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4,7-Diarylbenzo[c][1,2,5]thiadiazoles as fluorophores and visible light organophotocatalysts†‡

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Electron donor–acceptor (D–A) systems based on the benzo[c][1,2,5]thiadiazole (**BTZ**) motif have been extensively researched for use in photovoltaics or as fluorescent sensors. However, their use as potential visible-light organophotocatalysts has not received any in-depth study. Here we report the synthesis, characterisation, and application of a library of 26 D–A compounds based on the **BTZ** group. By varying the donor groups whilst keeping the **BTZ** acceptor group the same, we have been able to systematically modify the photocatalyst's optoelectronic and photophysical properties. These photocatalysts were then validated using a Minisci-type decarboxylative alkylation of electron deficient heteroarenes as a test reaction under both batch and continuous flow conditions.

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Introduction

Organic π -conjugated electron donor–acceptor (D–A) systems have firmly established themselves as a powerful design motif for applications such as organic photovoltaics, fluorophores and photocatalysts.^{1–4} Among the endless combinations of donor and acceptor groups, the benzo[c][1,2,5]thiadiazole (**BTZ**) motif has received appreciable attention as a strongly electron accepting moiety, primarily for use in photovoltaic applications but also as fluorescent sensors or bioimaging probes for lipid droplets, mitochondria and plasma membranes.^{5–12} The **BTZ** group has also been researched for photocatalytic applications although this has mainly been limited to heterogeneous systems involving **BTZ** containing metal–organic frameworks

(MOFs),^{13–15} covalent organic frameworks (COFs),^{16–18} and conjugated porous polymers (CPPs).^{19–23}

In recent years, anthropogenic climate change and globally depleting resources have stimulated discussion within the chemical community about methods to perform environmentally sustainable chemistry, including the use of light as a “traceless” reagent. Concordant with this has been the rise of photoredox catalysis as a powerful tool for enabling organic transformations, with current state of the art photoredox catalysts based on iridium and ruthenium polypyridyl complexes.^{24,25} However, the cost and scarcity of the precious metals they contain calls into question the long term environmental sustainability of photocatalysis and contradicts the green ideals that modern photochemistry strives towards.²⁶ Extensive research has been conducted into the use of organophotocatalysts as alternatives but these are mainly limited to a few chemical families including carbazoyl cyanobenzenes,²⁷ benzophenones, xanthenes, thiazines and acridiniums.^{25,28} Recent examples have highlighted that the properties of various families of organophotocatalysts can be tailored towards more favorable redox potentials and photophysical properties. For example, Miyake and coworkers have demonstrated that installation of electron donating groups to phenoxazine photocatalysts was more conducive towards high reduction potentials than electron withdrawing groups.²⁹ Similarly, Zeidler and coworkers have demonstrated that carbazoyl cyanoarenes can be tuned with the goal of achieving both strongly oxidizing and reducing photocatalysts from a common scaffold.²⁷

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†Dedicated to the memory of Professor John S. Fossey.

‡Electronic supplementary information (ESI) available: Full synthesis and characterisation of photocatalyst library and substrate scope, details of computational calculations, single crystal X-ray structures. CCDC 2179269–2179281 and 2179284. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d2qo01316a>



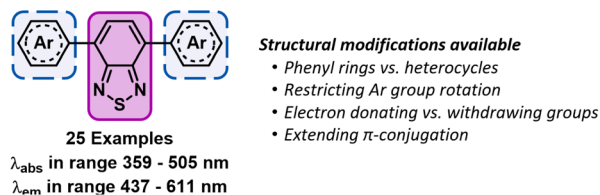


Fig. 1 Structure of the 4,7-diarylBTZ photocatalysts with the BTZ group highlighted in pink (solid line) and the aryl groups highlighted blue (dashed lines).

Inspired by these systematic approaches towards tailoring the properties of photocatalysts, we sought to perform a similar study of photocatalysts based on the 4,7-diarylBTZ motif (see Fig. 1).⁵ While there has been selected reports of homogeneous BTZ photocatalysis arising from intramolecular charge transfer between the donor and acceptor units, there has been no in-depth exploration of how their photophysical properties and excited state redox potentials can be altered by manipulating available structural levers.^{30–35} The reaction that was selected to test this library of BTZ photocatalysts was the decarboxylative C–H functionalisation of heteroarenes, previously explored by Glorius and coworkers.³⁶ This Minisci-type reaction was noted to provide a direct method to alkylate a wide variety of substrates, including pharmaceutical analogues, using visible light and without the need for pre-functionalisation of substrates.

While the research conducted into homogeneous BTZ photocatalysis is limited in scope, more in-depth research has been devoted towards their fluorescence properties.^{5,7} The focus of research into BTZ fluorescence includes methods to tune the wavelengths of emission (particularly towards lower energy red light) and spans applications such as sensors,³⁷ bioimaging agents,³⁸ thermally activated delayed fluorophores,³⁹ and organic LED devices.⁴⁰ With this in mind, we sought to examine the fluorescence properties of our library of BTZ molecules in parallel with assessing their photocatalytic performance.

Herein we describe the synthesis of a library of 26 organo-photocatalysts based on the BTZ motif and examine the effect that different structural modifications had on their optoelectronic and photophysical properties, with specific reference to tuning their absorption, fluorescence, and redox properties. The BTZ photocatalyst library was then validated using the decarboxylative C–H functionalisation of heteroarenes as a test reaction.

Results and discussion

Design and synthesis of photocatalyst library

In the design of this BTZ photocatalyst library (Fig. 1), we aimed to investigate how different structural modifications to the parent compound 4,7-diphenylbenzo[c][1,2,5]thiadiazole (pH-BTZ) influenced the compounds photophysical properties. In this study we elected to keep the BTZ core (the acceptor)

constant across all photocatalysts and focus solely on investigating the effects of changing the donor aryl groups. With this in mind, our general design considerations included targeting changes that we envisaged would allow access to a wide window of redox potentials and bathochromically shift the absorption of light from ultraviolet to visible wavelengths. This led us to investigate the effect that introducing heavy atoms or electron donating/electron withdrawing substituent groups, extending π -conjugation, restricting rotation of the aryl groups, and replacing the phenyl rings with heterocycles had on the photocatalyst. The structures of all of the photocatalysts investigated as part of this study are shown in Fig. 2 along with the abbreviations used throughout this study.

