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Ruthenium-catalyzed free amine directed (5+1) annulation of anilines with olefins: diverse synthesis of phenanthridine derivatives

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A ruthenium(II)-catalyzed cross-ring (5+1) annulation between 2aminobiphenyls and activated olefins is disclosed for succinct synthesis of valuable phenanthridine scaffolds. The protocol avails common organic functional group, free amine, as a directing group represents unique combination of а activation/annulation/C-C bond cleavage cascade that bodes well in the production of bioactive alkaloids including trisphaeridine and bicolorine.

Transition-metal-catalyzed annulation reactions exploiting ubiquitous and otherwise inactive C-H bonds represent an important synthetic strategy to fabricate polycyclic molecular frameworks.^{1,2} Over the years, chemists have compiled a ruthenium-catalyzed reaction compendium that consists of a series of $(4+2)^{3a-e}$, $(3+2)^{3f-h}$, $(2+2+2)^{3i}$, and $(4+1)^{3j}$ annulations, forging diverse carbocycles and heterocycles. Despite these achievements, till now, ruthenium-catalyzed (5+1) annulation has remained largely underdeveloped.⁴ In these annulation reactions, directing groups play fundamental roles to facilitate the C-H bond activation process and mitigate the problem of regioselectivity. Common organic functional groups like carboxylic acid, ester, amide, ketone, etc. are often employed as directing groups. 5 However, the free amine group (NH₂), one of the most valuable and widely abundant functionalities, has largely been ignored in ruthenium-catalyzed directed C-H bond activation reactions,6 probably owing to the challenges associated with its strong coordinating ability with metal catalyst along with the superior nucleophilic reactivity that result in pivotal issues of catalyst deactivation and unwarranted side reactions. 6c,9b Thus, there is an ample scope in free amine directed ruthenium(II)-catalyzed regioselective C-H bond activation/annulation manifold and importantly, it could potentially lead to high-value N-heterocycles when

amine directing group becomes the critical component of the ring structure.

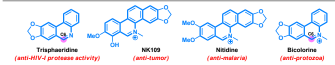
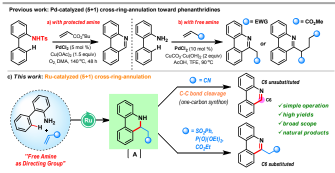


Figure 1. Biologically important phenanthridine alkaloids.

Phenanthridine and benzophenanthridine alkaloids signify an important class of organic molecules with promising biological activities.^{7,8} Some of the important natural products are presented in Figure 1. The biological activities of such alkaloids range from anti-cancer to anti-fungal, anti-bacterial, to name a few. Consequently, devising novel synthetic strategies towards such molecular frameworks is highly desirable.8 Arguably, a C-H bond activation based (5+1) cross-ring-annulation (CRA) reaction of biaryl-2-amines would be a succinct route to access these scaffolds (Scheme 1).



Scheme 1. Ru(II)-catalyzed free amine directed cross-ring (5+1)annulation towards

Further, majorities of the naturally occurring phenanthridine alkaloids do not possess any substitution at the C6-position and hence, challenges lie in strategic designing of a suitable one-carbon synthon for the CRA reaction. In 2012, Li group reported an intriguing Pd-catalyzed (5+1) CRA reaction of biaryl-2-amines with activated alkenes (butyl acrylate) that features the pivotal C-C bond cleavage to offer C6unsubstituted phenanthridines in high yields (Scheme 1a).9a In this case, the use of N-protected biaryl-2-amines was

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COMMUNICATION Journal Name

necessary as N-unprotected biaryl-2-amines gave poor yields. In parallel, Zhang group also reported (5+1) CRA reaction of biaryl-2-amines with alkenes under Pd-catalysis trifluoroethanol (Scheme 1b).9b This reaction is effective with unprotected amine, however, they did not observe any C-C bond cleavage phenomenon and, in case of acrylate coupling partner, a second Michael addition was proposed for the step en route to C6-substituted aromatization phenanthridines. Currently, such CRA reaction manifold for the production of phenanthridines is unknown with Ru-catalysis and herein, we disclose the first example of free amine directed (5+1) CRA reaction of biaryl-2-amines with activated alkenes under Ru-catalysis (Scheme 1c). When acrylonitrile was used as coupling partner, it acts as a C1-synthone and delivered C6-unsubstituted phenanthridines after the C-C bond cleavage. In contrast, other activated olefins, such as vinyl sulfone, vinyl phosphate, and acrylate, furnished C6substituted phenantridines in very high yields.

