Electrochemical Access to Aza-Polycyclic Aromatic Hydrocarbons: Rhoda-Electrocatalyzed Domino Alkyne Annulations

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Abstract: Nitrogen-doped polycyclic aromatic hydrocarbons (aza-PAHs) have found broad applications in material sciences. Herein, a modular electrochemical synthesis of aza-PAHs was developed via a rhodium-catalyzed cascade C–H activation and azide annulation. A multifunctional O-methylamidoxime enabled the high chemo- and regioselectivity. The isolation of two key rhodacyclic intermediates made it possible to delineate the exact order of three C–H activation steps. In addition, the metalla-electrocatalyzed multiple C–H transformation is characterized by unique functional group tolerance, including highly reactive iodo and azido groups.

Polycyclic aromatic hydrocarbons (PAHs) are broadly explored in material sciences and synthetic chemistry. The incorporation of heteroatoms, such as nitrogen, into these conjugated π-systems can tune or fundamentally alter their optoelectronic and catalytic properties. Therefore, the assembly of atomically precise aza-PAHs in an efficient and economic manner has received considerable attention. However, the synthesis of PAHs and aza-PAHs generally relies on stepwise elaborations, largely involving Diels–Alder cycloadDITIONS, dehydrogenative cyclizations, and transition-metal-catalyzed cross-couplings that require prefunctionalized substrates. Transition-metal-catalyzed oxidative C–H activation/annulation has been proven to be a powerful tool for PAH syntheses. However, the inherent sustainability of the C–H activation approach is compromised by the use of toxic and waste-generating stoichiometric oxidants, such as copper(II) and silver(I) salts.

The merger of metal catalysis and electrosynthesis, the incorporation of metal catalysis and electrochemical synthesis, has been identified as a powerful strategy towards sustainable synthesis, which was substantiated by electrochemical metal-catalyzed C–H functionalizations in recent years. Very recently, the groups led by Ackermann and Xu disclosed a new catalytic regime, namely anodic oxidation-induced reductive elimination, where the electricity not only functions as the terminal oxidant, but is also responsible for anodic oxidation-induced reductive elimination in rhodium-catalyzed C–N and C–P bond formations. These findings may lead to new catalytic activities and transformations, which are not possible with common chemical oxidants.

While most of the aza-PAHs synthesized by transition-metal-catalyzed C–H functionalization involve only one or two C–H activation steps, domino transformations in which three or more C–H bonds are activated and larger π-extended systems constructed continue to be rare. To this end, we envisioned a multifunctional O-methylamidoxime motif to facilitate threefold C–H activation by taking advantage of the imido group after the first internal oxidative C–H functionalization. With our ongoing interest in material syntheses by metalla-electrocatalysis, we have now developed a one-step electrochemical assembly of aza-PAHs via rhoda-electrocatalyzed cascade C–H annulations (Figure 1). Salient features of this electrocatalytic transformation include: a) electricity as a green oxidant, b) excellent functional group tolerance, c) three C–H bonds activated and six new bonds formed, and d) user-friendly scale-up.

We initiated our studies by using amidoxime 1a and diphenylacetylene (2a) for the envisioned rhoda-electrocatalyzed cascade C–H annulation (Table 1). When KOAc was used as the base and [Cp*RhCl₂]₂ (Cp* = C₅Me₅) as the catalyst precursor and with MeOH as the solvent and a constant current of 4.0 mA, the desired product 3aa was isolated in 38% yield (entry 1). Other solvents, such as trifluoroethanol (TFE), H₂O, and acetonitrile failed to deliver product 3aa (entries 2–5). Other bases showed inferior performance compared with KOAc (entries 5–8). The addition of catalytic amounts of carboxylic acids was found to be beneficial, with l-adamantanecarboxylic acid (AdOH) giving the best results (entries 9–11). The reaction gave a higher...
yield at a shorter reaction time, indicating that a high potential and prolonged electrolysis may lead to undesired side reactions or product decomposition (entry 12). Indeed, lowering the applied current to 2.0 mA increased the yield to 75% with a prolonged reaction time (entry 13). Cationic rhodium catalyst \([\text{Cp*Rh(CH}_3\text{CN)}_3(SbF}_6\text{)}_2\) further improved the efficacy, affording product \(3\text{aa}\) in 90% yield at 35°C (entries 14 and 15). Control experiments verified the necessity of electricity and the rhodium catalyst (entries 16–18). Chemical oxidant Cu(OAc)\(_2\) failed to afford \(3\text{aa}\) in effective yield (25%). \([13]\) In addition, when nickel foam was used as the cathode or platinum as the anode, the reaction proceeded, albeit in inferior yields (67% and 65%, respectively). \([13]\)