The starting point for the synthesis of the photocatalyst library was 4,7-dibromobenzo[c][1,2,5]thiadiazole (**Br₂BTZ**), as this can be cheaply synthesised through high yielding literature procedures on a large scale.^{41,42} With other common organophotocatalysts, such as acridiniums,²⁸ BODIPYs,⁴³ and phenothiazines,⁴⁴ derivatisation can often be difficult or labourious: this can be a lengthy process as the syntheses often involve multiple steps. In comparison, by mainly employing Suzuki–Miyaura cross coupling we were able to take advantage of the diversity of aryl boronic acids that were commercially available and synthesise a large proportion of our library in a single step with moderate to excellent yields. For selected photocatalysts, additional reactions such as bromination, iodination or Miyaura borylation with a subsequent Suzuki–Miyaura cross couplings were also utilised (see section 2.2 of the ESI†). Using these reactions, we were able to synthesise a total of 26 different prospective photocatalysts to investigate: in each case, UV-Vis. absorption/emission spectra and cyclic voltammograms were recorded and photostability assessed before moving onto testing their photocatalytic performance.

Absorption and emission properties

For each of the BTZ photocatalyst that was synthesised, their UV-Vis absorption and visible emission spectra were recorded in chloroform solution (see Table 1 and Fig. S30–S55 in the ESI†). In the case pH-BTZ, which as the parent photocatalyst was used as a reference for all modifications, the maximum of light absorption (λ_{abs}) and emission (λ_{em}) were located at 380 and 482 nm respectively. The introduction of electron donating substituent groups in the *para*-position led to slight bathochromic shifts in both absorption and emission. The largest bathochromic shift in both absorbance and emission was observed with the introduction of a *para*-methoxy group (λ_{abs} of 409 and λ_{em} of 525 nm respectively). This shift in absorption and emission can be rationalised in terms of an increase in both the lowest unoccupied molecular orbital (LUMO) and, to a greater degree, the highest occupied molecular orbital (HOMO) energy levels of the photocatalyst, with the overall effect being a reduction in the HOMO–LUMO energy gap. Electron withdrawing groups produced the opposite result, leading to hypsochromic shifts in light absorption and emission. Of the electron withdrawing groups tested, the largest hypsochromic shift in absorption was produced by the intro-



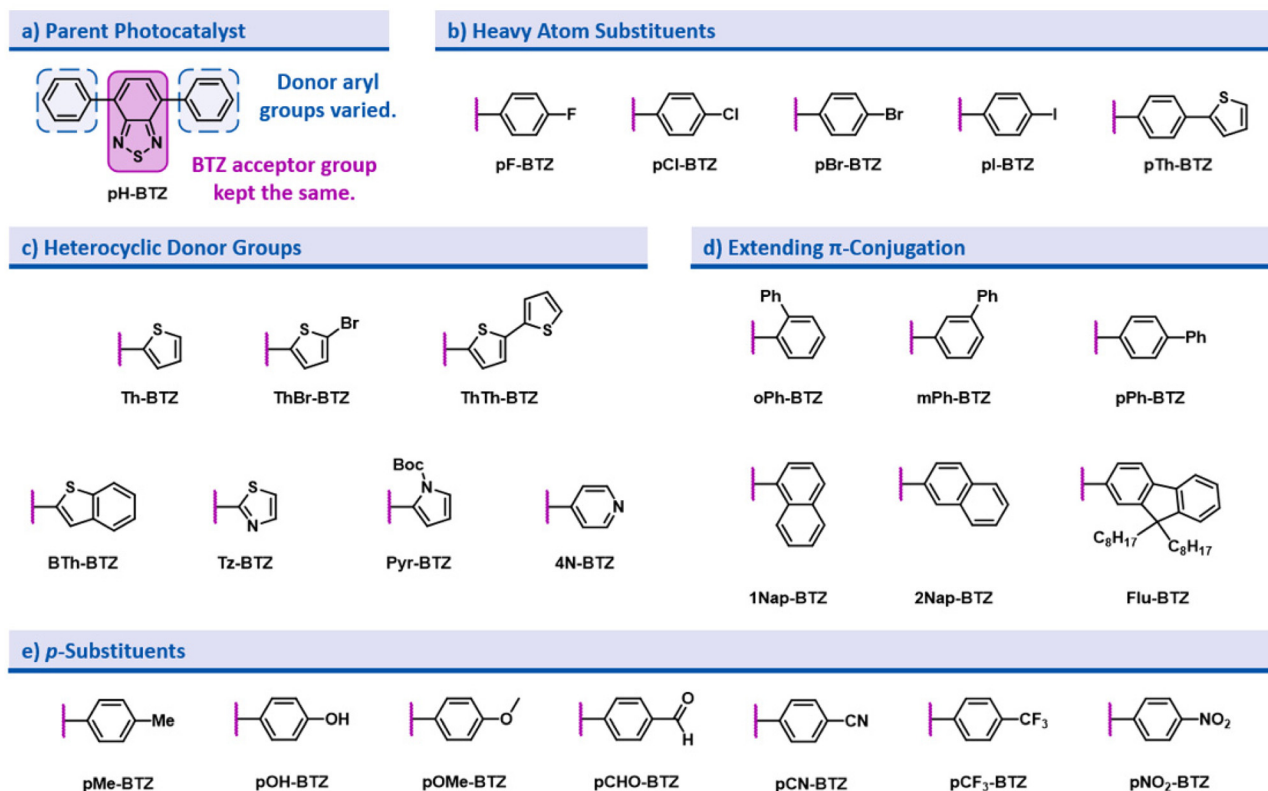


Fig. 2 Summary of the different derivative structures of the parent photocatalyst pH-BTZ (a) that were investigated, and the abbreviations used. The different types of modifications made to pH-BTZ were (b) the addition of heavy atoms, (c) replacing the phenyl ring with heterocycles, (d) extension of π -conjugation and (e) the addition of *p*-substituents.

