield 3aa/r.s.m. ^a	Solvent
1	Ac	OMe	Cfoam/Pt	21—	MeCN
2	Ac	Me	Cfoam/Pt	23/14	MeCN
3	Ac	Br	Cfoam/Pt	35—	MeCN
4	Ac	Ph	Cfoam/Pt	69/15	MeCN
5 ^b	Ac	Ph	Cfoam/Pt	18/8	MeCN
6	Me	Ph	Cfoam/Pt	72/—	MeCN
7	Me	Ph	Cfelt/Pt	53/—	MeCN
8	Me	Ph	Cfoam/Fe	70/—	MeCN
9	Me	Ph	Cfoam/Cu	88 ^c /—	MeCN
10	Me	Ph ^d	Cfoam/Cu	75/—	MeCN
11	Me	Ph	Cfoam/Cu	60/—	MeCN ^e
12	Me	Ph	Cfoam/Cu	—/—	DMF
13	Me	Ph	Cfoam/Cu	82/—	DCM

r.s.m., returned starting material. ^aYields determined by ¹H NMR using dibromomethane as an internal standard. ^b 0.35 W 400 nm LED.

^c Average of two replicates. ^d 5 mol% TPA. ^e LiClO₄ as electrolyte.

of e-PRC. Interestingly, tri([1,1'-biphenyl]-4-yl)amine (**TpBPA**) afforded a notably higher yield of **3aa** than the commercial tris (4-bromophenyl)amine (entry 4) despite having an appreciably lower $E_{1/2}$. After further optimization, using MeOH as proton source, carbon foam as WE and copper as CE gave the best result (entry 8). The yield of **3aa** tracked well with increasing applied constant potential (Fig. 3). For full optimization studies, see the ESI.† With optimal conditions in hand, the amination of arenes with a variety of pharmaceutically-relevant N-heterocycles was explored (Table 2). Halide-bearing and carbonyl (aldehyde, ketone and ester)-bearing pyrazoles, triazole, benzotriazole, and a functionalized derivative of benzimidazole afforded generally good to excellent (50–89%) yields of aminated mesitylenes **3aa–3aj**. We note that benzimidazole

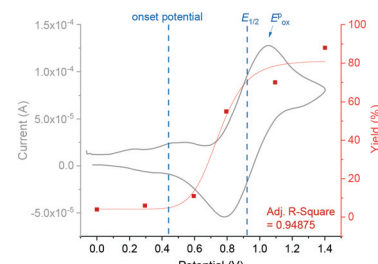
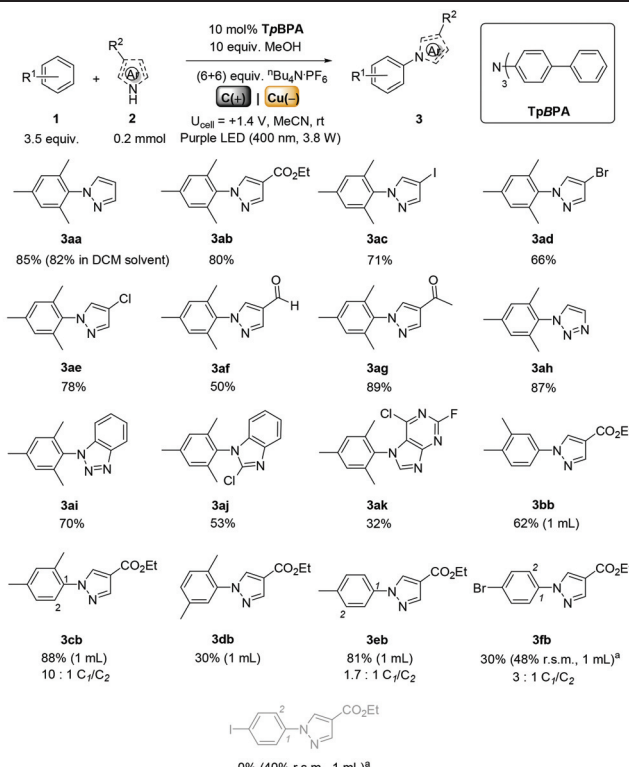


Fig. 3 Yield as a function of increasing applied U_{cell} (top). Development of color with increasing applied U_{cell} (bottom).

Table 2 e-PRC C–H heteroaminations using TpBPA



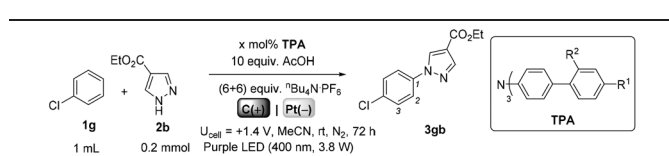
Unless otherwise stated, all reactions used 3.5 eq. arene; isolated yields. Yields in parenthesis determined by ¹H NMR. ^a Pt CE and AcOH were used.

derivatives have not been reported as nucleophiles in previous photoelectrochemical arene amination or conPET photo-catalytic methods.^{13d,16e,25} 6-Chloro-2-fluoropurine afforded a modest yield of **3ak** (32%). Xylenes and toluene were tolerated to afford **3bb–3fb** in moderate to excellent (30–88%) yields. Interestingly, the reactivity trend of xylenes followed the order



meta->*ortho*->*para*-xylene, despite E^P_{ox} following the opposite trend (see Mechanistic Study).²⁶ Toluene has a higher E^P_{ox} than xylenes but reacted to give 81% of **3eb**.²⁶ Bromobenzene afforded a 30% yield of **3fb** with notable r.s.m., while iodobenzene gave no reaction (60% r.s.m.). Benzene and PhCl were unsuccessful, presumably due to their notably higher E^P_{ox} (only a 10% yield of **3gb** was obtained, even with 72 h and an excess of PhCl). CE Substitution of Cu for Pt wire cathode and substitution of MeOH for AcOH increased the yield to 35% (Table 3). We could not improve the yield beyond this threshold. Leveraging the facile customization of TPAs, we synthesized **TpBPA** derivatives (Fig. 4) with electron-withdrawing groups to bolster their TPA^+ excited state potentials (entries 1–4). Of these, tris(4'-cyano-[1,1'-biphenyl]-4-yl)amine (**TCBPA**) increased the yield of **3gb** to 46% (Table 3). In contrast to **TpBPA**, the optimal **TCBPA** catalyst loading was 5 mol% (entries 4–8), increasing the yield of **3gb** to 69% (entry 6). Notably, the reaction was still efficient with only 1.5 mol% of **TCBPA** (entry 8). Under optimal conditions, reactions of PhCl, benzene and even fluorobenzene were enabled, affording **3gb**–**3ib** in modest to good (30–65%) yields (Table 4). Our conditions tolerated free carboxylic acids (**3hm**), prone to de-carboxylation under typical PRC. Interestingly, under the same applied constant potential U_{cell} as in Table 2 and in contrast to the use of **TpBPA**, here toluene underwent benzylic oxidation instead of amination, while a pyrazole-4-carboxaldehyde underwent oxidation to give ultimately **3hm**. Bromobenzene gave a lower yield of **3fb** than in Table 2. These observations indicate a less oxidizing TPA^+ excited state (${}^*\text{TpBPA}^+$) is beneficial for certain substrates and demonstrates the value of tunability presented by this class of e-PRCats. We further probed the limits of arene SET oxidations with **TCBPA**⁺, by targeting 1,2-dichlorobenzene and trifluorotoluene, and were encouraged to detect products, albeit in low yields, when

Table 3 e-PRC C–H heteroamination optimization with second generation TPA e-PRCats



Entry	R ¹	R ²	Yield 3gb/r.s.m. ^a	Catalyst loading 'x
1	H	H	35/33	10
2	tBu	H	38/22	10
3	F	F	36/18	10
4	CN	H	46 ^b /25	10
5	CN	H	40/18	20
6	CN	H	69/—	5
7	CN	H	52/—	3
8 ^c	CN	H	45/25	1.5

r.s.m., returned starting material. ^aYields determined by ¹H NMR using dibromomethane as an internal standard. ^bAverage of two replicates. ^cReaction conducted with **1g** (1 mL), **2b** (0.4 mmol), **TCPBA** (0.006 mmol), AcOH (4 mmol), ⁿBu₄N-**PF₆** ((1.2 + 1.2) mmol), MeCN (3.5 + 3.5 mL).

