Selective C(sp³)–H Arylation/Alkylation of Alkanes Enabled by Paired Electrocatalysis

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We report a combination of electrocatalysis and photoredox catalysis to perform selective C(sp³)–H arylation/alkylation of alkanes for the first time, in which a binary catalytic system based on earth-abundant iron and nickel is applied. Reaction selectivity between two-component C(sp³)–H arylation and three-component C(sp³)–H alkylation is tuned by modulating the applied current and light source. Importantly, an ultra-low anodic potential (~0.23 V vs. Ag/AgCl) is applied in this protocol, thus enabling compatibility with a variety of functional groups (> 70 examples). The robustness of the method is further demonstrated on a preparative scale and applied to late-stage diversification of natural products and pharmaceutical derivatives.

Electrosynthesis makes use of electrons as a sustainable and inherently safe redox reagent, representing an environmentally benign synthetic method.¹⁻¹³ In this context, the development of practical and convenient electrosynthesis methods for rapid construction of structurally complex and valuable molecules from readily available simple feedstock chemicals has been of long-standing interest, and has recently drawn renewed attention.¹⁻¹³ Hydrocarbons (e.g., alkanes) are among the most accessible materials available for synthesis; however, owing to their high redox potentials (often above 3.0 V vs SCE) and strong bond-dissociation energy (BDE ~96-101 kcal/mol) of the aliphatic C-H bonds, selective C(sp³)-H functionalization of alkanes remains a compelling synthetic challenge.14-17 Promoting the reactivity of simple hydrocarbons typically requires pre-functionalization via multistep reaction sequences, which result in stoichiometric amounts of waste, low efficiency, and poor atomic economy.

Electrophotochemistry integrates the merits of both electrochemistry and photocatalysis while overcoming their flaws. This method has recently emerged as a powerful platform for performing innovative chemical transformations.¹⁸⁻²⁰ In this technique, an inert substrate (e.g., C(sp³)–H bond) with a high oxidative/reductive potential can be activated at a much lower redox potential. In addition, electrophotochemistry negates requirements for stoichiometric amounts of oxidant/reductant for each turnover of photocatalyst, thus providing more economical and greener alternatives to alkane functionalization.

Since Xu and co-workers reported an elegant photoelectrochemical C-H alkylation of heteroarenes with organotrifluoroborates in 2019,21 electrophotochemistry has rapidly evolved over the past few years. Representative photoelectrochemical catalysts, including [Mes-Acr⁺]ClO₄⁻, trisaminocyclopropenium (TAC) ion, CeCl₃, dicyanoanthracene (DCA), and aryl ketones, has been developed and applied for a variety of transformations.²¹⁻⁵⁰ Despite excellent recent progress, electrophotochemical C(sp3)-H functionalization of inert alkanes remains exceedingly rare.^{29,32,34,37} For example, in 2020, Xu and co-workers described an electrophotochemical Minisci alkylation of N-heteroarenes with aliphatic C-H bonds.³² Later, the Lei group developed a Mn-catalyzed oxidative azidation of C(sp³)-H bonds under electrophotocatalytic conditions.³⁷ Reported examples have typically focused on net-oxidative



Figure 1. Introduction. Selectivity switch between twocomponent $C(sp^3)$ -H arylation and three-component $C(sp^3)$ -H alkylation enabled by paired electrocatalysis.

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functionalization of alkanes. Nevertheless, electrophotochemically driven redox netural $C(sp^3)$ –H arylation/alkylation of alkanes remains unexplored. More than that, selectivity switch between two-component $C(sp^3)$ –H arylation and three-component $C(sp^3)$ –H alkylation of alkanes in one catalytic system still remains elusive and has not been achieved yet (Figure 1).⁵¹⁻⁵⁵

