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Pyridylphosphonium salts as alternatives to cyanopyridines in radical—radical coupling reactions†

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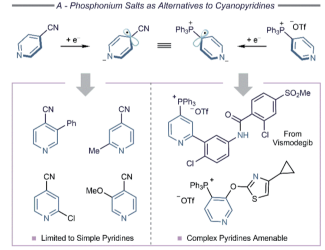
Radical couplings of cyanopyridine radical anions represent a valuable technology for functionalizing pyridines, which are prevalent throughout pharmaceuticals, agrochemicals, and materials. Installing the cyano group, which facilitates the necessary radical anion formation and stabilization, is challenging and limits the use of this chemistry to simple cyanopyridines. We discovered that pyridylphosphonium salts, installed directly and regioselectively from C–H precursors, are useful alternatives to cyanopyridines in radical–radical coupling reactions, expanding the scope of this reaction manifold to complex pyridines. Methods for both alkylation and amination of pyridines mediated by photoredox catalysis are described. Additionally, we demonstrate late-stage functionalization of pharmaceuticals, highlighting an advantage of pyridylphosphonium salts over cyanopyridines.

Modern photoredox catalysis and electrochemistry have enabled new synthetic methods that proceed *via* open-shell intermediates.¹ Under this regime, pyridine functionalization strategies have been developed where 4-cyanopyridines undergo single-electron reduction to form dearomatized radical species that couple with other stabilized radicals (Scheme 1A).² The cyano group is critical for efficient reactivity *via* pyridyl radical anions; alternatives such as 4-halopyridines more readily undergo elimination to pyridyl radicals after single-electron reduction resulting in a distinct set of coupling processes.³ We aimed to show that pyridylphosphonium salts could replicate the reactivity of cyanopyridines and allow a broader set of inputs into dearomatized pyridyl radical coupling reactions.⁴

Cyanopyridines have facilitated pyridine alkylation, allylation, and alkenylation reactions providing access to valuable building blocks for medicinal and agrochemical programs. The cyano group is essential for these methods, but a problem arises when applying this chemistry to complex pyridines, such as those found in pharmaceutical and agrochemical candidates. These structures are often devoid of pre-installed functional groups, and it is often challenging to install a cyano group from C–H precursors regioselectively. We envisioned pyridylphosphonium salts, regioselectively constructed from the C–H bonds of a diverse set of pyridines, could serve as alternatives to cyanopyridines. Herein, we report couplings between alkyl BF₃K salts and preliminary studies of carboxylic acids and amines with pyridylphosphonium salts, including late-stage

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B - Alkylation of Complex Pyridines via Phosphonium Salts

Scheme 1 Expansion of radical coupling reactions to complex pyridines.

functionalization of complex pyridine-containing pharmaceuticals using this strategy.

Recently, we reported a radical coupling reaction between a boryl-stabilized cyanopyridyl radical and a boryl-stabilized pyridylphosphonium radical.^{7a} The intermediate radicals arose via an unusual inner-sphere process that would be difficult to extend to other coupling reactions. A significant advance would be to show that pyridylphosphonium salts could function more generally as radical anion precursors and mimic the reactivity of cyano-pyridines. In particular, showing their viability in photoredox and electrochemical processes would translate to numerous synthetic transformations. To demonstrate this principle, we envisioned a redox-neutral alkylation reaction (Scheme 1B) via a radical coupling between radical zwitterion I, formed through single-electron reduction of a pyridylphosphonium salt ($E_{\rm p/2}^{\rm red} = -1.51$ V vs. SCE) and benzyl radical II, resulting from single-electron oxidation of a BF₃K salt $(E^{\text{red}} = +1.10 \text{ V } \nu \text{s. SCE for a primary benzylic salt).}^{8} \text{ Loss of }$ triphenylphosphine from dearomatized intermediate (III) would furnish the alkylated pyridine product. Notably, the redox events could invert, where the photocatalyst oxidizes the BF₃K salt first and reduces the pyridylphosphonium salt second, broadening the scope of amenable photocatalysts.