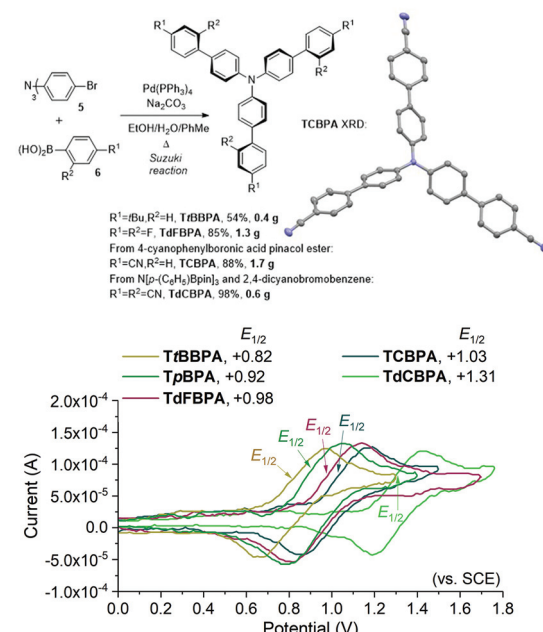
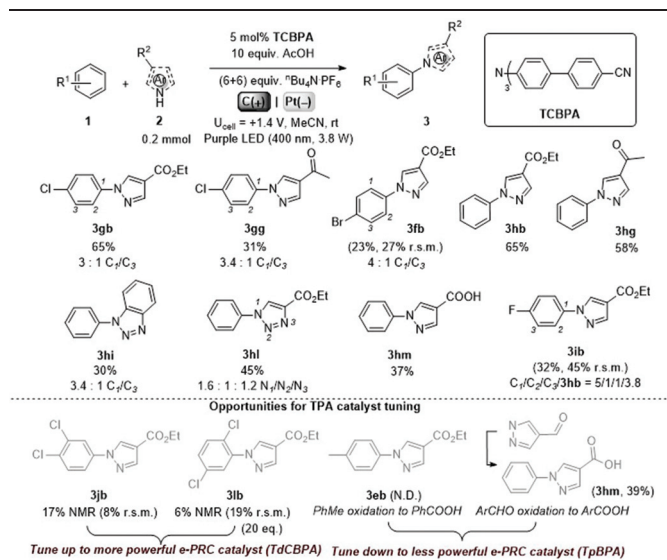


Fig. 4 Second Generation of TPA e-PRCats, and the XRD structure of TCPBA (top). Thermal ellipsoids are set at the 50% probability level. H atoms omitted for clarity. C atoms (grey), N atoms (blue). Cyclic voltammetry (bottom).

Table 4 e-PRC C–H heteroamination using TCBPA

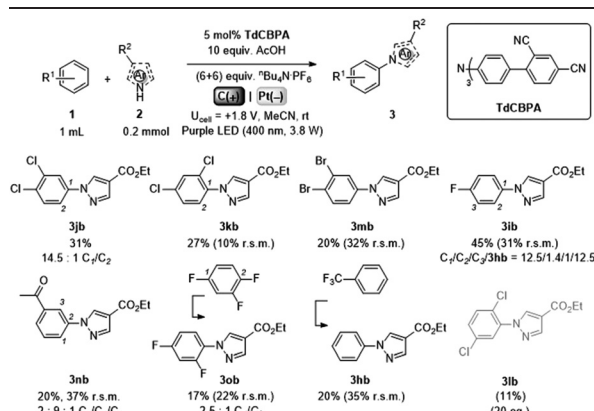


N.D. not determined. Unless otherwise stated, reactions used 1 mL arene; isolated yields. Yields in parenthesis determined by ^1H NMR.

using **TCBPA** (17% of **3jb** and 7%, respectively). Yields did not increase with extended time (96 h) or higher potential ($U_{\text{cell}} = +1.8$ V).

Gratifyingly, further e-PRCat tuning in the form of the even more electron-deficient tris(2',4'-dicyano-[1,1'-biphenyl]-4-yl) amine (**TdCBPA**) increased the yield of **3jb** to a satisfactory 31% (Table 5). Acetophenone could also be oxidized giving

Table 5 e-PRC activation of highly electron-deficient arenes using **TdCBPA**



Isolated yields. Yields in parenthesis determined by ^1H NMR.

3nb in 20% yield. Although polyfluorinated arenes gave **3ob** in modest yields (~20%), oxidative $\text{S}_{\text{N}}\text{Ar}$ -type activation of such a challenging substrate has not been previously accomplished. In competition with C–H activation, due to the role of F atom as good leaving group in $\text{S}_{\text{N}}\text{Ar}$, net redox-neutral C–F substitution occurred to give **3ob**.^{15c,27} Interestingly, C_6H_5 -bearing compound **3hb** was isolated from the reaction of trifluorotoluene, indicating $\text{S}_{\text{N}}\text{Ar}$ of a CF_3 group. While the role of CF_3 groups in promoting $\text{S}_{\text{N}}\text{Ar}$ reactions is well-known, no prior examples of a formal $\text{C}(\text{sp}^2)\text{–CF}_3$ to $\text{C}(\text{sp}^2)\text{–N}(\text{Het–Ar})$ substitution exist. The potentials (E_{ox}^{P}) of most arenes in this study have been measured by previous reports^{15c,16e} while those of acetophenone, trifluorotoluene and 1,2,4-trifluorobenzene exceed the solvent windows of MeCN and DMF ($>+3.0$ V vs. SCE). This reflects the remarkable oxidizing power of **TdCBPA**⁺'s excited state. In contrast to a previous report^{16e} and in line with reactivity of xylenes herein, product yields increased as a function of the substitution pattern on dichloroarenes (1,4- < 1,3- \leq 1,2-dichlorobenzene). This was surprising, given the corresponding increasing E_{ox}^{P} (1,4- < 1,2- \leq 1,3-dichlorobenzene).^{16e}

Mechanistic studies

Spectroelectrochemical analysis of **TpBPA** (Fig. 5) **TCBPA** and **TdCBPA** (see ESI†) revealed the formation of their respective TPA^+ 's by the disappearance of the TPA band ($\lambda_{\text{max}} = \text{ca}$.

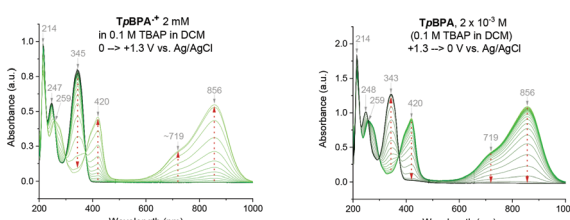


Fig. 5 Spectroelectrochemistry of **TpBPA**.