Owing to our on-going interest in electrosynthesis and given our recent success of electrochemical C-H functionalization,56-⁶⁰ herein we report a selective C(sp³)-H arylation/alkylation of alkanes via paired electrocatalysis (Figure 1). Earth-abundant iron and nickel are used as the anodic and cathodic catalysts, respectively. As illustrated in Figure 2. the electrophotochemical strategy involves the conversion of C(sp³)-H to carbon radicals by chlorine radicals, which are generated through ligand-to-metal charge transfer (LMCT) induced by light irradiation of [FeCl₄]^{-,61,62} The relatively large BDE of HCl (102 kcal/mol) ensures that Cl· can activate the strong aliphatic C-H bond (BDE 96-101 kcal/mol) of alkanes via a hydrogen-atom transfer (HAT) process.³² Concurrently, aryl bromides are used as the $C(sp^2)$ coupling partner, which reacts with low valent nickel species generated at the cathode to provide aryl-Ni^{II} species. Finally, aryl-Ni^{II} species intercept the alkyl radical to afford Ni^{III}(aryl)(alkyl) species. Further reductive elimination gives the desired cross-coupled product. Notably, if an alkene is used as a "linchpin" in this reaction, multicomponent C(sp3)-H alkylation reactions may be achieved, thus facilitating rapid buildup of structurally complex and valuable molecules with high atom- and step-economy.



Figure 2. Proposed electrophotochemical $C(sp^3)$ -H arylation/alkylation.

Importantly, owing to the low oxidative potential required to recycle Fe^{2+} to Fe^{3+} ($E_p = 0.25$ V vs. Ag/AgCl, Figure 5b), functional groups that are very sensitive to oxidative conditions would be compatible with this proposed electrophotochemical $C(sp^3)$ –H functionalization of alkanes.

In principle, owing to the large electrode separation in each cell and inefficient interelectrode transport rates, highly reactive intermediates generated simultaneously at the anode and cathode are usually not stable enough to migrate from one side to the other and radical decomposition occurs prior to the desired cross-coupling.⁶³ Hence, the concentration of radical anions/cations generated from paired electrolysis that would further participate their cross-coupling can be quite different, which leads to low reaction selectivity. This mismatch is even more challenging for multicomponent $C(sp^3)$ –H alkylation reactions. Accordingly, achieving control over the rate of reactive alkyl radical generation to match the relatively stable aryl-Ni^{II} speciation is key to ensure a high selectivity of the reaction.

Results and discussion

Investigation of the reaction conditions. On the basis of the above-mentioned challenges, we hypothesized that the LMCT process associated with the anodic half-reactions might be separately modulated by the light source while the cathodic half-reactions could be tuned by dialing in the current or electrode potential, to achieve the proposed rate-matched model of paired electrolysis. Reaction selectivity in electrophotochemistry would benefit from the use of such energy-input tuning. On this basis, our initial investigation started with the reaction of cyclohexane (1a) and 1-bromo-4-(trifluoromethyl)benzene (2a) in the presence of the commercially available FeCl₃•6H₂O and NiBr₂•3H₂O under irradiation by blue light emitting diodes (LEDs) (Table 1). Rewardingly, the desired $C(sp^3)$ -H arylation product 1 was obtained in 48% yield (entry 1). The use of chlorine salts, to increase the effective concentration of the anion [FeCl₄]-, markedly increased the yields and LiCl facilitated the best yield of 91% (entries 2-3). As expected, the light source was essential for the reaction, neither a larger power nor a smaller power, is detrimental to the reaction efficiency (entries 4-5). Further screening showed that the use of graphite felt as an electrode was also important, possibly because of its large specific surface area and the high onset overpotential of the hydrogen evolution

∧ Br		FeCl ₃ 6H ₂ O (10 mol %)						
+ F ₃ C +		+	undivided cell, conditions		\rightarrow F ₃ C	F ₃ C +		
1a	2a		3a (linkage)			1	130	41
entry	electrodes	linkage	light source	current	electrolyte	solvent	yield (%) / 1^{b}	yield (%) / 41^{b}
1	(+)GF/(-)GF	w/o	20 W 395-400 nm	4 mA	Bu ₄ NBF ₄	MeCN	48	-
2	(+)GF/(-)GF	w/o	20 W 395-400 nm	4 mA	Et ₄ NCl	MeCN	57	-
3	(+)GF/(-)GF	w/o	20 W 395-400 nm	4 mA	LiCl	MeCN	91	-
4	(+)GF/(-)GF	w/o	30 W 395-400 nm	4 mA	LiCl	MeCN	79	-
5	(+)GF/(-)GF	w/o	10 W 395-400 nm	4 mA	LiCl	MeCN	17	-
6	(+)GF/(-)Pt	w/o	20 W 395-400 nm	4 mA	LiCl	MeCN	69	-
7	(+)C-rod/(-)GF	w/o	20 W 395-400 nm	4 mA	LiCl	MeCN	27	-
8	(+)GF/(-)GF	w/o	20 W 395-400 nm	10 mA	LiCl	MeCN	56	-
9	(+)GF/(-)GF	w	20 W 395-400 nm	4 mA	LiCl	MeCN	41	22
10 ^c	(+)GF/(-)GF	w	20 W 395-400 nm	25 mA	LiCl	MeCN	11	59
11°	(+)GF/(-)GF	w	20 W 393 nm	25 mA	LiCl	MeCN	7	82
12 ^c	(+)GF/(-)GF	w	20 W 393 nm	25 mA	LiCl	MeCN/Acetone	trace	93
13 ^{c,d}	(+)GF/(-)GF	w	20 W 393 nm	25 mA	LiCl	MeCN/Acetone	n.d.	n.d.
14 ^d	(+)GF/(-)GF	w/o	20 W 395-400 nm	4 mA	LiCl	MeCN	n.d.	n.d.

