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Strategies for arene dissociation from transition metal  $\eta^6\text{-arene complexes}$ 

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Transition metal  $\eta^6$ -arene complexes have unique properties that facilitate a variety of arene substitution reactions, rendering  $\pi$ -activation a powerful approach for arene functionalization. For decades, these complexes have been studied in the context of coordination chemistry and synthetic methodology *via* stoichiometric reactivity; one central challenge in expanding the utility of arene functionalization *via* transition-metal- $\pi$ -activation is the dissociation of the arene product that remains bound to the transition metal. In this perspective, we highlight representative strategies and methods for the removal and/or exchange of arenes from such complexes. Recent studies that implement these strategies toward catalytic processes are discussed, along with remaining challenges in this area.

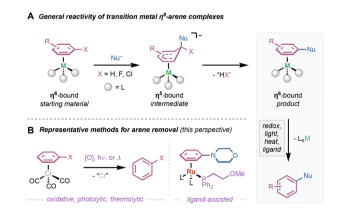
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#### Introduction

Organotransition metal complexes have served as powerful reagents and intermediates in a wide range of synthetic contexts. Complementary to conventional transition metal-catalyzed bond formations in which the metal is directly responsible for making and breaking carbon-carbon (C-C) and carbon-heteroatom (C-X) bonds (i.e. cross-coupling), 1-4 electronic activation of C-C  $\pi$ -bonds *via* coordination of transition metals is a powerful approach for introducing new functional groups (i.e. nucleometalation).<sup>5,6</sup> Comparatively less developed than alkene and alkyne counterparts, transition metal  $\pi$ -complexes with arenes offer a powerful platform for modular functionalization of aromatic compounds. For decades, metalarene  $\pi$ -complexes have been studied under the lens of coordination chemistry to probe the unique properties that  $\eta^6$ binding has on the bound arene.7 Depending on the identity of the metal center, oxidation state of the metal, and ligand environment, n<sup>6</sup>-bound arenes can be activated as substrates toward a variety of nucleophiles and/or electrophiles.8 This "π-activation" strategy has been demonstrated as a powerful method for accessing reactivity that traditional vicarious nucleophilic substitution (VNS), nucleophilic aromatic substitution (S<sub>N</sub>Ar), and electrophilic aromatic substitution (S<sub>E</sub>Ar) cannot. Using this reaction manifold, a wide range of C-C and C-N/O bonds can be installed in both S<sub>N</sub>Ar- and C-H functionalization-type pathways. Despite the attractiveness of this approach, this reaction manifold is largely limited to conditions that employ stoichiometric preformed metal-arene complexes.

When envisioning a catalytic cycle, arene exchange from the transition metal is critical. The first step in  $\pi$ -activation-mediated functionalization, addition of the desired coupling partner to the bound arene, generally leads to an  $\eta^5$ -bound intermediate that rearomatizes to the product  $\eta^6$ -bound arene (Scheme 1A). At this point, the bound arene must dissociate from the metal to yield the desired product. This key step presents a major challenge: selective dissociation of the product arene over starting material. This perspective highlights representative methods and strategies for arene dissociation from transition metal  $\eta^6$ -arene complexes, with an outlook toward further adapting and innovating these protocols in catalytic



Scheme 1 (A) General depiction of transition metal  $\eta^6$ -arene complex undergoing nucleophilic substitution followed by rearomatization and dissociation of the arene product. (B) Overview of highlighted methods for arene dissociation discussed in this perspective.

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contexts. Methods for arene decomplexation are generally organized based on: (1) photolytic or thermolytic cleavage in air, (2) use of chemical oxidants, and (3) ancillary ligand-assisted dissociation (Scheme 1B). Furthermore, we highlight recent efforts toward catalytic  $S_NAr$  that utilize a combination of these strategies. While some functionalization reactions will be discussed in the context of subsequent arene dissociation, this perspective will not include exhaustive details on the types of compatible nucleophiles/electrophiles in this reactivity mode.  $^{9,10}$ 

# Thermolytic and photolytic arene dissociation

In 1970, pioneering work by Helling and coworkers demonstrated that bis( $\eta^6$ -mesitylene)Fe(II) complexes could undergo addition reactions with organolithium reagents at room temperature; in the absence of bound Fe(II), even temperatures as high as 165 °C led only to slow conversion. Subsequent heating of the corresponding ( $\eta^6$ -mesitylene)( $\eta^5$ -arene)Fe(II) at 216 °C under air released the functionalized mesitylene product and mesitylene (Fig. 1). In this case, thermolysis under air served to both rearomatize the product and remove the bound Fe(II). Alternatively, the authors report that a solution of ( $\eta^6$ -mesitylene)( $\eta^5$ -adduct)Fe(II) in pentane could be treated with chemical oxidants to affect the same sequence (*vide infra*).