345 nm) and appearance of two broad absorption bands between 360–430 nm ($\lambda_{\text{max}} = \text{ca. } 420$ nm) and between 600–900 nm ($\lambda_{\text{max}} = \text{ca. } 719, 856$ nm), when the potential was increased from 0 to +1.3 V. Excellent reversibility was observed upon returning the potential to +0.0 V, indicating high stability of the TPA^+ 's as e-PRCats.

Neutral **TpBPA** possesses a strong absorption at $\lambda_{\text{max}} = 365$ nm, where **TpBPA**⁺ absorbs poorly. Performing our optimized synthetic procedure for **3aa** with 365 nm LEDs afforded only 11%, suggesting photoexcitation of neutral **TpBPA** is not a dominant mechanistic factor. The aforementioned control reaction without applied potential afforded only 4% of **3aa**. In contrast to **TpBPA**, **TCBPA** does absorb appreciably at 395 nm. Nevertheless, in the absence of an applied potential for the optimized synthesis of **3gb**, only a 12% yield of **3gb** was observed. The very substantial yield differences when applied potential is present or absent confirm the pivotal role of the TPA^+ 's and e-PRC as the main product-forming pathway. Inspecting the UV-visible absorption spectra of **TpBPA**⁺ and **TCBPA**⁺, we reasoned the longest wavelength bands (λ_{max} at *ca.* 719 and 856 nm, respectively) must contain their $\text{D}_0 \rightarrow \text{D}_1$ transitions. Based on the photochemical interpretation of Kasha's rule that prohibits photochemistry from higher order excited states, we irradiated the reactions forming **3aa** and **3gb** with 740 nm and 850 nm and were mystified to observe *no reaction* in either case (Fig. 6). That successful reaction was only observed at 395 nm implicated anti-Kasha behavior; a *higher-order excited state* participating in SET photooxidation. Such behavior is as surprising as it is intriguing, since (i) not only are examples of photochemistry violating Kasha's rule rarely reported in organic synthesis^{3b,c,28,29} but (ii) the reported lifetimes of photoexcited radical ion species are already ultra-short.³⁰ In fact, despite a number of conPET/e-PRC reports invoking their photochemistry in super-oxidations/reductions, the question of how photoexcited radical ion species could ever participate in photochemical processes has eluded chemists. The ultrashort lifetimes of these species lie beneath timescales of diffusion-controlled bimolecular quenching.¹⁰

Greatly aiding our mechanistic study was the fact that TPA^+ 's can be isolated as bench stable PF_6^- salts.²³ Their crystal structures revealed a common propeller-type structure consistent with parent TPAs (Fig. 7). However, attempts to investigate quenching of photoexcited TPA^+ 's were thwarted by the fact that they *do not exhibit steady-state fluorescence* (see ESI†). Consistent with this observation, reported lifetimes of related

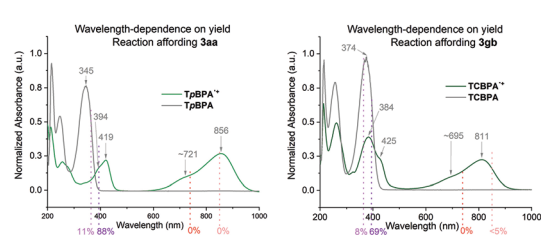


Fig. 6 Wavelength dependence on product yields.



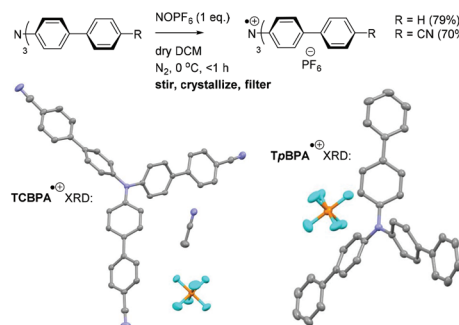


Fig. 7 Synthesis of **TpBPA**⁺, **TCBPA**⁺ and their XRD crystal structures. Thermal ellipsoids are set at the 50% probability level. H atoms omitted for clarity; C atoms (grey), F atoms (cyan), N atoms (blue), P atoms (orange).

excited N radical cation species³¹ lie within the femto- to picosecond timeframe. Transient absorption spectroscopy (TAS) was therefore employed to determine the lifetimes of excited **TpBPA**⁺ and **TCBPA**⁺. Pumping with a broadband visible light source (490–900 nm)³² revealed a ground state bleach in the 600–850 nm bands (Fig. 8) and an excited state absorption between 490–570 nm, indicative of absorption by the D_1 (or D_2) excited state. The lifetimes of the D_1 excited states of **TpBPA**⁺ and **TCBPA**⁺ were 4.6 ps and 8.6 ps, respectively, clearly ruling out diffusion-controlled bimolecular quenching. It is reasonable to assume higher order excited states possessing even shorter lifetimes.

Precomplexation could rationalize productive, unimolecular SET and the aforementioned anti-Kasha behaviour. Presuming that precomplexes may possess different UV-vis absorptions than their parent TPA^+ 's, we investigated quenching of the absorption of TPA^+ 's in the presence of representative concentrations of arenes (Fig. 9). In the presence of mesitylene however, the spectrum of **TpBPA**⁺ was unchanged. Gratifyingly, irradiation with 395 nm light effected gradual conversion of **TpBPA**⁺ to **TpBPA**, corroborating the expected

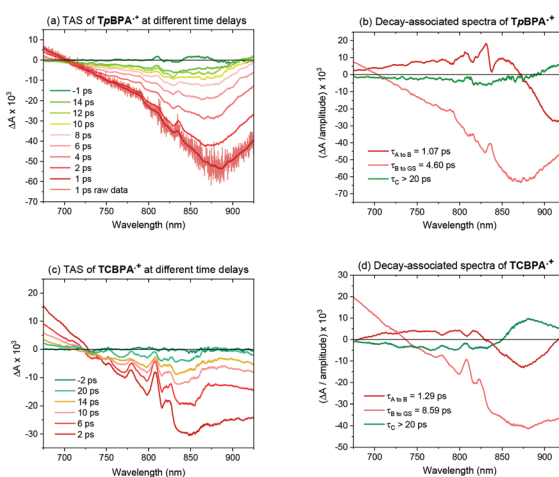


Fig. 8 Transient absorption and decay-associated spectra of TPA^+ 's.

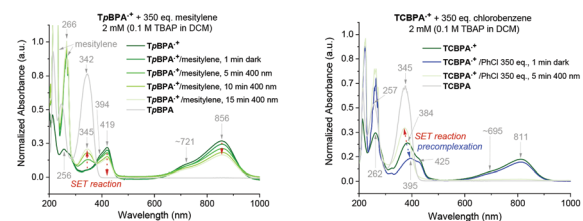


Fig. 9 UV-vis spectra of **TpBPA**⁺ (top) and **TCBPA**⁺ (bottom) in the presence of mesitylene and PhCl, respectively and after irradiation at 395 nm.

SET from mesitylene to the photoexcited TPA^+ . Interestingly and in contrast, the spectrum of **TCBPA**⁺ was altered by PhCl; a small bathochromic perturbation of the peak at 384 nm to 395 nm occurred. Irradiation with 395 nm completely converted **TCBPA**⁺ to **TCBPA** after minutes.