^aReaction conditions: **1a** (3 mmol), **2a** (0.3 mmol), **3a** (0.6 mmol), FeCl₃•6H₂O (10 mol%), NiBr₂•3H₂O (10 mol%), 2,2'-bipyridine (10 mol%), LiCl (2.0 equiv.), dry MeCN (6.0 mL), 4 mA, 12 h, 20 W 395-400 nm, argon, graphite felt (GF) as electrodes, undivided cell. ^bGC yields using biphenyl as an internal standard. ^c4,4'-di-tert-butyl-2,2'-bipyridine was used as ligand. ^dw/o paired catalyst or light or electricity. n.d. = not detected.

reaction on graphite felt (entries 6–7). Notably, the $C(sp^3)$ –H arylation became inefficient when a higher current was applied (entry 8). Such apparent correlation between the applied current and reactivity indicates the importance of matching cathodic aryl-Ni^{II} speciation with anodic half-reactions.

Inspired by these results, we further examined threecomponent $C(sp^3)$ -H alkylation using methyl acrylate as a linkage. As expected, a mixture of the two-component $C(sp^3)$ -H arylation product 1 and three-component $C(sp^3)$ -H alkylation product 41 was obtained with low selectivity under the optimized conditions for $C(sp^3)$ -H arylation (entry 9). In general, selectivity of the three-component $C(sp^3)$ -H alkylation arises from a competitive alkyl radical Giese addition (E to F step in Figure 2) versus the alkyl radical metalation process (**E** to **G** step). However, these two steps have a small energetic difference based on the product selectivity as has been previously reported.⁶⁴ In contrast, the more electron-deficient alkyl radical **F** has a faster alkyl radical metalation process (**F** to **L** step) than that of **E**. Moreover, considering that the relative concentration of methyl acrylate during catalytic turnover is significantly higher than that of alkyl radical species and the irreversible radical Giese addition, we expect that a higher rate for the C(sp³)–H alkylation process would match a faster aryl-Ni^{II} speciation. Accordingly, the current density plays an important role in controlling selectivity was reversed when

a much higher current (25 mA) was applied (entry 10), affording the corresponding $C(sp^3)$ –H alkylation product **41** in 59% yield along with only 11% of **1**. Modifications of the light source and solvent further increased the yield of the $C(sp^3)$ –H alkylation product **41** to 93% whereas the $C(sp^3)$ –H arylation reaction was retarded (entries 11–12). Control experiments indicated that the paired catalyst, ligand, electric current, and light irradiation were all essential, and their absence led to no enrichment (entries 13–14).