Table 1 Photophysical properties of the BTZ photocatalysts

Entry	Compound	$\lambda_{\text{abs}}^a/\text{nm}$	$\lambda_{\text{em}}^{a,b}/\text{nm}$	Stokes shift/nm	$\epsilon_M^{a,c}/\times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$	PLQY ^{a,d} /%
1	pH-BTZ	380	482	102	7.5 \pm 0.1	86.4
2	pF-BTZ	381	495	114	7.5 \pm 0.3	93.3
3	pCl-BTZ	383	490	107	10.4 \pm 0.2	94.3
4	pBr-BTZ	381	492	111	9.8 \pm 0.4	93.4
5	pI-BTZ	384	475	91	13.7 \pm 0.2	91.9
6	pTh-BTZ	409	525	116	17.9 \pm 0.6	86.9
7	Th-BTZ	446	552	106	12.1 \pm 0.4	88.1
8	ThBr-BTZ	455	556	101	17.4 \pm 0.5	91.4
9	ThTh-BTZ	505	611	106	22.8 \pm 0.2	58.0
10	BTh-BTZ	450	548	98	16.6 \pm 0.3	96.2
11	Tz-BTZ	428	514	86	17.3 \pm 0.6	92.7
12	Pyr-BTZ	397	529	132	5.6 \pm 0.1	73.4
13	4N-bTZ	359	437	78	12.3 \pm 0.3	66.1
14	oPh-BTZ	373	486	113	6.5 \pm 0.2	100
15	mPh-BTZ	381	479	98	10.2 \pm 0.4	92.3
16	pPh-BTZ	398	502	104	18.2 \pm 0.3	97.3
17	1Nap-BTZ	376	504	128	5.6 \pm 0.1	23.6
18	2Nap-BTZ	396	509	113	15.5 \pm 0.3	66.2
19	Flu-BTZ	420	526	106	19.9 \pm 0.1	96.8
20	pMe-BTZ	391	499	108	13.0 \pm 0.4	92.7
21	pOH-BTZ	404	524	120	11.3 \pm 0.1	89.1
22	pOMe-BTZ	409	525	116	9.8 \pm 0.4	94.0
23	pCHO-BTZ	377	464	87	25.2 \pm 0.2	85.2
24	pCN-bTZ	372	457	85	18.3 \pm 0.3	93.9
25	pCF ₃ -BTZ	369	459	90	9.4 \pm 0.3	88.2
26	pNO ₂ -BTZ	377	455	78	23.1 \pm 1.0	4.87

^a In CHCl₃ solution. ^b Excitation wavelength was the wavelength of maximum absorption of the photocatalyst. ^c The molar attenuation coefficient (ϵ_M) was measured as described in section 4.2 of the ESI.† ^d Absolute PLQY.



duction of trifluoromethyl group (λ_{abs} of 369 nm) while for emission this was produced by the nitro group (λ_{em} of 455 nm).

The second type of modification that produced interesting shifts in absorption and emission was extending the π -conjugated structure of **pH-BTZ**. For example, replacing the 4,7-phenyl groups with 2-naphthyl groups in **2Nap-BTZ** introduced a bathochromic shift in both light absorption and emission (λ_{abs} of 396 and λ_{em} of 509 nm). In comparison, the isomeric **1Nap-BTZ** demonstrated a slight hypsochromic shift in absorption (376 nm), which could arise from a disruption of π -conjugation by introducing the sterically bulky 1-naphthyl groups. Some evidence of this was provided by computational modelling of the energy barrier to rotation of **1Nap-BTZ** and **pH-BTZ** (see Fig. S88 and S90†). As can be seen in these figures, the lowest energy conformation for **pH-BTZ** was determined to be when the phenyl groups sat at around 40° to the **BTZ** group. By comparison, the minimum energy for **1Nap-BTZ** was around 70°, with an energy barrier to rotation of the 1-naphthyl groups of around 63 kJ mol⁻¹ (compared to 8 kJ mol⁻¹ for **pH-BTZ**). Similar trends were also observed by the introduction of phenyl groups in the *ortho*-, *meta*- and *para*-positions of **pH-BTZ**. In the case of **pPh-BTZ**, the phenyl group situated in the *para*-position extended π -conjugation and introduced a bathochromic shift in light absorption (with λ_{abs} of 398 nm). Conversely, the *ortho*-isomer **oPh-BTZ** featured a hypsochromic shift in absorption (λ_{abs} at 373 nm), which could also be explained by the steric bulk of the *ortho*-phenyl groups disrupting π -conjugation (see Fig. S91†). With the *meta*-isomer, **mPh-BTZ**, the phenyl group was neither in a position to suitably extend π -conjugation in a meaningful way or twist the molecule, leading to no significant change in either absorption or emission.

The most dramatic changes in the absorption and emission spectra were observed when the phenyl rings were replaced with various heterocycles. For example, replacing the phenyl rings of **pH-BTZ** with 2-thienyl groups, to give **Th-BTZ**, resulted in a significant bathochromic shift in λ_{abs} to 446 nm and λ_{em} to 552 nm, again largely driven by an increase in the HOMO energy relative to the LUMO. The addition of a second 2-thienyl group to form **ThTh-BTZ** further shifted λ_{abs} to 505 nm, which represents a total bathochromic shift in absorption from **pH-BTZ** of 125 nm. **ThTh-BTZ** also featured the lowest energy emission of the entire series, with λ_{em} at 611 nm. This can be contrasted with **4N-BTZ**, with the electron deficient 4-pyridyl group, which exhibited the highest energy absorption (λ_{abs} of 359 nm) and emission (λ_{em} of 437 nm) in the entire series.