Shortly after Helling's reports, a cyclopentadienyl (Cp) Fe(II) arene platform emerged as a more stable alternative to dicationic bis(arene)Fe(II) compounds. In 1982, Lee and coworkers showcased a double  $S_NAr$  reaction using a  $CpFe(\pi)(\eta^6-o-1)$ dichlorobenzene) complex with catechol to generate the corresponding  $\eta^6$ -9,10-dioxaanthracene complex (Fig. 2A). Arene removal was achieved via pyrolytic sublimation at 200-250 °C at 1 torr, delivering 9,10-dioxaanthracene in 91% yield. Under these conditions, as shown by Nesmeyanov and coworkers, 13 ferrocene and  $Fe(\Pi)$  salts were generated by disproportionation. In 1997, Roberts and coworkers utilized CpFe(II)(η<sup>6</sup>-fluorobenzene) as a building block for S<sub>N</sub>Ar with chiral amine substituents; they utilized flaked graphite in an 850 W microwave to yield N-phenyl-(s)- $\alpha$ -methylbenzylamine (Fig. 2B). Pearson and colleagues conducted a similar S<sub>N</sub>Ar reaction using CpFe  $(II)(\eta^6-p\text{-dichlorobenzene})$  with phenolates and amines, and achieved decomplexation of the desired product after light irradiation for 4 h in acetonitrile (Fig. 2C).<sup>15</sup>

Using a related  $CpRu(\pi)(\eta^6$ -arene) complex, Pearson later showcased the power of a  $\pi$ -activation strategy in an impressive

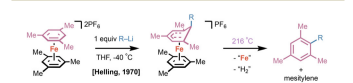


Fig. 1 Helling and coworkers' thermolysis under air to rearomatize and release Fe(n).

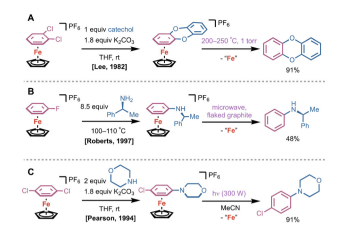


Fig. 2 (A) Lee and coworkers' synthesis of 9,10-dioxaanthracene using pyrolytic sublimation from the corresponding CpFe(II)( $\eta^6$ -arene). (B) Example of arene removal using flaked graphite in a microwave by Roberts. (C) Photolysis of a CpFe(II)( $\eta^6$ -arene) in MeCN solvent to yield desired *N*-aryl morpholine product.

multistep peptide coupling and macrocyclization *en route* to a ristocetin A model substrate. The synthesis relies on a CpRu(II) complex bound to a phenylalanine derivative which undergoes sequential peptide couplings and an intramolecular  $S_NAr$  reaction with a phenol from the D ring of the structure. The combined yield of the  $S_NAr$  reaction and demetallation using light was 46%, illustrating the robustness of this decomplexation protocol (Fig. 3A). In 2022, the same photolytic protocol was used with a related CpRu(II)( $\eta^6$ -arene) complex to dissociate  $S_NAr$  products from 1,3-dione nucleophiles (Fig. 3B).

Photolytic dissociation methods have also been employed for arene dissociation from Cr-based complexes, particularly with respect to  $(CO)_3Cr(\eta^6\text{-arene})$  systems. These typically involve initial photodissociation of a CO ligand, which promotes subsequent arene dissociation. In 1980, Trahanovsky demonstrated *ortho*-lithiation reactions of  $(CO)_3Cr(\eta^6\text{-anisole})$  with *n*-BuLi at  $-40~^{\circ}C$  followed by addition of iodomethane to yield  $(CO)_3Cr(\eta^6\text{-}2,6\text{-dimethylanisole})$ ; the reaction was also

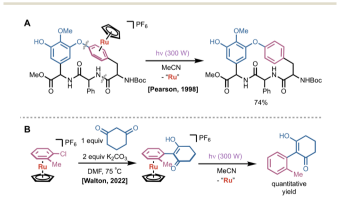


Fig. 3 (A) Photolytic dissociation of a  $CpRu(n)(\eta^6$ -arene) with a macrocyclic peptide after sequential peptide coupling and  $S_NAr$ . (B) Recent example of photolytic dissociation by Walton to generate substituted arenes *via* 1,3-dione addition.