Given the paramagnetic nature of TPA^+ 's, we reasoned that a change in their EPR spectra in the presence of representative arene concentrations would be more conclusive in corroborating precomplexation. The EPR signal of **TpBPA**⁺ showed a triplet ($a_N = 8.9$ G). Addition of mesitylene to **TpBPA**⁺ caused its EPR signal to shift ($\Delta B = 4.5$ G) to lower G values (Fig. 10), but the signal shape was largely unchanged. This indicates that the spin density of the **TpBPA**⁺ is largely unaffected when it undergoes precomplexation with mesitylene. A less pronounced shift in G value occurred in the presence of iodobenzene ($\Delta B = 2.0$ G) but flattening of the triplet shoulders was observed (see ESI†). Based on these spectroscopic differences, we hypothesize that a different type of precomplex occurs in this case that is ‘unreactive’, to rationalize the inability to engage this substrate under the reaction conditions despite its more accessible redox potential ($E_{\text{ox}}^{\text{P}} = +2.07$ vs. SCE) than toluene ($E_{\text{ox}}^{\text{P}} = +2.28$ V vs. SCE).³³ In contrast to **TpBPA**⁺, simulated fitting of the EPR signal from **TCBPA**⁺ revealed two radical species, one triplet and one superimposed singlet appearing as a large broad central feature. We presume **TCBPA**⁺ exists as a mixture of rotamers in solution, one propeller-type as observed in the solid state, and one in which a biphenyl unit falls into conjugation with the N radical cation, consistent with reported behaviour for similar compounds.^{34,35} Addition of PhCl caused a notable change in the signal shape, giving exclusively the triplet signal (Fig. 11), consistent with the change in UV-vis (Fig. 9) and indicating that the spin density

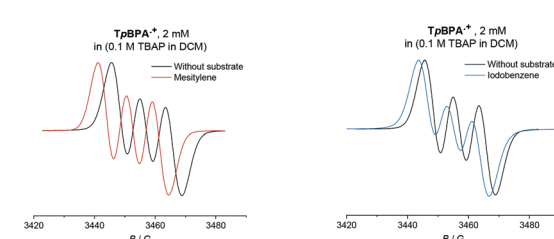


Fig. 10 EPR spectra of **TpBPA**⁺ with reactive mesitylene (left, 350 eq.) and unreactive iodobenzene (right, 350 eq.).



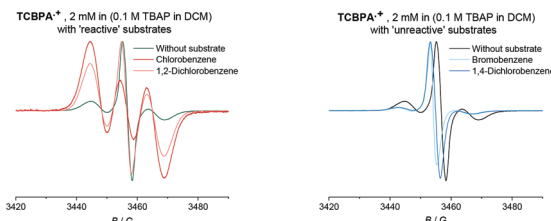


Fig. 11 EPR spectra of the TCBPA^+ with reactive (left) and unreactive (right) substrates (350 eq.).

of TCBPA^+ is notably affected by precomplexation with PhCl . A similar change was detected upon addition of 1,2-dichlorobenzene, but not for its 1,4-congener (17% yield of **3jb** vs. 6% yield of **3lb** under TCPBA e-PRC) and not for PhBr (22% of **3fb** from TCPBA e-PRC) which instead all gave spectra clearly favoring the ‘unreactive’ complex (singlet signal).³⁶

Density Functional Theory (DFT) calculations modelled precomplexation of various TPA^+ /arene combinations (Table 6). For unsymmetrical (halo)arene substrates, orientations of the complex with halogen facing both ‘in’ to the N radical cation and ‘out’ were explored (see ESI† for full investigations). We assumed that π -stacking interactions³⁷ at the TPA^+ ’s biphenyl unit could be responsible for precomplexation. Attempts to position PhCl or mesitylene substrates in a sandwich or parallel-displaced π - π stacking interaction (“ π - π ” complex) around the inner N-bearing ring of their respective TPA^+ ’s led to dissociation, whereas positioning of the substrates around the terminal aromatic ring identified local minima resembling T-type stacking interaction (“T- π ” complexes). For this

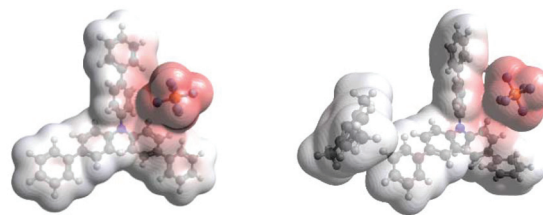


Fig. 12 DFT spin densities of TpBPA^+ without (left) vs. with (right) Mesitylene.

complex, minimal change in the spin density was detected for $\text{TpBPA}^+ + \text{mesitylene}$ (Fig. 12), whereas a large shift in spin density occurred for $\text{TCBPA}^+ + \text{PhCl}$ where the Cl atom was facing inwards (Fig. 13). This is consistent with the changes in EPR and UV-vis spectra, and so we assigned this T- π complex as the one responsible for the triplet EPR signal and successful reactivity, since. On the other hand, for less successful substrate PhBr and unsuccessful PhI (no product, 60% recovered **2b**), a π - π complex was presumed to be responsible for the broad singlet EPR signal. Delocalization of the N radical cation over the biphenyl aromatic system would lead to stabilization, presumably decreasing E^p_ox of the photoexcited TPA^+ . Free energies of precomplexation were endergonic at the levels of theory employed,³⁸ this was confirmed to be a result of the implicit solvation model (see ESI†). It is most important to consider relative trends. Intermolecular distances for T- π stacking and π - π stacking were close to previously-reported distances for simpler complexes/dimers.^{39–41}

Table 6 Free energies and intermolecular distances for T- π or π - π precomplexes

Complex ^a	Complexation ΔG (kcal mol ⁻¹)		Intermolecular distance (Å)	
	T- π	π - π	T- π	π - π
$\text{TpBPA}^+ + 1,3,5\text{-TMB}$	+7.2 ^b	+4.9 ^{c,f}	3.3–5.5 ^b	3.6–4.2 ^{c,f}
$\text{TpBPA}^+ + \text{PhI}$	+28.4 ^{b,d}	+28.3 ^{b,d}	4.7–6.5 ^{b,d}	5.2–5.5 ^{b,d}
	+28.1 ^{b,e}	+26.1 ^{b,e}	3.8–6.2 ^{b,e}	6.2–6.8 ^{b,e}
$\text{TCBPA}^+ + \text{PhCl}$	+4.5 ^{b,d}	+2.9 ^{c,e,f}	4.5–6.8 ^{b,d}	3.7–4.3 ^{c,e,f}
	+3.5 ^{c,d}		3.2–5.3 ^{c,d}	
	+5.1 ^{b,e}		4.8–7.4 ^{b,e}	
$\text{TCBPA}^+ + \text{PhBr}$	+30.8 ^{b,d}	29.7 ^{b,d}	3.4–6.2 ^{b,d}	4.9–5.4 ^{b,d}
	+31.4 ^{b,e}	N.D. ^{b,e}	3.7–6.9 ^{b,e}	N.D. ^{b,e}
$\text{TCBPA}^+ + 1,2\text{PhClCl}$	+4.5 ^{b,d}	+6.6 ^{b,d}	4.2–6.4 ^{b,d}	5.3–5.7 ^{b,d}
	+4.7 ^{b,e}	N.D. ^{b,e}	4.6–6.9 ^{b,e}	N.D. ^{b,e}
$\text{TpBPA}^+ + 1,3\text{-PhClCl}$	+4.9 ^{b,d}	N.D. ^{b,d,g}	N.D. ^{b,d}	N.D. ^{b,d,g}
	+5.2 ^{b,e}	N.D. ^{b,e,g}	4.9–7.4 ^{b,e}	N.D. ^{b,e,g}
$\text{TCBPA}^+ + 1,4\text{-PhClCl}$	N.D. ^{b,g}	+4.2 ^{b,d}	N.D. ^{b,g}	5.5–5.8 ^{b,d}

N.D. not determined. In all cases, MeCN solvent was modelled implicitly. Pseudopotentials were applied to I and Br atoms (see ESI†). Intermolecular distances quoted are not centroid-to-centroid of the arene rings and are defined in the ESI.† ^a Hypothesized orientation of the precomplex matching UV-vis/EPR data and reactivity patterns. ^b uB3LYP/6-31+g(d,p). ^c oB97XD/6-31+g(d,p). ^d Halogen atom(s) facing “out”. ^e Halogen atom(s) facing “in”. ^f Rearranged from the T- π complex. ^g Dissociated.