We began our investigation by examining a series of photocatalysts for the coupling reaction of phosphonium salt 1a, formed with complete regioselectivity for the 4-position from 2phenylpyridine, and benzylic BF₃K salt 2a under irradiation from a 455 nm Kessil light (Table 1A and available redox potentials and triplet state energies of photocatalysts shown in Table 1B). We discovered that both $Ir(ppy)_3$ and $[Ir(dF(CF_3))_3]$ ppy)₂(dtbbpy)]PF₆ catalyze the transformation despite markedly different redox properties (entries 1 and 2), suggesting that the order of redox events in Scheme 1B are potentially interchangeable.16 The Adachi-type photocatalyst 3DPAFIPN improved the yield to 77% with a further increase to 82% after increasing the reaction concentration (entries 3 and 4). Adding 2,6-lutidine, previously shown as an effective additive for photoredox cross-coupling reactions of BF₃K salts by the Molander group,9 had no impact on the yield of 2-phenylpyridine salt 1a (entry 5) and the [Ir(ppy)₂(dtbbpy)]PF₆ catalyst was marginally

Table 1 Optimization of pyridine alkylation, photocatalyst data and effect of BF₃·OEt₂ as an additive^a

A – Reactio	n Optimization			B – Photocatalyst Redox Properties, and Triplet State Energies ^h			
Photocatalyst, Additive OTf R Me Photocatalyst, Additive 1,4-dioxane, 455 nm Kessil light rt, 16 h R = 2-Ph, 1a R = 3-Ph, 1b 2a		Me R R = 2-Ph, 3a R = 3-Ph, 3b		X=CH, N, (Ir(dF(CF ₃)) X=N, R ₁ =C		Ir(ppy)] ₃ R ₁ =R ₂ =R ₃ =H ppy) ₂ (dtbbpy)]PF ₆ ^l CF ₃ , R ₂ =F, R ₃ =IBu l ₂ (dtbbpy)]PF ₆ ^l	
Entry Salt	Photocatalyst (2 mol%)	Additive (3 equiv) Concen	stration % 3ab	X=N, R ₁ =R ₂ =H, R ₃ =tBu			
1 1a 2 1a 3 1a 4 1a 5 1a	Ir(ppy) ₃ [Ir(dF(CF ₃)ppy) ₂ (dtbbpy)]PF ₆ 3DPAFIPN 3DPAFIPN 3DPAFIPN [Ir(ppy) ₂ (dtbbpy)]PF ₆	none 0.1 none 0.1 none 0.1 none 0.3 2,6-lutidine 0.3 2,6-lutidine 0.3	M 66 M 77 M 82 M 82 (74)° (74)°	Ph N Ph	R ₃	Me Me Me BF4	N. N. N. N. PF6
7 1b	3DPAFIPN	none 0.3	M 20 ^e	3DPAFIPN		[Mes-Acr]BF ₄	[Ru(bpy) ₃]PF ₆
8 1b	3DPAFIPN	2,6-lutidine 0.3	M 60 ^f (42) ^c	Photocatalyst	Ir(ppy) ₃	[lr(dF(CF ₃)ppy) ₂ (dtbbpy)]PF ₆	[lr(ppy) ₂ (dtbbpy)]PF ₆
9 1a	none	2,6-lutidine 0.3		E _{1/2} (PC+/PC*)	–1.73 V	-0.89 V	-0.96 V
10 ^g 1a	none	2,6-lutidine 0.3		E _{1/2} (PC*/PC-)	+0.31 V	+1.21 V	+0.66 V
in the property of				E _{1/2} (PC+/PC)	+0.77 V	+1.69 V	+1.21 V
C – BF ₃ Additive Enables a Previously Ineffective Photocatalyst + PPh ₃ BF ₃ BF ₃ BF ₃ OEt ₂ (1 equiv)			locatalyst	E _{1/2} (PC/PC ⁻)	–2.19 V	−1.37 V	–1.57 V
OTf Ru(bpy)		(PF ₆) ₂ , 2,6-lutidine	Me	E _{0,0} (kcal/mol)	58.1	61.8	49.2
1a Ph	1	Kessil light, rt, 16 h ,4-dioxane		Photocatalyst	3DPAFIPN	[Mes-Acr]BF ₄	[Ru(bpy) ₃]PF ₆
+ e-	2a		NPh	E _{1/2} (PC+/PC*)	–1.38 V		-0.81 V
+ BF ₃			3a	E _{1/2} (PC*/PC-)	+1.09 V	+2.18 V	+0.77 V
Ph ₃ P			% 1a % 3a	E _{1/2} (PC+/PC)	+1.30 V		+1.29 V
P) " _ K "	Without BF ₃ •OEt ₂	98% trace	E _{1/2} (PC/PC ⁻)	–1.59 V	-0.49 V	-1.33 V
	BF ₃	With BF ₃ •OEt ₂	51% 47%	E _{0,0} (kcal/mol)	61.8	44.7	49.0