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Et<sub>2</sub>O THE -40 °C [Trahanovsky, 1980] 2.4 equiv *n*-BuLi 2.4 equiv TMEDA then CO2 (dry ice) TMSC

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THF, -78 °C

[Semmelhack, 1979]

(in situ) THF, -30 °C [Rose, 1981]

Fig. 4 (A) Example of an *ortho*-lithiation/methylation sequence using 
$$(CO)_3Cr(\eta^6$$
-anisole) and decomplexation under photoirradiation. (B) Tetralone and benzoate synthesis using  $CO_2$  as an electrophile after *ortho*-lithiation of a  $(CO)_3Cr(\eta^6$ -arene) followed by photolysis to release product. (C) Formal  $[3 + 2]$  to access substituted indanones following photodissociation from product  $(CO)_3Cr(\eta^6$ -arene) complex.

shown to proceed with n<sup>6</sup>-bound fluorobenzene.<sup>17</sup> Exposure to UV light resulted in quantitative decomplexation of 2,6-dimethylanisole (Fig. 4A). Various developments using this ortho-lithiation protocol emerged shortly after; in 1983, Uemura and coworkers used CO2 as an electrophile with (CO)<sub>3</sub>Cr( $\eta^6$ -benzyl alcohol) to synthesize a variety of phthalides and tetralones, albeit as minor products compared to methyl benzoates (Fig. 4B). 18,19 Photoirradiation under air yielded the desired cyclization products in 65-71% yield depending on the substrate. Similar indanone-derived spirocyclic products could be generated via a stereocontrolled formal [3 + 2] annulation from *ortho*-trimethylsilyl benzaldehyde (CO)<sub>3</sub>Cr(η<sup>6</sup>-arene) complexes; in 2002, Moser and colleagues demonstrated that these spirocyclic products could be generated by sequential aldol condensation, Brook rearrangement and cyclization, followed by photoirradiation in Et<sub>2</sub>O (Fig. 4C).<sup>20</sup> In an interesting demethoxylation reaction using LiAlH4 reported by Hacksell in 1991, the resultant (CO)<sub>3</sub>Cr(η<sup>6</sup>-arene) is dissociated using a combination of photoirradiation and ligand-induced displacement with gaseous ammonia.21

## Reagent-driven arene dissociation

Along with photo- and thermolytic methods, subsequent addition of a chemical reagent is an alternative strategy for arene dissociation. These are most commonly in the form of stoichiometric oxidants or Lewis bases that drive dissociative ligand exchange. In 1979, Semmelhack and coworkers conducted a study on substituent effects of (CO)<sub>3</sub>Cr(η<sup>6</sup>-arene) complexes on lithium carbanion addition; the intermediate Meisenheimer-like (CO)<sub>3</sub>Cr(η<sup>5</sup>-bound) species could be simultaneously rearomatized and released from the Cr center by treatment with excess I<sub>2</sub> (Fig. 5A).<sup>22</sup> Notably, one of the first

examples of *ortho*-lithiation of (CO)<sub>3</sub>Cr(η<sup>6</sup>-arene) complexes and subsequent treatment of electrophiles reported by Semmelhack utilized the same strategy.<sup>23</sup> In 1991, Kündig and coworkers reported a similar protocol for decomplexation using I2 after nucleophilic addition of dithianyllithium to a (CO)<sub>3</sub>Cr(n<sup>6</sup>-naphthalene) complex.<sup>24</sup> Interestingly, this report disclosed that difunctionalized products could be accessed in a stepwise fashion starting from (CO)<sub>3</sub>Cr(η<sup>6</sup>-OMe-naphthalene) and treatment with dithianyllithium followed by iodomethane under CO atmosphere. To prevent rearomatization but release product, PPh3 was added to drive decomplexation via ligand exchange. In addition to I2, N-bromo succinimide (NBS) is commonly used as a chemical oxidant to rearomatize  $\eta^5$ -bound species and/or dissociate the desired arene for a variety of transition metal complexes. Trahanovsky demonstrated treatment with NBS at -40 °C as an alternative to photoirradiation on a series of (CO)<sub>3</sub>Cr(η<sup>6</sup>-arene) complexes.<sup>17</sup> In 1981, Rose and coworkers generated either CpFe(II)(η<sup>6</sup>-arene) or the dissociated arene from a CpFe(II)(η<sup>5</sup>-adduct) by controlling the stoichiometry of NBS; one equivalent of NBS carried out rearomatization of the bound arene, while excess NBS directly released product (Fig. 5B).25

Fig. 5 (A) Early report by Semmelhack utilizing I2 to rearomatize and

In cases where a halogen-based oxidant may limit functional group compatibility on the arene fragment, alternative oxidants have been employed. In Helling's 1970 report on the functionalization of [bis(n<sup>6</sup>-mesitylene)]Fe(II) complexes with organolithium nucleophiles (vide supra), it was shown that quenching with dilute aqueous KMnO4 or ceric ammonium nitrate (CAN) yielded desired mesityl derivatives in up to 95% yield.11 In 1989, Miles and coworkers reacted lithium enolate nucleophiles with  $(CO)_3Mn(I)(\eta^6$ -arene) complexes at 0 °C for 1 hour, yielding the  $(CO)_3Mn(I)(\eta^5$ -bound) addition intermediate. Importantly, this was extended to lithium enolates bearing chiral auxiliaries to generate enantiopure  $\alpha$ -arylation products. Upon treatment with 2 equivalents of 2,3-dichloro-5,6-dicyanobenzquinone (DDQ), rearomatization and Mn removal occurred to deliver the α-arylation product (Fig. 6A).<sup>26</sup> Woodgate reported a DDQ rearomatization/dissociation sequence on Fe( $\pi$ )- and Ru( $\pi$ )( $\eta$ <sup>6</sup>-arene) complexes after S<sub>N</sub>Ar with silyl-group-containing nucleophiles.<sup>27</sup> Recently in 2017, Miles and coworkers highlighted a unique siteselectivity of direct lithio-benzofuran addition to a (CO)<sub>3</sub>Mn(ι)(η<sup>6</sup>-