The dependence of λ_{abs} and λ_{em} in various polar and non-polar solvents was also assessed for **pH-BTZ** (see Table S1†). It was observed that, while λ_{abs} remained mainly unaffected by the solvent used λ_{em} varied by up to 31 nm, with more polar solvents leading to bathochromic shifts in emission (see Fig. S56 and S57†). The dependence of the spectroscopic properties of **pH-BTZ** upon solvent can be visualised through a Lippert–Mataga plot of Stokes' shift against the solvent

polarity parameter (see Fig. S58†), where a straight-line plot was obtained.⁴⁵ The increased Stokes' shift in more polar solvent suggests that the excited state of **pH-BTZ** has a higher dipole moment than its ground state, indicating a redistribution of charge following photoexcitation and the formation of an intramolecular charge transfer state.^{46–48} A linear relationship between the observed Stokes' shift and solvent polarity was also attained when the Reichardt $E_{\text{T}}(30)$ parameter was used as an alternative measure of solvent polarity (see Fig. S59†).^{49,50} Absolute photoluminescence quantum yields (PLQY) in chloroform solution were also measured for the entire library, with values in the range of 4.87–100% observed (see Table 1).

Photostability of BTZ photocatalysts

The photostability of each **BTZ** photocatalyst was assessed by irradiating a solution of the photocatalyst dissolved in oxygenated CDCl₃ for 18 hours and comparing ¹H NMR spectra recorded before and after. Most of the library exhibited good photostability, with little to no photobleaching observed during this time. Only **Pyr-BTZ**, **2Nap-BTZ** and **pCHO-BTZ** exhibited significant photodecomposition (see Table S3†).

Electrochemical potentials

To assess the prospective performance of the library towards photoredox catalysis, cyclic voltammetry was performed on each photocatalyst to extract their ground state oxidation and reduction potentials. The S_0 ($\nu = 0$) \rightarrow S_1 ($\nu = 0$) energy difference ($E_{0,0}$) was estimated from the intercept between normalised absorption and emission spectra, then used to predict the excited state redox potentials as described by Romero and Nicewicz.²⁵ Due to the low solubility of selected **BTZ** photocatalysts in acetonitrile (the typical solvent for cyclic voltammetry), DCM was used instead as this provided better dissolution of the photocatalysts whilst still allowing a wide range of potentials to be assessed (see Table S35 and Fig. S94–118†).⁵¹

In general, it was observed that the excited state reduction potentials (E_{red}^*) were far more sensitive towards changing the aryl donor groups than the excited state oxidation potentials (E_{ox}^*) were. In comparison to **pH-BTZ** ($E_{\text{red}}^* = 1.42$ V), E_{red}^* was increased by the addition of electron withdrawing substituents onto the phenyl ring (*p*-CHO, *p*-CN, *p*-CF₃ and *p*-NO₂). The most extreme example of this was with the nitro-substituted **pNO₂-BTZ**, which exhibited an E_{red}^* of 1.91 V. Similarly, replacing the phenyl rings with electron deficient heterocycles such as the 4-pyridyl groups of **4N-BTZ** or the 2-thiazolyl groups of **Tz-BTZ** also increased E_{red}^* from 1.42 V to 1.88 V and 1.51 V respectively. Introducing electron donating groups (*p*-OH, *p*-OMe and *p*-Me) or electron rich heterocycles (such as pyrrole or thiophene rings) produced the opposite effect and led to a reduction in E_{red}^* . Overall, the susceptibility of E_{red}^* towards aryl group derivatisation allowed a wide window of potentials in the range of 0.95–1.91 V to be covered. The maximum E_{red}^* of 1.91 V attained for **pNO₂-BTZ** is higher than some common organophotocatalysts, such as thiazines, BODIPYs and carbazoyl cyanobenzenes, and approaches the E_{red}^* displayed by



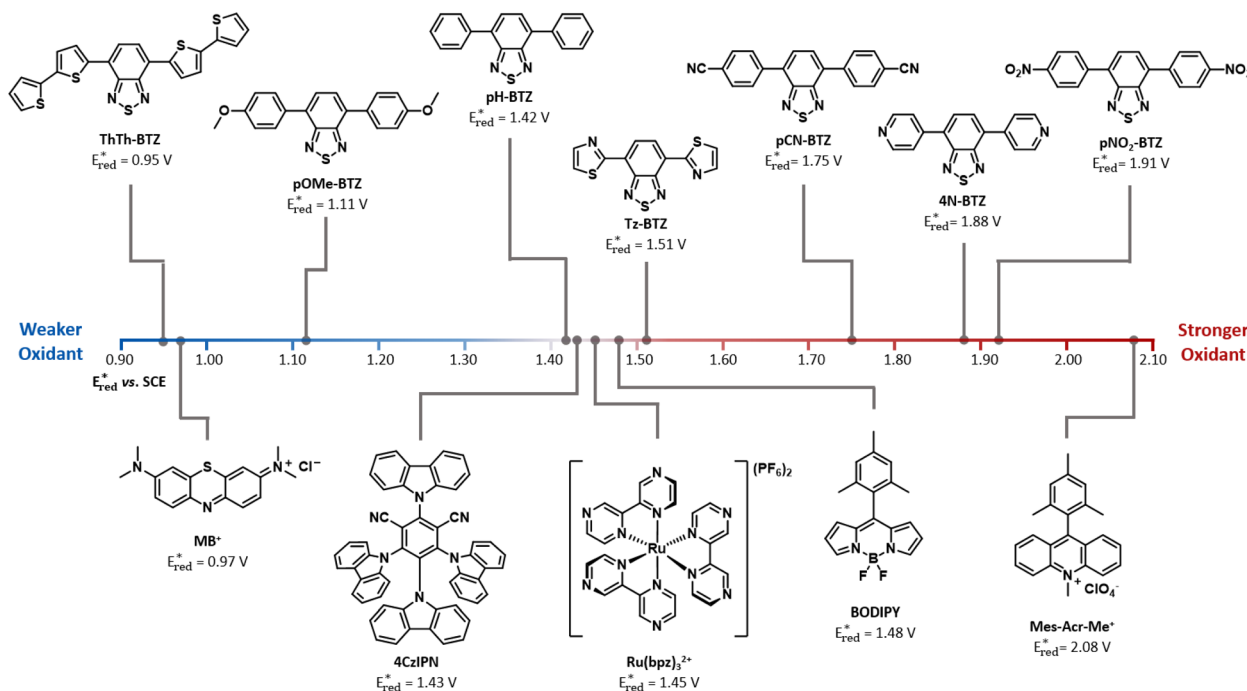


Fig. 3 Comparison between the excited state reduction potentials (E_{red}^*) of selected BTZ photocatalysts (above axis) and common photocatalysts discussed in literature (below axis).^{27,28,52–54}

some widely utilised acridinium photocatalysts (for comparison the E_{red}^* of **Mes-Acr-Me⁺** is 2.08 V) (Fig. 3).^{27,28,52–54}

In comparison to the wide range of E_{red}^* , donor group variation produced a narrower window of E_{ox}^* , with values in the range of -1.11 to -1.49 V, but no clear correlation between the type of modification made and the value of E_{ox}^* . In addition, several of the photocatalysts tested underwent chemical reactions at positive potentials during cyclic voltammetry.