We commenced our investigations following the model reaction of 2-aminobiphenyl ${\bf 1a}$ with acrylonitrile ${\bf 2a}$ (Table 1). The choice of acrylonitrile as an olefin coupling partner is intriguing as initially formed dihydrophenanthridine intermediate ${\bf A}$ bearing cyanomethyl ($-{\rm CH_2CN}$) functionality

Table 1. Optimization of (5+1) annulation reaction^a

MesCO₂H

MesCO₂H

MesCO₂H

MesCO₂H

entr

1

2 3

4

5

6 7

8

9

10

11^f

12

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	LI CN	acid additive (1.2 equiv) solvent, t °C, 48 h		I H
	1a 2a			3a
γ	acid additive	solvent	temp (°C)	yield (%) ^b
	AcOH	THF	80	52
	AcOH	DCE	80	32
	AcOH	DME	80	38
	AcOH	dioxane	80	46
	$MesCO_2H$	THF	80	72
	$MesCO_2H$	2-Me-THF	80	37
	1-AdCO ₂ H	THF	80	12

100 / 60

80

80

80

80

62/0

<5

59

16c/11d

[Ru(p-cymene)Cl₂]₂ (5 mol %) AgSbF₆ (20 mol %)

Cu(OAc) .: HaO (2 equiv)

13 MesCO $_2$ H THF 80 $16^j/0^k/0^l$ °Reaction conditions: 1a (0.3 mmol), 2a (0.36mmol), solvent (4.2 mL) for 48 h under argon atmosphere. bIsolated yields. 'AgBF $_4$ (20 mol %) was used as additive. 'CuO (2 equiv) was used as oxidant. 'Reaction without $[Ru(p\text{-cymene})Cl_2]_2$ catalyst or Cu(OAc) $_2$ ·H $_2$ O oxidant or AgSbF $_6$ additive. 'Reaction without MesCO $_2$ H (mesitoic acid) additive. '2 equiv of water was added. 'With Pd(OAc) $_2$ '-With (Cp*RhCl $_2$) $_2$ catalyst. 'With (Cp*IrCl $_2$) $_2$ catalyst.

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may experience a C–C bond cleavage phenomenon either through a radical pathway or a coordination assisted base promoted elimination mechanism to validate domino C–H activation based (5+1) annulation *en route* to C6-unsubstituted phenanthridine scaffold (Scheme 1c). Accordingly, when we treated 1a and 2a in presence of [Ru(p-cymene)Cl₂]₂ (5 mol %), Cu(OAc)₂·H₂O (2 equiv), AgSbF₆ (20 mol %), and CH₃CO₂H (1.2 equiv) in THF solvent, we were delighted to find the desired 6-unsubstituted phenanthridine product 3a in 52% yield (Table 1, entry 1). Switching the reaction solvent to DCE, dioxane, and

DME furnished inferior results (entries 2-4). Screening of the acid additives revealed mesitoic acid Pasi: the 3965 CEN 5762, delivering the desired product 3a in 72% isolated yield (entry 5). Change of the reaction solvent from THF to higher boiling 2methyl tetrahydrofuran (2-Me-THF) gave only 37% yield of 3a (entry 6). Further tuning of the reaction conditions, such as use of 1-AdCO2H acid (entry 7), increasing or decreasing of reaction temperature (entry 8), utilization of AgBF₄ additive and use of CuO oxidant (entry 9) had detrimental effects. Control experiments revealed that all the components were essential for the success of the reaction (entries 10-11). Yield also decreased in the presence of excess water in the reaction medium (entry 12). Other transition metals like Pd, Rh, and Ir based catalysts were ineffective under standard reaction conditions, highlighting the uniqueness of ruthenium in this protocol (entry 13).