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Fig. 6 (A) Decomplexation via DDQ oxidation to yield enantiopure α-arylation products derived from chiral enolate addition to (CO)<sub>3</sub>Mn(ı)  $(\eta^6$ -arene) complexes. (B) Stemofuran precursor synthesis by Miles and coworkers demonstrating CAN oxidation to rearomatize and dissociate product arene.

2,6-dimethoxytoluene) complex en route to the stemofuran family of natural products (Fig. 6B).<sup>28</sup> Subsequent treatment with CAN or DDQ delivered stemofuran precursors as a single isomer in 93% yield.

## Ligand-based arene dissociation

Given that many reagent-based arene dissociation strategies render the metal center unreactive toward further  $\pi$ -complexation, one attractive strategy to enable catalytic  $\pi$ -activation of arenes is the design of ligands that accelerate arene dissociation and exchange. The resulting ligand systems can be generally categorized in one of two ways: (1) hemilabile ligands that stabilize the transition from the  $\eta^6$ -to  $\eta^4$ -bound arene; or (2)  $\eta^5$ -ligands that enable direct dissociation or accelerate association of other donors.

In 2005 Semmelhack and coworkers reported their efforts in the design of tris(pyrrolyl)phosphine ligands bearing additional chelating groups on one pyrrole substituent.29 Based on the acceleration of arene exchange by coordinating solvents or additives (Fig. 7A), they hypothesized that incorporation of a coordinating side chain would greatly accelerate arene exchange while minimizing the possibility of deactivating the metal center via coordinative saturation.30,31 They found that ligands bearing groups with coordinating lone pairs and low-lying  $\pi^*$  orbitals—such as 2-pyridyl or an ester—greatly accelerated arene exchange in LCr(CO)<sub>2</sub>(η<sup>6</sup>-fluorobenzene) complexes compared to the parent tris(pyrrolyl)phosphine (Fig. 7B). While the unfunctionalized tris(pyrrolyl)phosphine provided slow exchange at 150 °C, the fastest chelating ligands enabled measurable rates at room temperature, and unmeasurably fast exchange at 70 °C.

In 2015, Walton and Williams applied a similar approach to the development of Cp-style ligands to accelerate the ruthenium-catalyzed nucleophilic aromatic substitution of chloroarenes (vide supra).32 They found that tethering donor ligands with an appropriately long carbon spacer, such as carbonyl

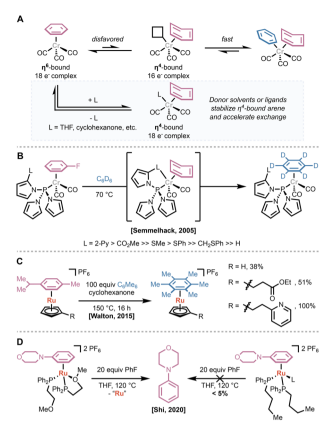


Fig. 7 (A) Rate limiting dissociation of the  $\eta^6$ -arene ligand to form an η<sup>4</sup>-bound species can be accelerated by addition of donor solvents or ligands. (B) Hemilabile tris(pyrrolyl)phosphine ligands accelerate arene exchange from 18-electron chromium complexes. (C) Incorporation of chelating groups to Cp ligands accelerates arene exchange in ruthenium(II) complexes. (D) Demonstration of Shi's hemilabile phosphine ligands that enable exchange of arylamines from ruthenium(II) complexes.

derivatives or a 2-pyridyl group—increased the rate of arene exchange by up to 18-fold compared to the unfunctionalized CpRu( $\pi$ )( $\eta^6$ -arene) catalyst (Fig. 7C). Despite this acceleration, they did not see a concomitant increase in the rate or yield of catalytic reactions. Finally, Shi and coworkers developed hemilabile phosphine ligands to promote arene exchange in LRu(π)<sup>I</sup>(η<sup>6</sup>-arene) complexes (Fig. 7D).<sup>33</sup> They found that incorporating a pendant methoxy group on the phosphine enables dissociation, even when the leaving arene is significantly more tightly binding than the incoming arene.