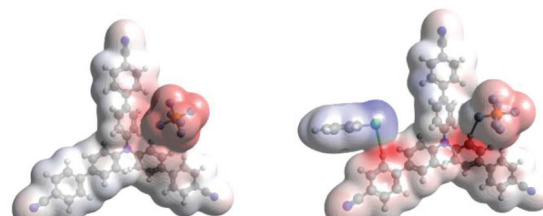


Fig. 13 DFT spin densities of TCBPA^+ without (left) vs. with (right) PhCl , T- π complex, Cl atom facing “in”.

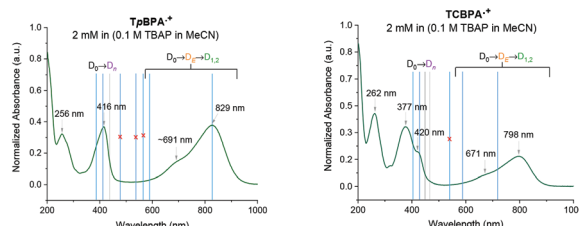


Fig. 14 Computed (CAM-B3LYP/6-31G(d,p), CPCM = MeCN, blue lines) vs. experimental UV-vis spectra of TpBPA^+ and TCBPA^+ in MeCN containing 0.1 M $\text{Bu}_4\text{N}^+\text{PF}_6^-$. Negligible TD-DFT excitations (coefficient $f < 0.00020$) in grey. Assignments based on the optical transitions.



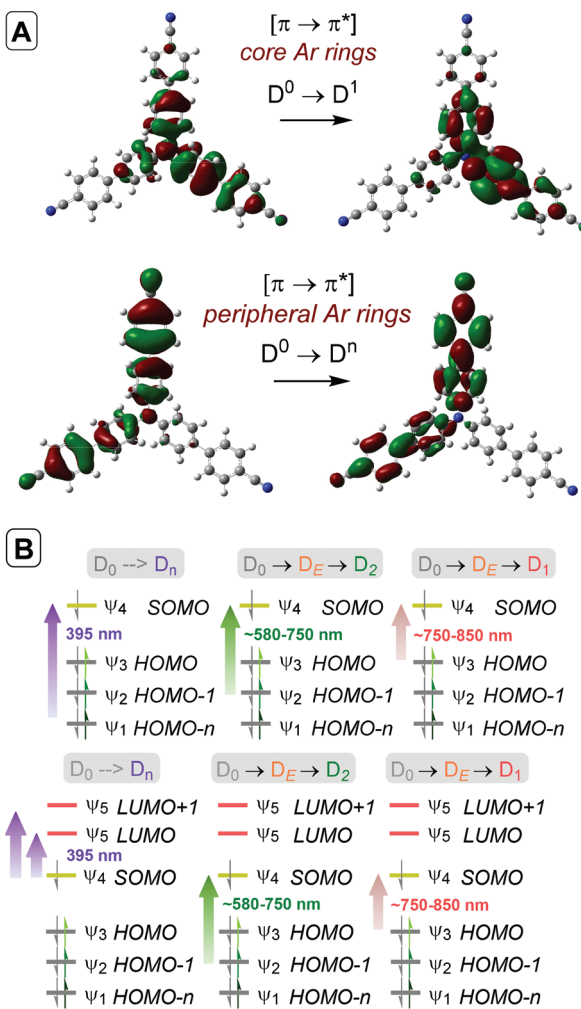


Fig. 15 (A) Natural Transition Orbitals (NTOs) depicting photoexcitation of TCBPA⁺ to D₁ or D_n. (B) TD-DFT predicted orbital transitions of TpBPA⁺ (top) and TCBPA⁺ (bottom). Wavelengths are derived from optical transitions.

Mesitylene, chlorobenzene, 1,2-dichlorobenzene and their respective TPA⁺'s all had accessible ΔG values (+3.5–7.2 kcal mol⁻¹) for the 'reactive' T-π complex. An accessible ΔG was also found for 1,3-dichlorobenzene (+4.9–5.2 kcal mol⁻¹) consistent with its reactivity (20% 3 kb under the TCBPA e-PRC conditions of Table 4). Iodo- and bromobenzene as substrates gave very high ΔG values for T-π complexes and their π-π complexes were more accessible albeit still highly endergonic. Attempts to obtain a T-π complex for TCBPA⁺ with 1,4-dichlorobenzene led to dissociation, while its π-π complex was accessible.

Time-dependent density functional theory (TD-DFT) calculations investigated the energies of higher order excited D_n states for TpBPA⁺, TCBPA⁺ and TdCBPA⁺, using both CAM-B3LYP/6-31G(d,p) and ωB97XD/6-31+G(d,p).^{31a,42} The calculated UV-visible spectra of TpBPA⁺ and TCBPA⁺ in MeCN reasonably agreed with experimental spectra (Fig. 14). The broad visible band (ca. 580–850 nm) is known to result from symmetry breaking^{35a} of the first excited state D_E in solution and in the excited state to give the D₁ and D₂ states, meaning TD-DFT transitions of the 580–850 nm region were less accurate than those of higher order excitations. Canonical molecular orbital (MO) calculations^{31a,43} revealed that excited states involved HOMO-'n' to SOMO transitions, typical of hole-particle excitations (Fig. 15A).^{16e,31a} Due to the complexity in interpretation of Canonical MO transitions of higher excited state transitions, Natural Transition Orbitals (NTOs) were employed to visualize the changes in 'hole density'.⁴⁴ For all TPA⁺'s studied, the first (and second) excited states (D₀→D_E→D_{1,2}) involved π → π* transitions around the core aromatics, while the higher order excited states (D₀→D_n) corresponding to 395 nm involved π → π* transitions around the peripheral aromatics (Fig. 15B). The concentration of 'hole density' at peripheral aromatics is exactly where it would be in closest proximity to substrate arenes in reactive T-π precomplexes. Ruling out participation of the first two excited states and states accessed at wavelengths <380 nm, we predict 'effective maximum' excited state potentials of TpBPA⁺ at

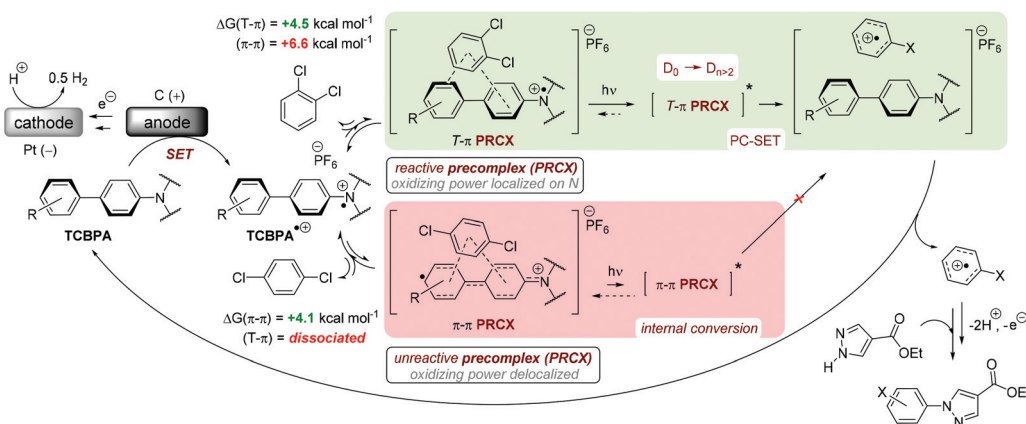


Fig. 16 Proposed e-PRC mechanism involving precomplexation modes for π-stacking and photochemistry from a higher order (than D₁/D₂) photo-excited state.