After establishing the optimized reaction conditions for this rhoda-electrocatalyzed cascade annulative C–H activation, we probed its versatility with various imidoximes \(1\) (Scheme 1). A broad range of aryl amidoximes \(1\) bearing electron-donating (\(1b, 1c, \text{ and } 1d\)) and electron-withdrawing groups (\(1e–1h\)) proved applicable for this electrocatalysis. For asymmetrically substituted substrates \(1i\) and \(1j\), corresponding products were isolated with high levels of regioselectivity and good yields (\(3i\text{a} \text{ and } 3j\text{a}\)). Notably, the highly sensitive iodo substituent was well tolerated under the electrochemical conditions, which facilitates further postsynthetic modifications (\(3k\text{a}\)).

Subsequently, a variety of alkynes \(2\) were evaluated (Scheme 2). Alkynes \(2\) with electron-donating substituents on the arenne motif delivered the desired products (\(3\text{ab}, 3\text{ac}, \text{and } 3\text{ad}\)). The trimethylsilyl group was well tolerated under the electrolysis conditions, serving as a handle for further transformations, such as Hiyama cross-coupling. \([14]\) The cascade annulative reaction proceeded equally well with meta-

| Entry | Base | Additive | Solv. | I [mA] | t [h] | \(3\text{aa}\) [%] |
|-------|------|----------|-------|-------|------|---------------|
| 1     | KOAc| –        | MeOH  | 4.0   | 10   | 38            |
| 2     | KOAc| –        | TFE   | 4.0   | 10   | 0             |
| 3     | KOAc| –        | H\(_2\)O| 4.0  | 10   | 0             |
| 4     | KOAc| –        | MeCN  | 4.0   | 10   | 0             |
| 5     | KOAc| –        | EtOH  | 4.0   | 10   | 4             |
| 6     | NaOAc| –      | MeOH  | 4.0   | 10   | 25            |
| 7     | NaOPiv| –    | MeOH  | 4.0   | 10   | 29            |
| 8     | CsOAc| –        | MeOH  | 4.0   | 10   | 9             |
| 9     | KOAc| \(\text{PivOH}\) | MeOH  | 4.0   | 10   | 43            |
| 10    | KOAc| AdOH     | MeOH  | 4.0   | 10   | 46            |
| 11    | KOAc| AcOH     | MeOH  | 4.0   | 10   | 43            |
| 12    | KOAc| AdOH     | MeOH  | 4.0   | 6    | 56            |
| 13    | KOAc| AdOH     | MeOH  | 2.0   | 12   | 75            |
| 14    | KOAc| AdOH     | MeOH  | 2.0   | 12   | 90(b)        |
| 15    | KOAc| AdOH     | MeOH  | 2.0   | 12   | 89(g)        |
| 16    | KOAc| AdOH     | MeOH  | 2.0   | 12   | trace(e)     |
| 17    | KOAc| AdOH     | MeOH  | 2.0   | 12   | 0(f)         |
| 18    | KOAc| AdOH     | MeOH  | 2.0   | 12   | 60(f)        |

[a] Undivided cell, graphite felt anode (GF) (10 \(\times\) 15 \(\times\) 6 mm\(^3\)), platinum plate cathode (Pt) (10 \(\times\) 15 \(\times\) 0.25 mm\(^3\)), \(1\) (0.2 mmol), \(2\) (0.7 mmol), \([\text{Cp*RhCl}_2]\_2\) (2.5 mol%), base (2.0 equiv), additive (0.1 equiv), solvent (4.0 mL), 25°C under air, yield of isolated product. [b] \([\text{RhCp*(CH}_3\text{CN)}_3(SbF}_6\text{)}_2\) (5.0 mol%), 35°C. [c] \([\text{RhCp*(CH}_3\text{CN)}_3(SbF}_6\text{)}_2\) (2.5 mol%), 35°C. [d] \([\text{RhCp*(CH}_3\text{CN)}_3(SbF}_6\text{)}_2\) (5.0 mol%), without electricity, 35°C. [e] Without rhodium catalyst, 35°C. [f] \([\text{RhCp*(CH}_3\text{CN)}_3(SbF}_6\text{)}_2\) (2.5 mol%), 35°C, under N\(_2\).