Other studies have focused on identifying  $\eta^5$ -, Cp-style ligands that promote arene dissociation and exchange. Loginov and coworkers used arene exchange as a model for catalyst initiation in rhodium-catalyzed reductive amination using carbon monoxide as a reductant.34 They found that more donating, permethylated Cp\* and indenyl ligands provided faster exchange than unfunctionalized variants (Fig. 8A). Given the large atomic radius of rhodium, they concluded that the increase in the rate of exchange is due to electronic factors rather than steric effects or indenyl slippage. Computational studies suggested that the more donating ligands stabilize the [LRh(III)]2+ fragment, decreasing the strength of the Rh- $(\eta^6$ -arene) bond.

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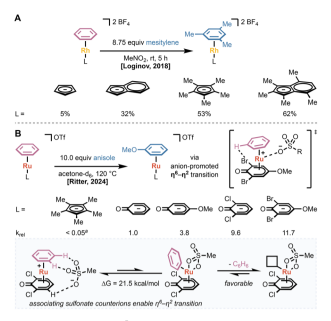


Fig. 8 (A) Electron-rich  $\eta^5$ -ligands provide improved rates of arene exchange. (B) Ritter's  $\eta^5$ -phenoxo ligands provide improved rates of arene exchange due to ligand-arene interactions and anion-promoted  $\eta^6 {-} \eta^2$  transition.

Recently, the Ritter group described the systematic study and design of η<sup>5</sup>-phenoxo ligands to promote arene exchange in LRu(II)( $\eta^6$ -arene) complexes.<sup>35</sup> They found that  $\eta^5$ -phenoxo ligands provide a greater than 20-fold increase in the rate of arene exchange when compared to the isolobal CpRu(π)<sup>I</sup>(η<sup>6</sup>arene) complex (Fig. 8B). Incorporating an electron-donating substituent in the 4-position of the ligand led to an increase in the rate of arene exchange due to a productive interaction between the donor substituent and the counterion. Adding halide ligands to the 2- and 6-positions of the phenoxo also led to an increase in the exchange rate due to a stabilizing hydrogen bond interaction between the halide and the dissociating arene proton in the transition state between  $\eta^6$ - and  $\eta^2$ arene binding. Additionally, they found that more coordinating sulfonate anions increased the rate of arene exchange by interacting with both the leaving arene and the metal center (Fig. 8B).

Other studies have highlighted the ability of specific ligand classes to promote rapid arene exchange or have compared the relative rates of arene dissociation between ligands in a specific class. The lower prevalence of many of these ligands compared to phosphines or cyclopentadienyl ligands and their unproven application to catalytic η<sup>6</sup>-activation make them less applicable to this perspective. 10,36,37

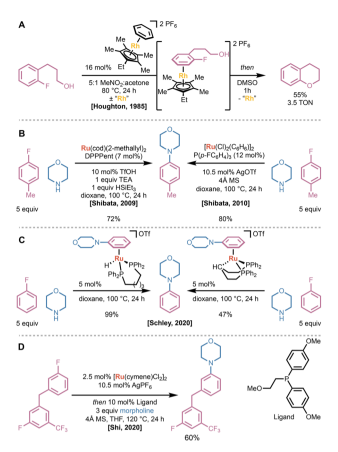
## Catalytic examples of arene dissociation

Despite the expansive study of stoichiometric activation and decomplexation of  $\eta^6$ -bound arenes, catalytic applications remain limited. Incompatibility inherent to solvolysis or chemical methods render the metal center inactive for additional turnover. As a result, many examples of catalytic  $\pi$ -activation of arenes leverage a combination of: (1) accelerated rates of arene exchange on metal centers with large atomic radii; (2) electron-rich Cp\*-style ligands; and (3) inherent thermodynamic favorability due to a decrease in the donicity of the arene following functionalization. One area that experienced exciting developments from a  $\pi$ -activation standpoint is benzylic functionalization; however, we elected to instead focus on a rapidly emerging area that builds on catalyst development strategies.38-43

Beyond α-functionalization of arenes, the other category of reactions in which  $\pi$ -activation has seen significant interest is in promoting S<sub>N</sub>Ar reactions. 44 The strong electron-withdrawing properties of η<sup>6</sup>-bound metal cations enable substitutions that are typically thermodynamically inaccessible or require alternative strategies involving C-X oxidative addition or arene oxidation. While attractive, catalytic  $\eta^6$ -activation is complicated by possible deactivation of the metal center by the nucleophile and a strong thermodynamic preference for binding the product arene over the starting material due to the increase in donicity following substitution of the nucleofuge. 45 The earliest example of catalytic  $\eta^6$ -activation was reported by Houghton, Voyle, and Price in 1984.46 They described the rhodium-catalyzed cyclization of 3-(2-fluorophenyl)propanols to generate chromanes (Fig. 9A). While previous work had found that simple tricarbonyl-metal complexes promoted the desired S<sub>N</sub>Ar reaction under basic conditions, the necessity of strong associating bases prevented catalytic turnover. 47 They found that ethyl-tetramethylcyclopentadienyl rhodium(III) salts were capable of promoting cyclization even under neutral conditions, enabling catalytic turnover. Notably, the identity of the counterion impacted the activity of the catalyst significantly, with the PF<sub>6</sub> salts often providing a 1.2-1.8 times increase in turnover compared to the BF<sub>4</sub> equivalent. Altogether, this early report demonstrated that careful selection of the metal, supporting ligand, and solvent can enable catalytic  $\pi$ -activation strategies, even in the presence of nucleophilic functionalities.