+4.02 V, **TCBPA**⁺ at +4.19 V and **TdCBPA**⁺ at +4.41 V *vs.* SCE from the Rehm-Weller equation.⁴⁵ A mechanism is proposed consistent with spectroscopic and computational studies herein (Fig. 16). Anodic SET oxidation generates the **TPA**⁺ from its **TPA**. Photoexcitation of the **TPA**⁺ to its D₁/D₂ or higher D_n states followed by bimolecular SET reductive quenching is prohibited by the **TPA**⁺'s picosecond lifetime. Instead, preassociation occurs to give a reactive T- π or an unreactive π - π precomplex (**PRCX**), depending on the sterics of the arene substrate.^{46,47} In the latter case, conjugative stabilization of the N radical cation decreases *E*⁰_{ox} of the ***TPA**⁺ below the threshold for productive unimolecular SET such that photoexcitation of the **PRCX** leads simply to non-radiative photochemical relaxation processes (such as internal conversion). In the former case, photoexcitation of the **PRCX** yields unimolecular SET reductive quenching of the ***TPA**⁺, regenerating the **TPA** and generating the arene radical cation to be intercepted by the N-heterocyclic nucleophile **2b** followed by loss of protons and further SET (anodic or by the **TPA**⁺) to yield product **3**. The prerequisite for precomplexation rationalizes the typical requirement for an excess of arene (3.5 eq. up to 1 mL, ~40 eq. herein) to drive precomplexation equilibrium in arene amination reactions mediated by radical cations.^{15c,16e}

Conclusions

We report tri(*p*-substituted)biarylaminium (TPAs) as a novel class of tunable, electroactivated photoredox catalyst. Photoexcited tri(*p*-substituted)biarylaminium radical cations (TPAs⁺) are demonstrated as highly potent SET oxidants. Straightforward customization of TPAs allows tuning of UV-vis absorptions, redox potentials and handles for precomplexation within the TPAs⁺. We report the first evidence of dispersion controlled (π -stacking) precomplexation in synthetic photoelectrochemistry and in the photochemistry of excited radical ions, which serves as a unique control element. Precomplexation enables remarkable photochemical phenomena: (i) circumvention of the ultrashort lifetimes of excited radical ion states for their use in photocatalysis, (ii) anti-Kasha fashion engagement of *higher-order* excited states in photocatalysis, (iii) overturning of thermodynamic selectivity dictated by redox potentials by virtue of steric/electronic factors involved in the precomplex. We are excited to witness future opportunities in reactivity and selectivity that dispersion precomplexation may provide to photocatalysis, in addition to recent developments in the control of photochemical outcomes through confined spaces and ordering of solvent.^{48–50}

Author contributions

S. W. performed the optimization study, conducted all photoelectrochemical reactions, synthesized **TPA**⁺s and measured spectroelectrochemistry and UV-vis spectra of compounds. S. W., J. P. B. and J. Ž. designed TPAs while J. Ž. and M. D. synthesized second generation TPAs. Under the gui-

dance of J. R., P. H. measured and analyzed all EPR spectra of **TPA**⁺s and precomplexes. V. B. and J. P. B. shared the DFT calculations of precomplexes, spin densities and TD-DFT calculations. D. J. S. performed XRD studies on **TPA**⁺s. Under the guidance of E. T. and J. H., A. K. measured TAS of **TPA**⁺s. E. T. measured fluorescence, EEM spectra and TCSPC of **TPA**⁺s. J. H. and E. T. analyzed and interpreted TAS and luminescence data. J. B. and S. W. together wrote the manuscript and analyzed all other spectroscopic data. J. B. conceived and guided the study, designed photoelectrochemical cells, conducted all CV measurements, guided the overall project and facilitated collaborations. All authors checked the manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- Selected reviews: (a) K. D. Moeller, *Tetrahedron*, 2000, **56**, 9527–9554; (b) J.-I. Yoshida, K. Kataoka, R. Horcajada and A. Nagaki, *Chem. Rev.*, 2008, **7**, 2265–2299; (c) B. A. Frontana-Uribe, R. D. Little, J. G. Ibanez, A. Palma and R. Vasquez-Medrano, *Green Chem.*, 2010, **12**, 2099–2119; (d) M. Yan, Y. Kawamata and P. S. Baran, *Chem. Rev.*, 2017, **117**, 13230–13319; (e) A. Weibe, T. Geishoff, S. Mohle, E. Rodrigo, M. Zirbes and S. Waldvogel, *Angew. Chem., Int. Ed.*, 2018, **57**, 5594–5619.
- Selected reviews: (a) K. Zeitler, *Angew. Chem., Int. Ed.*, 2009, **48**, 9785–9789; (b) J. M. R. Narayanan and C. R. J. Stephenson, *Chem. Soc. Rev.*, 2011, **40**, 102–113; (c) C. Prier, D. Rankic and D. W. C. MacMillan, *Chem. Rev.*, 2013, **113**, 5322–5363; (d) S. Fukuzumi and K. Ohkubo,



Chem. Sci., 2013, **4**, 561–574; (e) N. A. Romero and D. A. Nicewicz, *Chem. Rev.*, 2016, **116**, 10075–10166; (f) K. L. Skubi, T. R. Blum and T. P. Yoon, *Chem. Rev.*, 2016, **116**, 10035–10074.

3 (a) I. Ghosh, T. Ghosh, J. I. Bardagi and B. König, *Science*, 2014, **346**, 725–728; (b) I. A. MacKenzie, L. Wang, N. P. R. Onuska, O. F. Williams, K. Begam, B. D. Duneitz, A. M. Moran and D. A. Nicewicz, *Nature*, 2020, **580**, 76–80; (c) J. P. Cole, D.-F. Chen, M. Kudisch, R. M. Pearson, C.-H. Lim and G. M. Miyake, *J. Am. Chem. Soc.*, 2020, **142**, 13573–13581.

4 (a) M. Majek, U. Faltermeier, B. Dick, R. Pérez-Ruiz and A. J. von Wangelin, *Chem. – Eur. J.*, 2015, **21**, 15496–15501; (b) C. G. López-Calixto, M. Liras, V. A. de la Peña O’Shea and R. Pérez-Ruiz, *Appl. Catal., B*, 2018, **237**, 18–23; (c) C. Kerzig and O. S. Wenger, *Chem. Sci.*, 2019, **10**, 11023–11029; (d) B. D. Ravetz, A. B. Pun, E. M. Churchill, D. N. Congreve, T. Rovis and L. M. Campos, *Nature*, 2019, **565**, 343–346.