^a Conditions: **1a** (1.0 equiv.), **2a** (2.0 equiv.), photocatalyst (2 mol%), additive (3.0 equiv.), rt. ^b Yields determined by ¹H NMR analysis using 1,3,5-trimethoxybenzene as internal standard. ^c Isolated yield on 0.50 mmol scale. ^d Isolated yield on 2.00 mmol scale. ^e 3 : 1 4- νs. 2-regioisomeric ratio determined from the crude ¹H NMR. ^g Used 365 nm LEDs instead of 455 nm Kessil light for 89 h. ^h All redox potentials reported νs. SCE and all values compiled from previous literature reports. ¹ Counterion omitted in structure for simplicity.

less efficient under the same conditions (entry 6). We observed that 2,6-lutidine did substantially improve the yield when isomeric 3-Ph salt **1b** was employed (entries 7 and 8); without 2,6-lutidine, the crude ¹H NMR indicates significant amounts of decomposition occurred, including 3-phenylpyridine, and the 4-vs. 2-position product ratio was 3:1. This outcome suggests that protiodephosphination and non-selective Minisci-type pathways can occur under these conditions. With 2,6-lutidine,

We conducted a series of further experiments to explore the effect of light and photocatalyst type on the reaction (Table 1, entries 9–11). Irradiating the reaction at 455 nm without photocatalyst resulted in traces of **3a**, but we did observe 66% yield of the product when we used a 365 nm light (entries 8 and 9). No evidence of an electron donor–acceptor (EDA) complex was observable by UV-Vis spectroscopy, and we propose that the reaction starts by an overlap of the tails of the LEDs emission with the absorption of **1a** at 345 nm (see ESI†).¹¹ Furthermore, a photocatalyst with a redox potential window misaligned with

Table 2 Scope of heterocyclic phosphonium salt coupling partners^a

the crude reaction pathway is cleaner, and the 4- vs. 2-position

ratio improved to 8:1. At this point, we have not established

the role of 2,6-lutidine, although it is conceivable that it reacts

with BF₃ produced as the reaction progresses. In 2-substituted

systems, steric hindrance around the pyridine N-atom of the

salt would deter BF3-coordination, whereas, in 3-substituted

systems, such as salt 1b, coordination is more likely and may

^a Isolated yields of single regioisomers. Conditions: 1 (1.0 equiv.), 2a (2.0 equiv.), 3DPAFIPN (2 mol%), 2,6-lutidine (3.0 equiv.), 1,4-dioxane (0.3 M), rt. ^b 11: 1 crude regioisomeric ratio. Isolated as a single regioisomer. Grey circle denotes the site of alkylation for the minor regioisomer. ^c With 1 equiv. TfOH.

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the redox events in Scheme 1B, [Mes-Acr]BF₄, is also competent (entry 11). An energy transfer mechanism was considered based on entry 9, but the low triplet state energies for [Mes-Acr]BF₄ make this pathway unlikely (Table 1B). However, this result suggests that other factors could influence the reaction mechanism. The results in Table 1C show that pyridine N-activation can play a significant role in some instances. Using Ru(bpy)₃ and 455 nm irradiation resulted in trace products but adding 1 equivalent of BF3·OEt2 formed 3a in 47% yield. Here, we hypothesize that 1a reacts with BF3 to enable single-electron reduction, and radical zwitterions of type IV are intermediates in the reaction. At this stage of our investigations, we conclude that radical zwitterions I and IV are likely present to varying extents depending on the photocatalyst and reaction conditions employed. Using optimal catalyst 3DPAFIPN and adding 2,6lutidine to the reaction may favor I over IV and potentially indicate that I is more reactive in the C-C bond-forming step. Stern-Volmer quenching of 3DPAFIPN by pyridylphosphonium salt **1a** occurs $(K_{SV} = 14.9 \text{ M}^{-1} \text{ and } K_q = 3.55 \times 10^9 \text{ L mol}^{-1} \text{ s}^{-1})$, indicating that type I intermediates are accessible under the reaction conditions without BF₃.12-14

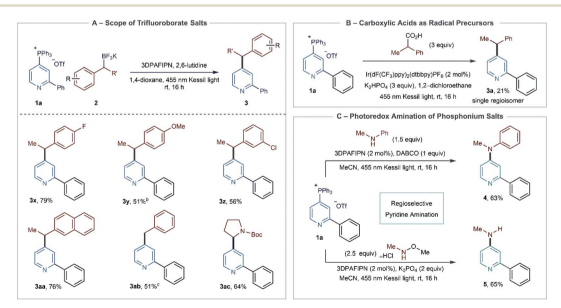
Employing the optimized conditions, we investigated the scope of pyridylphosphonium salts in this coupling process (Table 2). Starting with building block-type pyridines, 2-substituted pyridines with electron-withdrawing and electron-donating groups couple effectively (3c-3e). Aryl and heteroaryl groups are also compatible at the 2-position (3f & 3g), and we did not observe any undesired reactivity of the carbon-iodine bond in 3h. An 11:1 regioisomeric mixture of compounds formed when a 3-ethyl substituent was present (3i), but single isomers formed from 2,3-and 2,5-disubstituted pyridines (3j-3l). At present, 2-substituted pyridylphosphonium salts are not successful coupling partners; as a representative example, when we attempted to synthesize 3m, we observed significant decomposition of the starting material