Despite the early success of Houghton and Price, no advances in catalytic π-activation-promoted S<sub>N</sub>Ar were published for 25 years. In 2009, Shibata and coworkers published the first ruthenium-catalyzed amination of unactivated fluoroarenes (Fig. 9B). 48 This system overcame many obstacles: the large thermodynamic preference for binding of the product over the starting material; possible deactivation of the ruthenium(II) center via saturation with the nucleophile; and deactivation due to the production of HF. They found that DPPPent-a wide-bite-angle bisphosphine ligand-in combination with stoichiometric triethylamine and triethylsilane provided the highest yield. To overcome the thermodynamic preference for binding of the product N-arylmorpholine over the starting fluoroarenes, they employed an excess of the fluoroarene starting material. Shibata later published an alternative catalytic system that utilized two equivalents of the monodentate tris(4-fluorophenyl)phosphine rather than DPPPent, and

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**Fig. 9** (A) Houghton's synthesis of chromanes via  $\eta^6$ -activation using Rh(ιιι) catalysts. (B) Shibata's two catalyst systems enable amination of unactivated fluoroarenes using either a silane/base mixture or molecular sieves. (C) The active catalyst in Shibata's DPPPent system is the L<sub>2</sub>Ru(ιι) (H)( $\eta^6$ -arene), which is stable and catalytically active, while the cyclometallated species formed in the absence of a silane is not. (D) An example of how  $\pi$ -activation using Shi's hemilabile ruthenium(ιι) catalyst system provides S<sub>N</sub>Ar orthogonal to traditional thermochemical S<sub>N</sub>Ar reactions.

4 Å molecular sieves to adsorb the formed HF. 49 Notably, both wide-bite-angle bisphosphines and bis-monodentate phosphinyl ligand systems have been shown to accelerate arene dissociation in related rhodium(1) complexes.<sup>50</sup> In-depth mechanistic analysis by Schley and Mueller demonstrated that the silane additive functioned not as an HF or F adsorbent, but to convert the ruthenium catalyst into the  $\eta^6$ -product-bound ruthenium(II) hydride (Fig. 9C). 45 This ruthenium(II) hydride was identified as the resting state of the catalyst, and was stable and soluble under the reaction conditions, unlike the  $\kappa^3$ C-H-activated ruthenocycle. Kinetic and equilibrium measurements established that the rate-limiting step of the reaction was arene exchange, and that the product arene had a binding affinity approximately 2000 times greater than the fluoroarene. Finally, they established that triethylamine was unnecessary, only functioning to generate the triflate anion, which is critical to enable arene exchange. Overall, the work by Shibata revitalized interest in catalytic  $\pi$ -activation and provided a basis for further advancements.

In 2020, Shi and coworkers published the development and application of hemilabile phosphine ligands that accelerate arene exchange in  $L_2Ru(II)(\eta^6$ -arene) complexes (*vide supra*).<sup>33</sup> The use of these ligands enabled amination of un- or deactivated fluoroarenes with excess amine rather than fluoroarene. As a result, Shi's system could be used to maximize the yield of amination products with respect to highly functionalized arenes. Notably, neither bisphosphines nor non-hemilabile phosphine ligands provided significant yield of the desired product (<2% yield). Additionally, the preference for complexation of electron-rich arenes enabled reactivity orthogonal to traditional  $S_NAr$  reactions (Fig. 9D).

Other studies focused on conversion of chloroarenes rather than fluoroarenes, utilizing Cp-style ligands to promote the desired substitution. In 2015, Walton and Williams described the use of CpRu(II)(n<sup>6</sup>-arene) catalysts to afford amination of 4-chlorotoluene.<sup>32</sup> They found that the efficiency of the reaction depended greatly on the solvent employed. While reactions in cyclohexanone provided 16% conversion after 18 h, using 1-octanol improved conversion to 25%. Increasing the reaction time to 14 days led to 90% conversion of 4-chlorotoluene. In the same year, Grushin and coworkers published the fluorination of haloarenes with catalytic Cp\*Ru(η<sup>6</sup>-naphthalene) BF<sub>4</sub>. 51 They found that reactions provided the highest yield when run neat at 180 °C. While turnover numbers were relatively low (3-9 TON in many cases), these results demonstrate that  $\pi$ -activation can be used with diverse arene electrophiles.