Scope of substrates. Having assessed the feasibility of the paired redox strategy, we next evaluated the reaction scope with respect to C(sp³)-H arylation (Figure 3). Cycloalkanes of various ring sizes ranging from five to twelve carbon atoms are effective coupling partners, gave the desired products 1-4 in good to excellent yields (64%-90% yield). For acyclic alkanes bearing multiple reactive sites, as an example of isopentyl benzoate, only the primary C(sp³)-H-functionalized products were isolated in good yields (5). Pivalonitrile was also tolerated in this reaction, affording the corresponding product 6 in 63% yield. Cyclic ethers were feasible substrates and delivered α oxygen C-H functionalized products in good yields (7). Linear ethers-like 1,2-dimethoxyethane gave exclusively the ethereal coupling product at the primary $C(sp^3)$ -H bonds (8). This protocol was also effective for electron-rich aryl-substituted ethers that are sensitive to oxidative conditions, as exemplified by anisole and 1,3,5-trimethoxybenzene, which gave the desired products in 66% and 75% yields, respectively (9 and 10). These results highlight the extremely mild 'redox-neutral' nature of this electrosynthesis. Ethyl acetate, which possess multiple hydridic C-H bonds, resulted in mixtures of regioisomers (11). Nitrogen-containing hydrocarbons, as exemplified by tert-butyl pyrrolidine-1-carboxylate, were also compatible in this catalytic system, giving the corresponding products (12) in moderate yield. The reaction of benzylic C(sp³)-H bonds was also efficient, delivering the desired products in 71%-81% yields (13 and 14). Notably, in the case of ethylbenzene with multiple hydridic C-H bonds, the reaction proceeded predominantly at the primary C(sp³)-H bonds, with the weaker benzylic C-H bond remaining intact (15). A substrate with allylic C(sp³)-H bonds bearing multiple reactive sites, here exemplified by carvone, gave the coupling product (17) at the primary C(sp³)– H bond. Organosilanes are versatile building blocks in synthesis and extensively used in fields such as materials science and organic synthesis. Various silanes with thiophene, alkenyl, and chloride substituents, were well tolerated under the standard reaction conditions, affording the desired organosilanes in 31%–69% yields (18–20). Electron-withdrawing groups, such as nitriles usually render adjacent hydrogen atoms both stronger and less hydridic, resulting in lower reactivity. Remarkably, this protocol could also be used as a modular synthetic route for functionalization of less hydridic and more challenging C–H bonds. For example, acetonitrile was tolerated in this reaction, affording the corresponding product (21) in 71% yield.

Next, the scope of the reaction with respect to aryl bromide component was explored. As shown in Figure 3, a wide range of aryl bromides, with electron-rich and electron-poor substituents on the aromatic ring, were viable in this transformation. Electron-deficient aryl bromides gave better efficiencies than did electron-rich aryl compounds. Notably, functional groups that are typically sensitive to the electroreductive conditions, such as nitrile (22), ketone (23), ester (24), sulfone, formyl (26) and chlorine (29) were welltolerated,⁶⁵ furnishing the corresponding products in 49%–96% yields. Moreover, functional groups that are typically sensitive to the oxidative conditions, such as methoxyl, methylthio, aryltrimethylsilane and naphthyl also reacted well, giving the desired products in 52%-77% yields. Pharmaceutically relevant aryl halide, as ememplified by a precursor to a beclobrate analogue, was also well tolerated in this reaction, delivering the corresponding product 37 in 51% yields. Further application of this method to heteroaromatic bromides was successful, thereby delivering valuable pyridine, quinoline and benzothiophene derivatives 38-40 in 56% to 93% yields. Remarkably, Miniscitype addition of the alkyl radical to heteroarenes did not compete with C(sp³)-H arylation of N-heteroarene-containing substrates in this protocol, despite the mildly acidic conditions.

Subsequently, we turned our attention to the applicability of three-component $C(sp^3)$ -H alkylation. As shown in Figure 4a, strong aliphatic $C(sp^3)$ -H bonds and α -heteroatom $C(sp^3)$ -H



Figure 3. Unless otherwise specified, all reactions were performed under standard conditions. For details, see the Supporting Information. ^aNiBr₂•3H₂O (20 mol%) and 2,2'-bipyridine (20 mol%) were used as the [Ni] catalyst.

bonds (e.g., 1,4-dioxane) were alkylated efficiently with excellent regioselectivity and 49%–89% yields (41–45). A wide array of aryl bromides, bearing either electron-donating groups or electron-withdrawing groups on the aromatic ring, were viable in this transformation, affording the desired products (46–52) in 58%–83% yields. Heteroaromatic bromides, as exemplified by 4-bromo-2-fluoropyridine, were also amenable to this protocol, and gave the desired product (53) in 79% yields. In addition to methyl acrylate, a series of acrylates with different

substituents and other electron-deficient alkenes, such as acrylonitrile, vinyl amides, were compatible and yielded the corresponding products (54–59) in 53%–85% yields. Furthermore, this electrophotochemical strategy could be readily scaled up with high efficiency. For example, 0.97 g of 1 was isolated in 71% yield (Figure 4b, for details, please see the Supporting Information).