and only minor amounts (<5%) of the desired compound among other unidentified products.

Next, we converted a series of drug-like fragments and pharmaceuticals into phosphonium salts in this alkylation reaction. These examples represent the most significant advantage of this chemistry as installing a cyano group would be challenging from the C-H bond and limits the ability to make analog compounds. In addition, these structures contain multiple reactive sites and functional groups that could interfere with the coupling process. Nevertheless, we synthesized benzylated fragments 3n-3r without difficulty. Notably, other heterocycles are compatible, such as thiazoles and protected piperidines and pyrrolidines. The pyridine-pyrimidine biaryl 3p is particularly interesting as the phosphonium salt formed siteselectively on the pyrimidine ring, and the photoredox coupling proceeded in good yield on this heterocycle. Lastly, we demonstrated coupling with four FDA-approved pharmaceuticals and an agrochemical that illustrates functional group tolerance for protonated tertiary amines, amides, aryl halides, benzyl ethers, and sulfones (3s-3w). These examples validate this tactic for late-stage functionalization of complex pyridines.

Scheme 2A shows the scope of the BF₃K salts in the photoredox alkylation reaction. Secondary benzylic salts with electron-withdrawing and electron-donating groups are suitable coupling partners (3x–3z). In the case of 3y, we added a 1.2 : 1 mixture of benzylic and homobenzylic BF₃K salts but only observed the benzylated product, presumably because the primary isomer is more difficult to oxidize. Secondary naphthyl and primary benzylic BF₃K salts are proficient, resulting in 3aa and 3ab. The reaction also tolerates α -amino BF₃K salts as evidenced by heterobenzylic amine derivative 3ac. At this stage, non-stabilized radicals were not successful in this process.

Finally, we investigated whether pyridylphosphonium salts are competent with other radical precursors. In Scheme 2B, we



Scheme 2 Scope of radical coupling partners. alsolated yields of single regioisomers. Conditions: 1a (1.0 equiv.), 2 (2.0 equiv.), 3DPAFIPN (2 mol%), 2,6-lutidine (3.0 equiv.), 1,4-dioxane (0.3 M), rt. $^{b}BF_{3}K$ starting material is 1.2 : 1 mixture of regioisomers (benzylic : primary). $^{c}>20$: 1 regioisomeric ratio and 5.7 : 1 mono : bis alkylated product in crude ^{1}H NMR spectrum. Isolated as single monoalkylated regioisomer.

obtained a preliminary result (unoptimized) of coupling with a carboxylic acid. These abundant compounds would improve the scope of radical coupling partners, and further studies are currently underway in our laboratory. In addition, Wu recently reported a method for photoredox catalyzed amination using cyanopyridines as coupling partners, and we attempted to replicate this transformation using pyridylphosphonium salts (Scheme 2C). Applying salt 1a to the reaction protocol with N-methyl aniline resulted in diaryl amine 4. Similarly, using N,O-dimethylhydroxylamine as a coupling partner, followed by *in situ* cleavage of the N–O bond, formed aniline 5 in reasonable yield. Consistent with the results in Table 2, we expect that this reaction will be compatible with more complex pyridine phosphonium salts and further suggests that phosphonium ions can serve as surrogates for cyanopyridines in other radical anion

Conclusions

coupling reactions.

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In conclusion, we report that pyridylphosphonium salts behave as alternatives to cyanopyridines to extend the utility of radical-radical coupling reactions to more complex substrates. We showed that two distinct reactions, pyridine alkylation, and amination, can proceed *via* phosphonium-stabilized radical intermediates. Our lab is currently investigating the capacity of pyridylphosphonium salts to participate in other open-shell reactions, as well as the mechanisms described in this study.

Data availability

All experimental procedures and data related to this study can be found in the ESI.

Author contributions

AMC and JWG conceptualized the work. JWG and BTB performed the experiments in this project. AMC and JWG prepared the manuscript.

Conflicts of interest

There are no conflicts to declare.

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