Computational results

In order to supplement the experimental characterisation, density functional theory (DFT) calculations were also carried out on all members of the photocatalyst library. Investigation of the rotational barriers around the donor-acceptor dihedral angles were undertaken *via* calculation of fully relaxed scans to determine global geometric minima (Fig. S88–92[†]), revealing the distinct difference in the geometric behavior of phenyl and thiophene based donor groups. Photocatalysts with phenyl-based donor groups possessed a barrier of approximately 8 – 10 kJ mol^{−1} with geometric minima at 45 and 135° , showing a preference to avoid a planar geometry. In comparison the thiophene-based donor groups show a geometric preference for a planar geometry in which the sulphur atom is orientated at a 180° dihedral angle to the sulphur of the acceptor group with a rotational barrier of 15 – 20 kJ mol^{−1}. A noted outlier in this trend was the 2-thiazolyl donor group (**Tz-BTZ**) which, while maintaining planarity, shows a preference to have the sulphurs orientated in the same direction (a dihedral of 0°) with a barrier of approx. 27 kJ mol^{−1}; this is likely due to the less favorable interactions between in-plane lone pairs situated

on the nitrogen atoms in both donor and acceptor groups when an angle of 180° was obtained. Alteration of the chalcogen atoms of the donor groups of **Th-BTZ** to either selenium or tellurium also resulted in a steady change in preference for a 0° dihedral angle with barriers of approximately 17 and 22 kJ mol^{−1} respectively (see Fig. S92[†]).

In addition to the experimental spectra presented, time dependent-DFT (TD-DFT) spectra were also obtained for each photocatalyst to further assess both their absorption and emission properties. A summary of the predicted absorption and emission maxima and a comparison to the experimental properties can be found in Table S4.[†] Computational absorption spectra were shown to be in strong agreement with those derived experimentally while the emission spectra obtained showed marginally less agreement (see Fig. S60–85 and Table S4[†]). Computational analysis revealed that the lowest energy absorption band observed in each spectrum was characterised by a single state, S_1 , of the photocatalyst. In comparison, the second strong absorption band was characterised by states ranging from S_2 to S_{10} , depending on the photocatalyst involved.

Natural transition orbitals (NTOs) corresponding to the excitations to the bright states responsible for the first two peaks of the absorption spectra were calculated for all synthesised photocatalysts and can be found in Tables S5–S30.^{†55} A general observation was that the hole orbital for the vast majority of bright transitions presented character on both the aryl donors and the benzo-portion of the **BTZ** acceptor group (see Fig. 4). In comparison, the particle orbital for the $S_0 \rightarrow S_1$ transition shows movement of charge from the aryl groups,



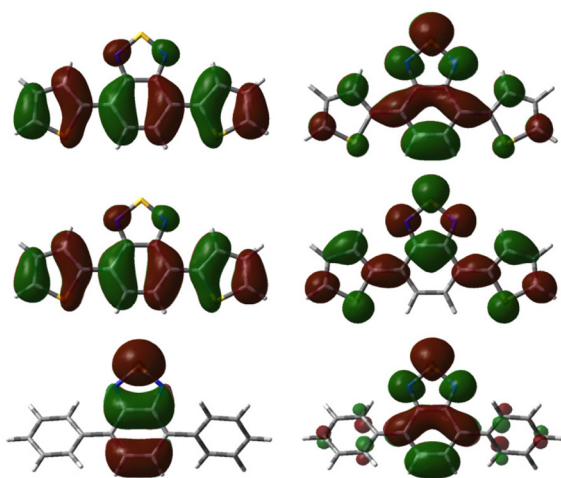


Fig. 4 NTO representations of the hole (left) and particle (right) orbitals of the bright states responsible for the two lower energy peaks observed in the absorption spectra. The S_1 state of **Th-BTZ** (top) represents the common transition responsible for the lowest energy peak while the S_2 transitions of **Th-BTZ** and **pTh-BTZ** (middle and bottom, respectively) represent the two different transitions that may contribute to the second peak.

becoming localised over the entirety of the **BTZ** acceptor group (e.g., $S_0 \rightarrow S_1$ **Th-BTZ**). The particle orbital of the $S_0 \rightarrow S_3$ transition of the **Th-BTZ** compound provides an example of the dominant transition in the second spectral peak in which less character is removed from the aryl groups and, instead of consolidating to the entire **BTZ** moiety, isolates to the 5-membered portion of the moiety. These results further reinforce the presence of an intramolecular charge transfer mechanism during light absorption, with electron density transferring from the aryl donor groups that constitute part of the hole orbital to the particle orbital localised on the **BTZ** group.

Single crystal X-ray structures

For a number of the **BTZ** photocatalysts investigated as part of this library, we were able to grow single crystals suitable for X-ray diffraction and subsequently determine their crystal structure. The crystal structures of **pF-BTZ**, **pCl-BTZ**, **pI-BTZ**, **ThTh-BTZ**, **BTh-BTZ**, **oPh-BTZ**, **mPh-BTZ**, **pPh-BTZ**, **1Nap-BTZ**, **2Nap-BTZ**, **pMe-BTZ**, **pOH-BTZ** and **pCF₃-BTZ** are all presented in the ESI (see Fig. S2–S29†) and stored in the Cambridge Structural Database (CCDC 2179269–2179281†). Unsurprisingly, most of the *para*-substituted **BTZ** photocatalysts possessed fairly similar structures. Due to the low solubility of **pOH-BTZ** in most solvents, the single crystal had to be grown from THF solution, which led to the formation of a solvate consisting of **pOH-BTZ** molecules engaged in hydrogen bonding with the THF solvent molecule.