5 Electrode materials can be modified with chiral additives to perform enantioselective reductions or oxidations, see: (a) M. Ghosh, V. S. Shinde and M. Rueping, *Beilstein J. Org. Chem.*, 2019, **15**, 2710–2746; (b) Q. Lin, L. Li and S. Luo, *Chem. – Eur. J.*, 2019, **25**, 10033–10044.

6 C. Schotten, T. P. Nicholls, R. A. Bourne, N. Kapur, B. N. Nguyen and C. E. Willans, *Green Chem.*, 2020, **22**, 3358–3375.

7 (a) T. Fuchigami, M. Atobe and S. Inagi, *Fundamentals and Applications of Organic Electrochemistry: Synthesis, Materials, Devices*, Wiley-VCH, Weinheim, 1st edn, 2015, p. 217; (b) N. Elgrishi, K. J. Rountree, B. D. McCarthy, E. S. Rountree, T. T. Eisenhart and J. L. Dempsey, *J. Chem. Educ.*, 2018, **95**, 197–206.

8 C. Costentin and J.-M. Savéant, *Proc. Natl. Acad. Sci. U. S. A.*, 2019, **116**, 11147–11152.

9 (a) A. M. Couper, D. Pletcher and F. C. Walsh, *Chem. Rev.*, 1990, **90**, 857–865; (b) D. M. Heard and A. J. J. Lennox, *Angew. Chem., Int. Ed.*, 2020, **59**, 18866–18884.

10 For a dedicated review, see: J. P. Barham and B. König, *Angew. Chem., Int. Ed.*, 2020, **59**, 11732–11747.

11 For reviews comparing photo-, electro- and photoelectrochemical strategies, see: (a) R. H. Verschueren and W. M. De Borggraeve, *Molecules*, 2019, **24**, 2122; (b) J. Liu, L. Lu, D. Wood and S. Lin, *ACS Cent. Sci.*, 2020, **6**, 1317–1340.

12 For highlights, see: (a) L. Capaldo, L. L. Quadri and D. Ravelli, *Angew. Chem., Int. Ed.*, 2019, **131**, 17670–17672; (b) Y. Yu, P. Guo, J.-S. Zhong, Y. Yuan and K.-Y. Ye, *Org. Chem. Front.*, 2020, **7**, 131–135.

13 (a) H. Tateno, Y. Miseki and K. Sayama, *Chem. Commun.*, 2017, **53**, 4378–4381; (b) T. Li, T. Kasahara, J. He, K. E. Dettelbach and G. M. Sammis, *Nat. Commun.*, 2017, **8**, 390; (c) H. Tateno, Y. Iguchi, Y. Miseki and K. Sayama, *Angew. Chem., Int. Ed.*, 2018, **57**, 11238–11241; (d) L. Zhang, L. Liardet, J. Luo, D. Ren, M. Grätzel and X. Hu, *Nat. Catal.*, 2019, **2**, 366–373.

14 (a) R. Scheffold and R. Orlinski, *J. Am. Chem. Soc.*, 1983, **105**, 7200–7202; (b) F. Wang and S. Stahl, *Angew. Chem., Int. Ed.*, 2019, **58**, 6385–6390.

15 (a) H. Yan, Z.-W. Hou and H.-C. Xu, *Angew. Chem.*, 2019, **131**, 4640–4643; (b) X.-L. Lai, X.-M. Shu, J. Song and H.-C. Xu, *Angew. Chem., Int. Ed.*, 2020, **59**, 10626–10632; (c) H. Huang and T. H. Lambert, *Angew. Chem., Int. Ed.*, 2019, **59**, 658–662; (d) W. Zhang, K. L. Carpenter and S. Lin, *Angew. Chem., Int. Ed.*, 2020, **59**, 409–417; (e) Y. Qiu, A. Scheremetjew, L. H. Finger and L. Ackermann, *Chem. – Eur. J.*, 2020, **26**, 3241–3246.

16 (a) J.-C. Moutet and G. Reverdy, *Tetrahedron Lett.*, 1979, **20**, 2389–2393; (b) J.-C. Moutet and G. Reverdy, *J. Chem. Soc., Chem. Commun.*, 1982, 654–655; (c) S. S. Shukla and J. F. Rusling, *J. Phys. Chem.*, 1985, **89**, 3352–3358; (d) B. R. Egging and P. K. J. Robertson, *J. Chem. Soc., Faraday Trans.*, 1994, **90**, 2249–2256; (e) H. Huang, Z. M. Strater, M. Rauch, J. Shee, T. J. Sisto, C. Nickolls and T. H. Lambert, *Angew. Chem., Int. Ed.*, 2019, **58**, 13318–13322; (f) H. Kim, H. Kim, T. H. Lambert and S. Lin, *J. Am. Chem. Soc.*, 2020, **142**, 2087–2092; (g) N. G. W. Cowper, C. P. Chernowsky, O. P. Williams and Z. K. Wickens, *J. Am. Chem. Soc.*, 2020, **142**, 2093–2099.

17 R. I. Walter, *J. Am. Chem. Soc.*, 1955, **77**, 5999–6002.

18 For selected reviews, see: (a) Z. Ning and N. Tian, *Chem. Commun.*, 2009, 5483–5495; (b) P. Cias, C. Slugovc and G. Gescheidt, *J. Phys. Chem. A*, 2011, **115**, 14519–14525.

19 For selected reviews, see: (a) E. Steckhan, *Angew. Chem., Int. Ed. Engl.*, 1986, **25**, 683–701; (b) R. Francke and R. D. Little, *Chem. Soc. Rev.*, 2014, **43**, 2492–2521.

20 For selected examples, see: (a) T. Fuchigami, M. Tetsu, T. Tajima and H. Ishii, *Synlett*, 2001, 1269–1271; (b) X. Wu, A. P. Davis and A. J. Fry, *Org. Lett.*, 2007, **9**, 5633–5636; (c) C.-Y. Cai and H.-C. Xu, *Nat. Commun.*, 2018, **9**, 3551.

21 (a) R. A. Pabon, D. J. Bellville and N. L. Bauld, *J. Am. Chem. Soc.*, 1983, **105**, 5158–5159; (b) D. W. Reynolds, K. T. Lorenz, H. S. Chiou, D. J. Bellville, R. A. Pabon and N. L. Bauld, *J. Am. Chem. Soc.*, 1987, **109**, 4960–4968.

22 (a) P. S. Engel, A. K. M. M. Hoque, J. N. Scholz, H. J. Shine and K. H. Whitmire, *J. Am. Chem. Soc.*, 1988, **110**, 7880–7882; (b) L. Eberson and B. Olofsson, *Acta Chem. Scand.*, 1989, **43**, 698–701; (c) F. Ciminale, L. Lopez, A. Nacci, L. D’Accolti and F. Vitale, *Eur. J. Org. Chem.*, 2005, 1597–1603.

23 J. P. Barham, M. P. John and J. A. Murphy, *J. Am. Chem. Soc.*, 2016, **138**, 15482–15487.

24 N. A. Romero, K. A. Margrey, E. N. Tay and D. A. Nicewicz, *Science*, 2015, **349**, 1326–1330.

25 S. Das, P. Natarajan and B. König, *Chem. – Eur. J.*, 2017, **23**, 18161–18165.

26 S. Fukuzumi, J. Yuasa, N. Satoh and T. Suenobu, *J. Am. Chem. Soc.*, 2004, **126**, 7585–7594.

27 For detailed mechanistic discussions on the preference of certain substrates to undergo C–H activation *vs.* C–F substitution, see: V. A. Pistrutto, M. E. Schutzbach-Horton and D. A. Nicewicz, *J. Am. Chem. Soc.*, 2020, **142**, 17187–17194; also see ref. 15c.