To further extend the synthetic applications of this electrophotochemical bimetallic catalysis protocol, we

investigated late-stage modification of structurally complex substrates, including natural products and pharmaceutical derivatives (Figure 4c). A series of aryl bromides derived from cholesterol, allofuranose, probenecid, fenbufen, menthol, ketoprofen, indomethacin, naproxen, and ibuprofen were coupled to furnish the $C(sp^3)$ -H arylation or $C(sp^3)$ -H



alkylation in moderate to good yields (60–71). These successful outcomes generated a library of structurally diverse molecular architectures and demonstrate the versatility of the protocol.

Mechanistic studies. We next performed a series of experiments to gain insight into the reaction mechanism. First, the electrode voltage over the course of electrolysis for the reaction of **1a** and **2a** was monitored under standard conditions

(Figure 5a). Anodic oxidation was maintained at around 0.23 V vs. Ag/AgCl while the cathodic reduction was kept around -1.1 V vs. Ag/AgCl. This result is in accordance with the oxidative catalytic cycle of the Fe^{II}/Fe^{III} couple (Figure 5b) at the anode and reductive catalytic cycle of the Ni^{III}/Ni^{II} or Ni^{II}/Ni^I (Figure 5c) at the cathode; thus, the direct anodic oxidation of chloride or bromide to give an electrophilic chloride or bromide radical that initiates the reaction can be excluded.



Figure 5. Mechanistic studies. a) Electrode voltage over the course of electrolysis for the reaction of **1a** and **2a**, b) CV of FeCl₃•6H₂O, c) CV of the mixture of NiBr₂•3H₂O and bpy.

In addition, the reaction was completely inhibited if a radical scavenger TEMPO was added to the standard reaction, and a cyclohexylated TEMPO was observed by NMR analysis (Figure 6, for details, see the supporting information). Hence, a carbon radical is likely formed via an anodic catalysis process. Furthermore, the use of a Ni⁰ source, Ni(COD)₂, was found to provide significantly lower efficiency in our system, suggesting that a Ni⁰ species was likely not involved in the main catalytic cycle (eqs b, Figure 6). Subsequently, 4-CF₃C₆H₄Ni(II), complex 73, was synthesized to explore its catalytic efficiency(Figure 6c-d). The desired $C(sp^3)$ -H arylation product was obtained in 37% yield in the stoichiometric reaction of 73 with cyclohexane and one equivalent FeCl₃•6H₂O through light irradiation. Of note, a Ni(I) complex was observed by HRMS analysis in this process, confirming the formation of Ni¹ in the reaction (eqs c, Figure 6). Furthermore, a 46% yield of 23 was isolated in the model reaction with the use of 73 as the nickel catalyst precursor (eqs d, Figure 6). These results reveal that the Ar-Ni^{II} complex might be a catalytic species in this protocol. The proposed mechanism in Figure 2 is thus further confirmed.

Generally, rate-matching modulation plays a crucial role in the selective coupling of reactive intermediates generated from



Figure 6. Mechanistic studies.

paired electrodes. In the three-component $C(sp^3)$ –H alkylationreaction, the high reaction selectivity arises from the effective matching of anodic reactions with cathodic aryl-Ni^{II} speciation. Separate modulation of the LMCT process associated with the anodic half-reactions plays an important role, which is further supported by comparing the reaction efficiency with varying light intensity. For example, $C(sp^3)$ –H alkylation reaction exhibits a higher reaction rate and selectivity (Figure 7) at a high light intensity than that of a low light intensity. These results highlight the unique features of electrophotochemical

systems, which would facilitate reactivities that would not otherwise be possible with a single catalytic system alone.



Figure 7. Rate-matching modulation tuned by the light.

Conclusions

Here, we successfully develop a selective $C(sp^3)$ -H arylation/alkylation of alkanes. Earth-abundant iron and nickel are used in the paired catalyst system that drives alkane functionalization at a low oxidation potential. The reaction selectivity between $C(sp^3)$ -H arylation and alkylation was tuned by separate modulation of the anodic and cathodic reactions in the undivided cell. We report application of the protocol to more than 70 compounds, including late-stage functionalization of natural products and pharmaceutical derivatives, at room temperature. Thereby we demonstrate the broad utility and functional-group tolerance of this protocol.

Data availability

The authors declare that the data supporting the findings of this study are available within the article and its Supplementary Information file.

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Author contributions

Q.L. conceived and directed the project. L.Z. conducted most of the experimental studies. S.X and R.S supported performance of synthetic experiments. Q.L. wrote the manuscript. All authors discussed the results, analyzed the data, and prepared the manuscript.

Competing interests

The authors declare no competing interests