BTh-BTZ and **ThTh-BTZ** were both observed to be completely planar in the crystal state, which is consistent with the previously reported crystal structure of **Th-BTZ**.⁵⁶ The planar structure of **BTh-BTZ** also echoes the preference to also be planar in solution as we determined using DFT calculations

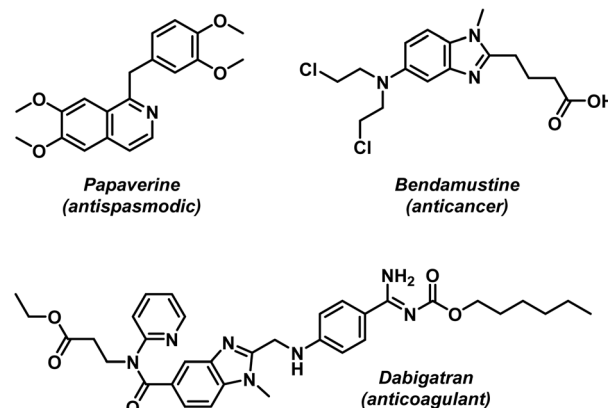
(see above). The structures of **BTh-BTZ** and **ThTh-BTZ** also exhibited some disorder regarding the relative orientation of the thiophene rings.

Photocatalytic application of BTZ library

To test our **BTZ** photocatalysts, we selected the alkylation of electron deficient heteroarenes *via* a Minisci-style C–H radical functionalisation as a test photoredox reaction.⁵⁷ Nitrogen containing heteroarenes, such as quinolines, pyridines, quinazolines *etc.*, are abundant structural motifs in nature that have become commonplace features among various pharmaceutical compounds.⁵⁸ As many of these heteroarenes feature alkyl regions, including examples such as Papaverine, Bendamustine and Dabigatran (Fig. 5a), several methods for selective alkylation have been explored by various groups (Fig. 5b).

The classical Minisci reaction generates alkyl radicals (R^\bullet) *via* a Ag(I) catalysed decarboxylation of carboxylic acids using $S_2O_8^{2-}$ as the oxidant.⁵⁹ The scope of possible R^\bullet sources has since been expanded to include a wide variety of functional groups, including aryl boronic acids and sulfinates.^{60–62} However, such methods often present harsh conditions, like the conventional Minisci coupling, or require pre-functionalisation of substrates. Barriault and co-workers have developed a method to alkylate heteroarenes using R^\bullet generated photocatalytically from widely available alkyl bromides.⁶³ However, this

a) Alkylated Heteroarenes in Pharmaceutical Compounds



b) Previous work

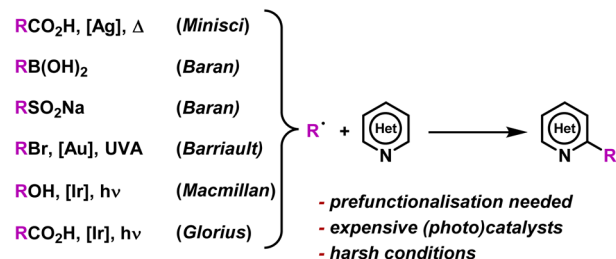


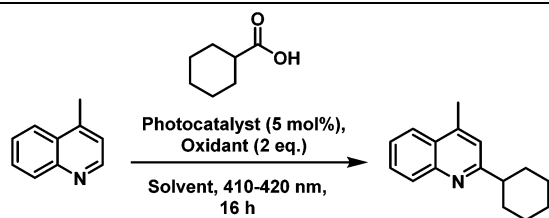
Fig. 5 (a) Common pharmaceutical molecules containing alkylated heterocycles. (b) Examples of previous methods to alkylate electron deficient heterocycles.^{36,59–64}



method utilised high loadings of gold catalysts and UVA irradiation (315–400 nm). The high energy UV light could lower the functional group tolerance of this protocol, a problem that would be remedied by the switching to lower energy visible light. Methods for alkylation of heteroarenes *via* visible light photoredox catalysis have been reported by both Macmillan and coworkers and Glorius and coworkers, using alcohols and carboxylic acids as the R^\bullet sources respectively.^{36,64} While these methods both address the issues of harsh reaction conditions, low functional group tolerance and required pre-functionalisation associated with other protocols, expensive iridium-based photocatalysts were employed.

In order to test the suitability of **BTZ** photocatalysts towards a Minisci-type reaction, we adapted a protocol developed by Glorius and coworkers.³⁶ The reaction conditions were first optimised using **pH-BTZ** as the photocatalyst (Table 2, entry 1). Using lepidine as the substrate, cyclohexanecarboxylic acid (10 equivalents) as the alkyl radical source, ammonium persulfate (2 equivalents) as the oxidant and DMSO as the solvent, we were able to attain a conversion of 48% in 16 h using a 12 W LED module (410–420 nm). From these initial conditions, we then varied the choice of solvent and oxidant, although we were unable to improve from the initial conditions. Replacing the ammonium persulfate oxidant with potassium persulfate ($K_2S_2O_8$, entry 2) still permitted the reaction to occur, although with significantly diminished conversion.

Table 2 Optimisation of reaction conditions for the decarboxylative C–H functionalisation of lepidine using cyclohexanecarboxylic acid



Entry	Photocatalyst	Oxidant	Solvent	Conversion ^b (%)
1	pH-BTZ	$(NH_4)_2S_2O_8$	DMSO	48
2	pH-BTZ	$K_2S_2O_8$	DMSO	15
3	pH-BTZ	$(NH_4)_2S_2O_8$	DMF	N.R.
4	pH-BTZ	$(NH_4)_2S_2O_8$	MeCN	4
5	pH-BTZ	—	DMSO	<1
6 ^c	pH-BTZ	$(NH_4)_2S_2O_8$	DMSO	N.R.
7	—	$(NH_4)_2S_2O_8$	DMSO	N.R.
8 ^d	pH-BTZ	$(NH_4)_2S_2O_8$	DMSO	4
9	oPh-BTZ	$(NH_4)_2S_2O_8$	DMSO	53
10	mPh-BTZ	$(NH_4)_2S_2O_8$	DMSO	25
11	pPh-BTZ	$(NH_4)_2S_2O_8$	DMSO	N.R.
12	1Nap-BTZ	$(NH_4)_2S_2O_8$	DMSO	57
13	2Nap-BTZ	$(NH_4)_2S_2O_8$	DMSO	16
14 ^{e,f}	1Nap-BTZ	$(NH_4)_2S_2O_8$	DMSO	>99 (73)