28 M. Fujitsuka and T. Majima, *J. Photochem. Photobiol. C*, 2018, **35**, 25–37.

29 A. P. Demchenko, V. I. Tomin and P.-T. Chou, *Chem. Rev.*, 2017, **117**, 13353–13381.

30 Selected examples: (a) D. Gosztola, M. P. Niemczyk, W. Svec, A. S. Lukas and M. R. Wasielewski, *J. Phys. Chem. A*, 2000, **104**, 6545–6551; (b) C. Lu, M. Fujitsuka, A. Sugimoto and T. Majima, *J. Phys. Chem. C*, 2016, **120**, 12734–12741.

31 (a) J. A. Christensen, B. T. Phelan, S. Chaudhuri, A. Acharya, V. S. Batista and M. R. Wasielewski, *J. Am. Chem. Soc.*, 2018, **140**, 5290–5299; (b) J. Grilj, P. Buchgraber and E. Vauthey, *J. Phys. Chem. A*, 2012, **116**, 7516–7522; (c) J. Köhler, T. Quast, J. Buback, I. Fischer, T. Brixner, P. Nuernberger, B. Geiß, J. Mager and C. Lambert, *Phys. Chem. Chem. Phys.*, 2012, **14**, 11081–11089.

32 Detailed TAS studies are ongoing to further elucidate the nature of higher order excited states and the influences of precomplexation.

33 W. C. Neikam, G. M. Dimeler and M. M. Desmond, *J. Electrochem. Soc.*, 1964, **111**, 1190–1192.

34 L. Fajari, R. Papoular, M. Reig, E. Brillas, J. L. Jorda, O. Vallcorba, J. Rius, D. Velasco and L. Juliá, *J. Org. Chem.*, 2014, **79**, 1771–1777.

35 Here ‘symmetry breaking’ refers to the rotation of one N-aryl unit into conjugation with the N radical cation, resulting in a symmetry change of $C_3 \rightarrow C_2$ as proposed in the following references: (a) S. Amthor, B. Noller and C. Lambert, *Chem. Phys.*, 2005, **316**, 141–152; (b) Y. Su, X. Wang, L. Wang, Z. Zhang, X. Wang, Y. Song and P. P. Power, *Chem. Sci.*, 2016, **7**, 6514–6518. In ref. 35b the thermally-excited triplet state therein represents a dimeric analog of the TPA^+ s herein and, like $TCBPA^+$, displays a broad central singlet-type signal.

36 Drastic changes in the EPR spectra of TPA^+ s cannot be rationalized by solvent effects. Signals did not change in shape or shift when compared in DCM *vs.* MeCN *vs.* 0.1 M $^nBu_4N\cdot PF_6^-$ in MeCN. Arenes with similar viscosities give rise to different signals in the EPR of $TCBPA^+$ and anisotropy typical of solvent effects is not observed, see ESI.†

37 (a) C. A. Hunter and J. K. M. Sanders, *J. Am. Chem. Soc.*, 1990, **112**, 5525–5534; (b) C. R. Martinez and B. L. Iverson, *Chem. Sci.*, 2012, **3**, 2191–2201.

38 Endergonic binding is reported for systems involving π - π dispersion interactions under a similar level of theory, see: (a) L. M. da Costa, S. R. Stoyanov, S. Gusalov, X. Tan, M. R. Gray, J. M. Stryker, R. Tykwinski, J. W. de M. Carneiro and P. R. Seidl, *Energy Fuels*, 2012, **26**, 2727–2735; (b) A. Muraoka and M. Hayashi, *Chem. Phys. Lett.*, 2020, **748**, 137393. Reactions with a 20 kcal mol⁻¹ barrier are spontaneous at room temperature, see ref. 23.

39 (a) D. E. Williams and Y. Xiao, *Acta Crystallogr.*, 1993, **49**, 1; (b) M. O. Sinnokrot and C. D. Sherrill, *J. Am. Chem. Soc.*, 2004, **126**, 7690–7697; (c) M. Hajji, H. Mtiraoui, N. Amiri, M. Msaddek and T. Guerfel, *Int. J. Quantum Chem.*, 2019, **119**, e26000.

40 The arene centroid-centroid model (ref. 37a) was deemed lacking in the case of these complexes where arenes bridged both aromatics of the TPA^+ biphenyl unit. For rationalization of distances, see ESI.†

41 Similar DFT levels were successfully used to model π -stacking interactions of similar or very large systems, see ref. 38. The long-range-corrected functional ω B97XD includes a dispersion model and provides comparable results to expensive calculations for dispersion-type interactions, see: (a) T. Gao, T. Li, W. Li, L. Li, C. Fang, H. Li, L. Hu, Y. Lu and Z.-M. Su, *J. Cheminf.*, 2016, **8**, 24; (b) K. Wang, J. Lv and J. Miao, *Theor. Chem. Acc.*, 2015, **134**, 5.

42 CAM-B3LYP and ω b97XD have been used for TD-DFT of similar carbazole radical cations. Agreements between calculated and experimental UV-vis spectra are reasonable. For example, see: Z. Zara, J. Iqbal, K. Ayub, M. Irfan, A. Mahmood, R. A. Khera and B. Eliasson, *J. Mol. Struct.*, 2017, **1149**, 282–298.

43 Calculated canonical MOs resembled similar reported compounds, see: B. C. Lin, C. P. Cheng and Z. P. M. Lao, *J. Phys. Chem. A*, 2003, **107**, 5241–5251.

44 R. L. Martin, *J. Chem. Phys.*, 2003, **118**, 4775–4777.

45 Calculated from the sum of the ground state redox potential (*vs.* SCE) and the transition energy (eV) closest to 395 nm predicted by computational methods (average of both methods), see ESI.†

46 Dispersive precomplexation in this way requires a precursor photocatalyst, and is fundamentally distinct from formation of EDA complexes where the chromophore is generated *in situ* from two colorless precursors. For a review, see: (a) G. E. M. Crisenza, D. Mazzarella and P. Melchiorre, *J. Am. Chem. Soc.*, 2020, **142**, 5461–5476.

47 Further refinements to our working mechanistic proposal of precomplexation, including explicit solvent molecules, the possibility of multiple substrate arene binding and the excited state dynamics of precomplexes are ongoing and will be reported in due course.

48 N. Berg, S. Bergwinkl, P. Nuernberger, D. Horinek and R. M. Gshwind, Extended Hydrogen Bond Networks for Effective Proton-Coupled Electron Transfer (PCET) Reactions: The Unexpected Role of Thiophenol and Its Acidic Channel in Photocatalytic Hydroamidations, *J. Am. Chem. Soc.*, 2021, **143**, 724–735.

49 M. Giedyk, R. Narobe, S. Weiß, D. Touraud, W. Kunz and B. König, Photocatalytic activation of alkyl chlorides by assembly-promoted single electron transfer in microheterogeneous solutions, *Nat. Catal.*, 2020, **3**, 40–47.

50 J. Kaur, A. Shahin and J. P. Barham, Photocatalyst-Free, Visible-Light-Mediated C(sp³)-H Arylation of Amides via a Solvent-Caged EDA Complex, *Org. Lett.*, 2021, DOI: 10.1021/acs.orglett.1c00132.