Extension of existing ruthenium(II) catalyst systems to engage more diverse nucleophiles was hampered by the low overall electrophilicity imparted by the monocationic ruthenium(II) center in CpRu(II)( $\eta^6$ -arene) complexes.<sup>52</sup> In 2021, Shi and coworkers demonstrated that neither ruthenium(II) catalysts utilizing their hemilabile phosphine ligands nor Cp\* were capable of activating fluorobenzene such that substitution with water occurred under neutral conditions.<sup>53</sup> While using a hydroxide nucleophile did provide the substituted product, the resulting  $\eta^6$ -bound phenol was rapidly deprotonated to form the non-exchangeable η<sup>5</sup>-phenoxo complex, preventing possible catalytic turnover (Fig. 10A). They found that a strategy analogous to that published by Houghton-the use of highly withdrawing rhodium(III) catalysts—allowed for the substitution of bound fluoroarenes under neutral conditions. This strategy allowed for the catalytic hydroxylation or alkoxylation of a variety of fluoroarenes with only three equivalents of water or an alcohol coupling partner (Fig. 10B). Later work by Shi leveraged the strong activating effects of rhodium(III) centers to allow for hexafluoroisopropoxylation of less electrophilic chloro- or bromoarenes using the typically non-nucleophilic hexafluoroisopropanol (HFIP) as a coupling partner.<sup>54</sup> Since binding of the nucleofuge would deactivate the rhodium(III) catalyst, they found that adding stoichiometric basic silver salts, such as Ag<sub>2</sub>CO<sub>3</sub> or Ag<sub>2</sub>O, was necessary to provide the desired product.

In 2024, Ritter demonstrated that utilizing more withdrawing ruthenium(II) catalysts can also provide alkoxylation of fluoroarenes, obviating the need for expensive rhodium-based

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Fig. 10 (A) Many ruthenium(III) catalyst systems are insufficiently activating to promote hydroxylation of fluoroarenes under neutral conditions. Electrophilicity calculations demonstrate that using a more oxidized rhodium(IIII) center is significantly more activating. (B) Examples of hydroxylation and alkoxylation of fluoroarenes using rhodium(IIII) catalysts. (C) Ritter's phenoxo ligands are sufficiently withdrawing such that ruthenium(IIIII) catalysts can promote alkoxylation of fluoroarenes, even with secondary alcohols.

catalysts.<sup>35</sup> They found that  $\eta^5$ -phenoxo ligands were sufficiently activating to enable substitution of bound fluoroarenes even with secondary alcohols under neutral conditions. When used in concert with catalytic amounts of tetrabutyl-ammonium mesylate (TBAOMs)—which accelerated arene exchange compared to triflate—their optimized catalyst provided arylation of epandrosterone, in significantly higher yield than [Cp\*Rh(III)( $\eta^6$ -anisole)]OTf<sub>2</sub> (Fig. 10C). Further, they demonstrated that the same catalyst can promote thermal decarboxylation of phenylacetic acid derivatives at significantly lower temperatures than traditional methods.

Recently, the Shi group has shown that substitution of 2-aminopyridines with amines can occur in the presence of 1,2,4-tri-*tert*-butylcyclopentadienyl ruthenium(II) catalysts. 
<sup>55</sup> While the identity of the ligand is necessary to modulate the efficiency of the substitution, it is unclear if the electronic and steric properties of the  $Cp^{(t-Bu)3}$  ligand promote arene exchange, or if a combination of  $\kappa$ -N binding of the starting material or nucleophile accelerate arene dissociation. Binding studies suggest that, while  $\kappa$ -N coordination of the starting 2-aminopyridine can occur, transition to the  $\eta^6$ -coordination mode occurs rapidly under the reaction conditions.

While deprotonation of phenols to form  $\eta^5$ -phenoxo complexes is degradative in direct  $S_NAr$  reactions, it provides an alternative pathway for functionalization. In 2022, the Shi group showed that, when bound to a rhodium(III) center,  $\eta^5$ -phenoxo compounds can undergo dehydrative condensation with amines to give an  $\eta^5$ -iminium, which tautomerizes to

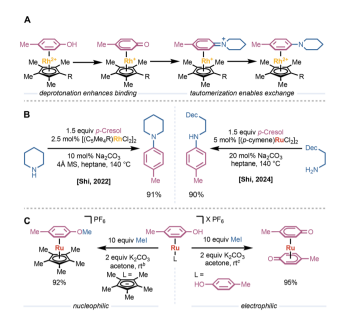


Fig. 11 (A) Outline of the strategy that promotes arene exchange and dehydrative amination of phenols using  $(Me_4RCp)Rh(|||)(\eta^6$ -phenol) complexes. (B) Examples of Shi's dehydrative amination of phenols using both rhodium and ruthenium catalysts. (C) Comparison of the relative nucleophilicity and electrophilicity of  $LRu(|||)(\eta^6$ -p-cresol) complexes.  ${}^3R = 3,5$ -bis(trifluoromethyl)phenyl.  ${}^bX = 1$ .  ${}^cX = 2$ .