^a Reaction conditions: lepidine (0.3 mmol), oxidant (0.6 mmol), cyclohexanecarboxylic acid (3 mmol), photocatalyst (0.015 mmol), solvent (3 mL), 12 W 410–420 nm LED, 16 h. ^b Conversion determined by ¹H NMR. ^c Reaction performed in the dark. ^d Reaction was performed without degassing the reaction mixture with nitrogen. ^e Reaction performed for 40 h. ^f Isolated yield in brackets. N.R. = no reaction.

Similarly, replacing DMSO with other solvents severely lowered the conversion (entries 3 and 4). A potential reason for this could be due to $S_2O_8^{2-}$ reportedly decomposing more readily in DMSO compared to other solvents to generate the reactive species.⁶⁵ Control experiments were also performed, showing that the reaction did not proceed in the absence of light, oxidant and photocatalyst (entries 5–7). Additionally, not degassing the reaction also drastically reduced the conversion attained (entry 8). We also observed that the conversion was drastically reduced by the addition of TEMPO or $CeCl_3$, which would act as radical traps for R^\bullet and the sulfate radical anion respectively.^{66,67} This would suggest the involvement of both the cyclohexyl and sulfate radicals in the reaction, in agreement with the mechanism previously proposed by Glorius and coworkers.³⁶

Using these initial conditions, we then proceeded to screen our photocatalyst library: the key results are summarised in Table 2, while all reaction conditions tested can be found in Table S36.† In most cases, derivatisation of **pH-BTZ** had a detrimental effect on the conversion of the reaction. Furthermore, no correlation could be observed between the conversion and the excited state oxidation or reduction potentials. This suggested that the reaction was not solely dependent on the optoelectronic properties of the photocatalyst but could also be influenced by other factors, such as the lifetime of the excited state. We observed that, of all of the modifications made, only limiting rotation of the aryl side group by introducing significant steric bulk resulted in any increase in conversion. Replacing the 4,7-phenyl groups of **pH-BTZ** with 1-naphthyl groups to form **1Nap-BTZ** increased the conversion attained to 57%. For comparison, the 2-naphthyl analogue, **2Nap-BTZ**, resulted in a conversion of 16%. These results were echoed by the installation of a phenyl ring in either the *ortho*-, *meta*- or *para*-positions. The highest conversion of 53% was produced by **oPh-BTZ** while its *meta*- and *para*-isomers, **mPh-BTZ** and **pPh-BTZ**, gave conversions of 25% and 0% respectively. To further increase the conversion obtained using **1Nap-BTZ**, we subsequently increased the reaction time from 16 hours to 40 hours, leading to a conversion of >99%.

These initial optimisation results can be directly compared with the work performed by Glorius and coworkers to gauge how **BTZ** compared to established photoredox catalysts.³⁶ Using **Mes-Acr-Me⁺** and $[Ru(bpy)_3](PF_6)_2$ (at 2 mol% loading), the authors reported attaining conversions of 36 and 28% in 16 hours, suggesting that **1Nap-BTZ**, along with **pH-BTZ** and **oPh-BTZ**, are comparable to some well investigated organic and metal-based photoredox catalysts. However, the authors also reported that by employing $[Ir(dF(CF_3)ppy)_2(dtbbpy)](PF_6)_3$ (at 0.5 mol% loading) as the photocatalyst, 94% conversion could be obtained within 90 minutes. This indicates that **1Nap-BTZ** was not the optimum photocatalyst for the reaction shown in Table 2, however, this could be offset by the lower costs of using **BTZ** based organophotocatalysts compared to the more expensive $[Ir(dF(CF_3)ppy)_2(dtbbpy)](PF_6)_3$.

Using these optimised reaction conditions and employing **1Nap-BTZ** as our photocatalyst, we then proceeded to investi-