**Scheme 1.** Rhoda-electrocatalyzed C–H activation with amidoxime \(1\). Rh cat. = \([\text{RhCp*(CH}_3\text{CN)}_3(SbF}_6\text{)}_2\)
substituted alkyne 2f, affording the desired product 3af in high yield and selectivity. Remarkably, the challenging unsymmetrical alkyne 2g delivered the corresponding product with only two regioisomers and good selectivity. When alkyne 2h bearing pendant hydroxyl groups was applied, the micelle-like PAH 3ah with hydrophilic hydroxyl groups surrounding the PAH moiety was obtained. Surprisingly, the highly functional azide group was well tolerated in this metalla-electrocatalysis (3ai), setting the stage for expedient postmodification and major potential in functional materials and bioimaging.

The high efficacy of this rhoda-electrocatalyzed cascade C–H activation for the synthesis of aza-PAHs motivated us to delineate its mode of action. To this end, intermediates 4 and 5 were isolated, which arise after the first and second C–H activation step, respectively (Scheme 3a). Both rhodacycles 4 and 5 showed catalytic reactivity for the electrocatalysis (Scheme 3b). This evidenced that the three C–H activation steps took place in the order of 1 → 2 → 3 (Scheme 3c). As N-methoxymide was widely used in rhodium- and ruthenium-catalyzed C–H annihilations and functioned as an internal oxidant,[13] a similar pathway might proceed for our new O-methylamidoxime directing group, as N–O bond cleavage was observed in rhodacycle 5 and the products 3. In addition, when cyclic voltammetry studies of 5 in the presence of alkyne 2b were conducted, a low oxidation peak at 0.23 V vs. ferrocene implied that an oxidatively induced reductive elimination pathway was involved for the second and third C–N bond formations (Figure 2). Based on these results, a detailed mechanism was proposed.[13]

The obtained aza-PAHs 3 could be easily transformed to further valuable functional molecular analogues. Treating aza-PAH 3aa with iodomethane afforded a cationic nitrogen-doped nanographene 6 in 93% yield (Scheme 4), which showed reversible redox behavior with its radical form at a low potential of $E_{1/2} = -1.72$ V vs. ferrocene (Figure 3). This promises applicability as a novel anolyte material in organic redox-flow batteries.[16] Intrigued by the six pendant azide groups on 3aj, we performed an azide–alkyne Huisgen cycloaddition reaction with a terminal alkyne linked to protected $d$-lactone 7 (Scheme 5). Thus, a dendrimer with hydrophobic core and hydrophilic periphery 8 was assembled in high yield, which has potential applications in imaging and drug delivery.[17]
substantiated by its ease of scale up. Hence, the gram-scale synthesis of product 3aa was realized with a reduced amount of the rhodium catalyst under a constant current of 12.0 mA (Scheme 6).

In summary, we have herein reported on a modular assembly of aza-PAHs enabled by rhoda-electrocatalyzed cascade C/H functionalization. A multifunctional and transformable O-methylamidoxime was designed to guarantee the reactivity and selectivity. The isolation of two C/H-activated rhodacycles revealed the order of the cascade C/H activation steps. The versatility of the electrosynthesis was demonstrated by its broad substrate scope and excellent functional group tolerance, including iodo and azido groups. The practicality of this reaction was reflected by its mild conditions, user-friendly setup, and easy scale up. The obtained aza-PAHs and their postransformation derivatives were characterized in terms of their photophysical and electronic properties, which point to potential applications in optoelectronics, biomaterials, and energy storage.

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Scheme 5. Synthesis of dendrimer 8 via sixfold aza-alkyne addition between 3aj and 7. DIPEA = N,N-diisopropylethylamine.

Scheme 6. Gram-scale synthesis of 3aa.

Conflict of interest

The authors declare no conflict of interest.

Keywords: aza-PAHs - C=H activation - domino reactions - metalla-electrocatalysis - rhodium catalysis

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