form an η<sup>6</sup>-bound aniline (Fig. 11A).<sup>56</sup> The resulting catalytic reaction functions as a pseudo-S<sub>N</sub>Ar of phenols with amines (Fig. 11B). Notably, while exchange of an aniline for a phenol is electronically unfavorable, deprotonation to from the  $\eta^5$ phenoxo provides a thermodynamic driving force for the desired exchange. Later work from Shi and coworkers expanded this type of reaction to employ ruthenium(II) catalysts.<sup>57</sup> Concurrent with the Ritter group, they discovered that the use of η<sup>5</sup>-phenoxo ligands both accelerated arene exchange and improved the electrophilicity of the bound arene (Fig. 11C). They demonstrated the change in the electrophilicity of the bound phenol ligand by reacting both Ru(II)(pcresol)<sub>2</sub> and CpRu(1)(p-cresol) with iodomethane under basic conditions. They found that the Cp\* complex exclusively provided the anisole derivative, indicating that the phenol was nucleophilic. Contrastingly, the bis(p-cresol) complex yielded the neutral  $Ru(II)(\eta^5-p$ -Me-phenoxo) compound. Methylation was only possible upon heating the complex to 80 °C with excess iodomethane. Interestingly, the use of  $\eta^6$ -phenol/ $\eta^5$ phenoxo ligands allowed Shi and coworkers to provide amination products in an effectively ligandless system, where the starting phenol derivative (used in slight excess) functioned as both the ligand and the substrate. This method simplified the reaction setup and prevented any possible conversion of an exogenous phenoxo ligand to the substituted aniline.

Together, the advances in nucleophilic substitution reactions catalyzed by  $\eta^6$ -binding of arenes to ruthenium(II) and rhodium(III) centers highlight many of the approaches used to promote arene exchange. The design of new hemilabile or

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phenoxo ligands, careful selection of solvent, and changes in thermodynamic preference due to favorable tautomerization all provide interesting approaches that enable their specific reaction. Moving forward, it seems likely that further advances will continue to improve the scope, practicality, and applicability of these strategies to existing and new functionalizations.

#### Conclusions

Arene removal and exchange from transition metal  $\eta^6$ -arene complexes serves as a critical enabling feature of modular functionalization via transition metal  $\pi$ -complexes. The examples mentioned here highlight the strategies by which dissociation of the functionalized arene from a transition metal  $\eta^6$ -arene complex is promoted.

Early work done by Helling and Trahanovsky showcased rearomatization and dissociation of the functionalized arene product via thermolysis or photolysis. Alternatively, chemical reagents-most commonly chemical oxidants such as iodine or NBS-may also be used to rearomatize and dissociate the arene product. While these early strategies helped develop the understanding of the conditions necessary to promote arene dissociation and allowed for the use of stoichiometric  $\pi$ -activation in synthetic campaigns, milder strategies that were compatible with diverse metal centers and reagents were necessary to enable applications of  $\pi$ -activation in catalytic contexts.

Guided by ligand electronics and chelating effects, recent advances in the design of catalysts that accelerate arene displacement have revitalized interest in catalytic S<sub>N</sub>Ar enabled by  $\pi$ -activation of arenes. Early work by Houghton demonstrated that leveraging the synergistic effects of metal identity, ligand electronics, counterion effects, and solvent identity could overcome the unfavorable thermodynamics of product inhibition in  $\pi$ -arene S<sub>N</sub>Ar. Following pioneering work by Shibata in 2009, the understanding of each of these effects has grown due to fundamental and catalytic studies by many groups. Most recently, Shi and Ritter demonstrated that new scaffolds such as η<sup>5</sup>-phenoxo ligands can accelerate arene exchange via new factors, such as favorable tautomerization and ligand-arene interactions. Together, the field is poised to largely overcome the issues of product inhibition, slow exchange rates, and insufficient activation that have plagued the development of new  $\pi$ -activation methods.

Contributions over the past decade in S<sub>N</sub>Ar promoted by catalytic  $\pi$ -activation have yielded insight into underexplored strategies of catalytic arene displacement. While these advances have laid the groundwork for future applications of catalytic  $\pi$ -activation, additions to and careful navigation of the existing schema will be necessary to afford efficient and novel catalytic functionalization of arenes *via* transition metal  $\eta^6$ arene complexes. However, further advances are still required to extend catalytic insights summarized herein toward the development of first-row transition metals as competent  $\pi$ -activation catalysts.

## Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

#### Conflicts of interest

There are no conflicts to declare.

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