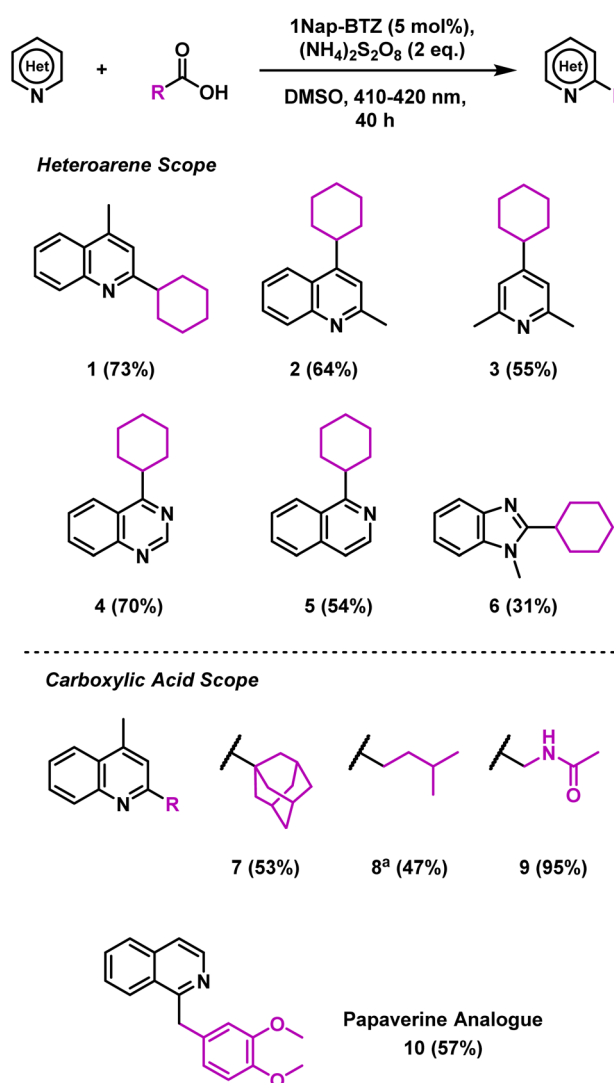


gate the scope of the reaction with selected heteroarenes and carboxylic acids (Scheme 1). Under the reaction conditions, various electron deficient nitrogen-based heterocycles including quinolines, pyridines, quinazolines and benzimidazoles (1–6) were successfully alkylated (31–73% yield). Expanding the scope of carboxylic acids revealed that the reaction was suitable for installing primary, secondary and tertiary alkyl groups (1, 7 and 8) in good yields (47–73%). However, in the case of 8, additional oxidant was required to attain full conversion, possibly due to the lower stability of the primary radicals generated. *N*-Acetyl glycine was also employed as a carboxylic acid in this reaction, with a yield of 95% obtained (9), demonstrating that this reaction could also be extended to amino acids. To demonstrate the pharmaceutical relevance of this reaction, we also applied 1Nap-BTZ to the synthesis of ana-

logue of the antispasmodic drug Papaverine using isoquinoline and homoveratric acid (10, 57% yield).

Under the optimised reaction conditions 1Nap-BTZ required 40 hours to achieve complete conversion. To lower this reaction time, we also investigated performing this reaction under continuous flow conditions. By flowing the reaction mixture through narrow transparent tubes, a smaller path-length for the incident light was achieved in comparison to the performing the reaction in a vial. This resulted in increased and more uniform photon flux, which has been well reported to drastically accelerate photochemical reactions.^{68,69}

Using a commercial flow reactor (Easy-Photochem E-Series, Vapourtec), we began by testing the alkylation of lepidine using cyclohexanecarboxylic acid under analogous conditions to the optimised batch conditions (Table 2, entry 12). The reaction mixture was continually recirculated through a 5 mL coil of tubing at a rate of 1 mL min⁻¹ under irradiation from the same LED that was utilised in the batch reactions (Fig. 6). After 16 hours, a conversion of >99% conversion was achieved, which was a significant improvement compared to the 40 hours required under batch conditions. It should be noted that the external LED module could only irradiate the 5 mL coil from a single side, which is not the most effective means of delivering light. We were further able to reduce the time required by switching to the flow machines built in LED module (UV-150 reactor module equipped with a 420 nm LED, Fig. S121†), which has a power of 60 W compared to 12 W for the in-house built LED array. This light source is designed to irradiate the tubing from the middle of the reactor coil, which presents a more efficient 360° method of supplying photons to the reaction. Employing this module reduced the time required to achieve >99% conversion to 100 minutes, which is



Scheme 1 Scope of heteroarenes and carboxylic acids. Reaction conditions: heteroarene (0.3 mmol), (NH₄)₂S₂O₈ (0.6 mmol), carboxylic acid (3 mmol), 1Nap-BTZ (0.015 mmol), DMSO (3 mL), 12 W 410–420 nm LED, 40 h. Isolated yields shown in brackets. ^a4 equivalents of (NH₄)₂S₂O₈ used.

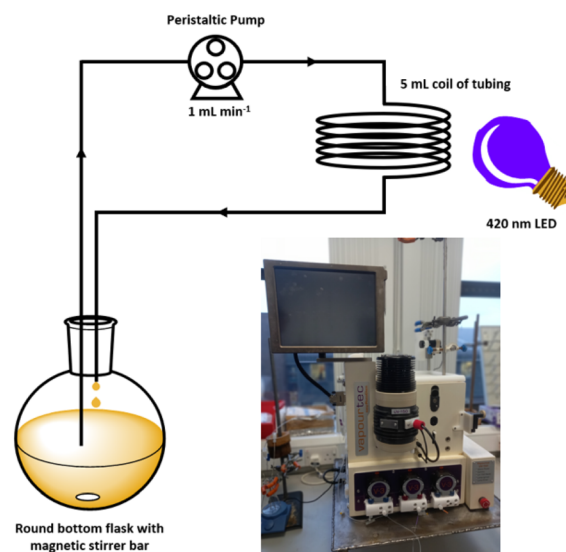


Fig. 6 Schematic representation of the continuous flow set up used to reduce the time required for the Minisci-type reaction. Inset picture shows the Easy-Photochem flow system from Vapourtec equipped with the UV-150 photoreactor.



an overall 24-fold decrease in the reaction time needed under the initial batch conditions.

Conclusions

In conclusion, a library of 26 electron donor–acceptor photocatalysts based on the **BTZ** group have been synthesised and studied, both experimentally and computationally. By varying the electron donor groups whilst keeping the **BTZ** constant, systematic variations in the photophysical and optoelectronic properties of the photocatalysts were obtained. This included tuning λ_{abs} to cover a range of 359–505 nm and λ_{em} to cover a range of 437–611 nm, with the introduction of various heteroarene groups producing the starkest variation in properties. Furthermore, E_{red}^* was observed to be sensitive to the nature of the electron donor groups present on the photocatalyst: the presence of electron rich donor groups reduced the value of E_{red}^* while electron deficient donor groups instead raised E_{red}^* . This ultimately allowed E_{red}^* to cover a range of values from 0.95 V for **ThTh-BTZ** to 1.91 V for **pNO₂-BTZ**. These photocatalysts were then implemented in a Minisci-type alkylation of heteroarenes *via* carboxylic acid decarboxylation, resulting in good to excellent yields. Although the reaction under batch conditions required a long time for completion (>99% in 40 hours using **1Nap-BTZ**), the use of continuous flow conditions allowed this time to be reduced to less than two hours, which further highlights the benefits of combining photoredox catalysis with flow technologies. In this study, we selected to solely focus on investigating the effect that changing the electron donor aryl groups had on the properties of the photocatalyst. This leaves the door open for further exploration of the strength of the acceptor **BTZ** core as a means to tune the properties of the photocatalyst, including substitution of the 5,6-positions, replacement of the sulphur atom with other chalcogen elements and/or extending the **BTZ** core with fused aromatic ring(s).

Conflicts of interest

There are no conflicts to declare.

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