Having acquired the optimal conditions, we sought to explore the scope of the (5+1) annulation reaction varying the electronic and steric nature in the arene ring (Table 2). The presence of electron-releasing groups such as alkyl (**3b-d**) and alkoxy (**3e-f**) at the *para*-position gave desired products in uniformly high yields (75-84%). Substrates bearing electron-withdrawing groups, for examples halogens (**3g-i**), trifluoromethyl (**3j**), and ester (**3k**) were smoothly reacted to produce C6-unsubstituted phenanthridines in good yields.

 Table 2. Substrate scope of (5+1) annulation with respect to amines

Pleasingly, coordinating free-hydroxyl group did not hamper the reaction, furnishing compound **3I** in 70% yield. When unsymmetrical *meta*-substitution was considered, annulations proceeded selectively at the sterically less hindered site to forge products **3n-p** in good yields. The protocol also worked efficiently with 2-naphthyl derivative, generating important fused polyaromatic heterocycle benzo[j]phenanthridine (**3m**) in 72% yield. The effect of substituents in the aniline ring was also examined; A host of electron-rich and electron-deficient aniline were effective for this reaction, delivering **3q-t** in 62-75% isolated yields. Synthetically useful yield was also obtained with sensitive ketone functionality (**3u**). Under the standard conditions, annulations did not take place with substrates having strongly electron withdrawing cyano (**3v**)

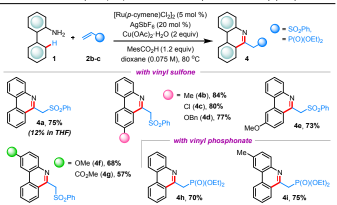
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and nitro (**3w**) groups as well as with anilines derived from heterocycles (**3x-y**).

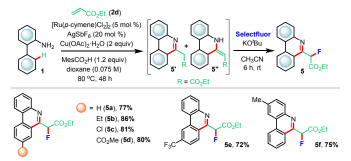
After successful implementation of our hypothesis, we questioned whether other activated olefinic coupling partners would participate in this ruthenium(II)-catalyzed CRA reactions (Table 3).¹⁰ When phenyl vinyl sulfone **2b** was reacted with 2-aminobiphenyl **1a** under the conditions established with acrylonitrile **2a**, the desired (5+1) annulation reaction did not take place effectively with the recovery of starting materials, indicating that a revision of the reaction conditions was necessary. Delightfully, the same reaction proceeded smoothly when the reaction solvent was changed to dioxane; however, we did not observe the concomitant C–C bond cleavage in this case and 6-substituted phenanthridine derivative **4a** was isolated in 75% yield. Other substituted 2-arylanilines also rendered products **4b-g** in good to high yields (57-84%). Similarly, reactions with diethyl vinylphosphonate **2c** were

Table 3. Substrate scope of (5+1) annulation with vinyl sulfone and vinyl phosphonate^a



"Reaction conditions: $\mathbf{1}$ (0.3 mmol), $\mathbf{2b}$ or $\mathbf{2c}$ (0.36 mmol), $Ru(p\text{-cymene})Cl_2l_2$ (5 mol %), $Cu(OAc)_2 \cdot H_2O$ (2 equiv), $MesCO_2H$ (1.2 equiv), $AgSbF_6$ (20 mol %), dioxane (4.2 mL) at 80 °C for 48 h under argon.

Table 4. Substrate scope with acrylate followed by fluorination^a



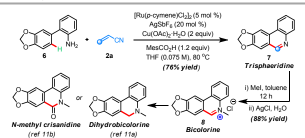
°Reaction conditions:1 (0.3 mmol), 2d (0.36 mmol), Ru(p-cymene)Cl $_2$ l $_2$ (5 mol %), Cu(OAc) $_2$ ·H $_2$ O (2 equiv), MesCO $_2$ H (1.2 equiv), AgSbF $_6$ (20 mol %), dioxane (4.2 mL) at 80 °C for 48 h under argon. Then, KO'Bu (1.2 equiv) and Selectfluor (1.2 equiv) were used in dry acetonitrile at room temperature for 6 h.

fruitful to offer alkyl phosphonate hinged phenanthridines **4h** and **4i** in 70% and 75% yields, respectively (Table 3). These findings reinforce the uniqueness of acrylonitrile in (5+1) crossring-annulation (CRA) for exclusive access of 6-unsubstituted phenanthridines.

Further, reaction of 2-aminobiphenyl **1a** with ethyl acrylate **2d** afforded a mixture of two products which were inseparable

by column chromatography (Table 4). ¹H-NMR aranalysis inferred the presence of desired (5+1) amulated product 5a² along with its tautomer 5a². At this juncture, we posited to use a suitable electrophile to functionalize the acidic C–H bond adjacent to carboxylate group (R = CO₂Et) that might compel the formation of phenanthridine moiety. We focused on electrophilic fluorination since fluorinated analogues of phenanthridine might exert interesting pharmaceutical properties. Consequently, the crude reaction mixture thus obtained from the (5+1) annulation step was exposed to Selectfluor in the presence of KOtBu in anhydrous acetonitrile at room temperature and, to our satisfaction, the desired product 5a was formed in 77% yield (Table 4). Following the same sequence, fluorinated analogues 5b-f were prepared in very high yields (72-86%).

The synthetic utility of this protocol was highlighted in the preparation of phenanthridine-based natural products. For example, trisphaeridine that displays excellent antiproliferative effects on both human and mouse cell was rapidly prepared from the reaction of 2-phenylaniline¹¹ **6** with acrylonitrile **2a** under the standard conditions in 76% yield (Scheme 2). Subsequent methylation gave the natural product bicolorine **8** in 88% yield and synthesis of dihydrobicolorine and *N*-methyl crisanidine from bicolorine is a known process (Scheme 2). ^{12a-b}



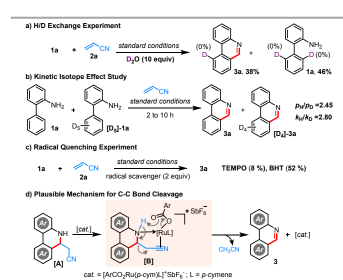
Scheme 2. Synthesis of bioactive alkaloids trisphaeridine and bicolorine

To gain mechanistic insights, we performed few control experiments. No significant deuterium incorporation was observed when bench-mark reaction of 1a and 2a was performed in presence of excess D₂O, approving an irreversible C-H metalation step (Scheme 3a). Kinetic isotope effect (KIE) studies through independent parallel (k_H/k_D = 2.80) and competitive ($p_H/p_D = 2.45$) experiments suggested that the C–H metalation could be the rate-determining step (Scheme 3b). Further, reaction was ineffective in the presence of TEMPO, but 3a was isolated in 52% yield in the presence of BHT, inferring that TEMPO might hamper the Ru-catalysis and the involvement of radical pathway is rather unlikely (Scheme 3c). While the exact reaction mechanism must await further investigations, we believe, in contrast to other alkenes, the unique C-C bond cleavage in case of acrylonitrile is facilitated through the coordination of cationic Ru-catalyst followed by carboxylate assisted deprotonation as shown in Scheme 3d.¹³

In conclusion, an efficient (5+1) cross-ring-annulation (CRA) reaction using readily available 2-aminobiphenyls and activated olefins under common functional group free amine assisted ruthenium(II) catalysis has been accomplished to prepare a library

COMMUNICATION Journal Name

of high-value functionalized phenanthridines in very high yields. Identification of acrylonitrile as a one-carbon synthon was a critical



Scheme 3. Control experiments

parameter for achieving the concomitant C–C bond cleavage, furnishing 6-unsubstituted phenanthridines in a succinct manner. Also, the applications of this methodology in syntheses of bioactive alkaloids like trisphaeridine and bicolorine add to the fruitfulness of the protocol. Further applications of Ru(II)-catalyzed annulation are currently ongoing in our laboratory.

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Conflicts of interest

There are no conflicts to declare.

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Graphical Abstract

(5+1) Cross-Ring-Annulation (CRA) Reaction

