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Edited by Indranil Chatterjee
Imprint

Beilstein Journal of Organic Chemistry
www.bjoc.org
ISSN 1860-5397
Email: journals-support@beilstein-institut.de

The Beilstein Journal of Organic Chemistry is published by the Beilstein-Institut zur Förderung der Chemischen Wissenschaften.

Beilstein-Institut zur Förderung der Chemischen Wissenschaften
Trakehner Straße 7-9
60487 Frankfurt am Main
Germany
www.beilstein-institut.de

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C–H bond functionalization: recent discoveries and future directions

Indranil Chatterjee

The process of C–H bond functionalization can be defined as the replacement of an activated or nonactivated C−H bond with a functional group. This discipline surfaced within the last few decades and proved to be a powerful synthetic tactic due to some remarkable advantages. It has drawn immense attention from the scientific community, thanks to some significant opportunities, such as the use of readily available feedstocks, the introduction of functionality at specific positions of molecules without requiring any prefunctionalized precursors, and the conversion of light alkanes to higher-value analogues [1,2]. The nonnecessity of prefunctionalization provides a step-economic alternative to classical reactions as well as famous Noble-prize-winning cross-couplings, therefore approaching another step up towards sustainability.

Likewise, a free-radical process is also a classical way to functionalize nonactivated C−H bonds in which site selectivity arises either from the relative strength of the C−H bonds or from the abstraction of intramolecular hydrogen atoms. Radical chemistry is a viable alternative to the two-electron process, involving C–H bond functionalization in the absence of any ligand and using low-cost redox-active metals (Fe, Cu, Mn, etc.) rather than heavy metals (Rh, Ir, etc.). Although radical strategies are age-old processes, they were initially cumbersome due to the stoichiometric use of heavy metal salts, peroxides, and other toxic materials as well as the generation of heavy organic and inorganic wastes. In modern days, new strategies are being developed, dealing with photoredox chemistry and its combination with organometallic chemistry for site-selective C−H bond functionalization [3,4]. Recent years have witnessed many viable strategies for the synthesis of complex targets utilizing photoredox catalysis, electroorganic catalysis, Lewis acid catalysis, and transition-metal-free techniques. Some energy-economic reactors such as ball mill, microwave, ultrasound and, most importantly, flow reactors have also evolved towards a more sustainable future.

To showcase the modern approaches in this domain, this thematic issue in the Beilstein Journal of Organic Chemistry gathers recent reports from several research groups, including
photochemical as well as transition-metal-mediated C–H bond functionalization. This mixing of traditional and classical with modern-day research will surely encourage synthetic chemists to sketch new methodologies.

Indranil Chatterjee

Rupnagar, September 2023

ORCID® iDs
Indranil Chatterjee - https://orcid.org/0000-0001-8957-5182

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https://doi.org/10.3762/bjoc.19.114
NaI/PPh$_3$-catalyzed visible-light-mediated decarboxylative radical cascade cyclization of $N$-arylacrylamides for the efficient synthesis of quaternary oxindoles

Dan Liu, Yue Zhao and Frederic W. Patureau$^*$

Abstract
A practical NaI/PPh$_3$-catalyzed decarboxylative radical cascade cyclization of $N$-arylacrylamides with redox-active esters is described, which is mediated by visible light irradiation. A wide range of substrates bearing different substituents and derived from ubiquitous carboxylic acids, including $\alpha$-amino acids, were synthesized and examined under this very mild, efficient, and cost-effective transition-metal-free synthetic method. These afforded various functionalized oxindoles featuring a C3 quaternary stereogenic center. Mechanistic experiments suggest a radical mechanism.

Introduction
Radical-initiated cascade reactions constitute a powerful synthetic approach to construct multiple C–C or C–X bonds in one pot. As such, these tend to allow facile access to many complex natural molecules and drugs [1-6]. Recently, radical-initiated cascade cyclizations involving acrylamides have attracted considerable attention due to their propensity to build important oxindole scaffolds. These are broadly found in natural products, pharmaceuticals, and bioactive molecules (Figure 1) [7-13]. Although a number of synthetic approaches have already been explored [14-20], these existing methods generally require stoichiometric, often onerous reagents [21-28], and/or high temperatures [29-38].

In the past few years, photocatalytic processes have become one of the most powerful tools in developing radical-initiated addition/cyclization cascades from acrylamides for the synthesis of oxindoles [39-41]. The radicals are typically generated from alkyl halides [42-44], carboxylic acids [45-47], simple alkanes [48], alkylboronic acids [49], isocyanides [50], or other [51-53]. In this context, the group of Fu reported a Ru(bpy)$_3$Cl$_2$-cata-
lyzed synthesis of N-Boc proline oxindole derivatives under visible-light assistance [47]. Therein, N-hydroxyphthalimide (NPhth) esters were utilized as alkyl radical precursors, which can be readily prepared from highly available carboxylic acids. In 2015, Cheng and co-workers disclosed a visible light-mediated radical tandem cyclization of N-arylacrylamides with N-(acyloxy)phthalimides to access 3,3-dialkylated oxindoles in the presence of [Ru(bpy)₃Cl₂]·6H₂O [46]. However, these seminal methods remain limited by the need of noble-metal-based photocatalysts, excess additives and limited substrate scopes (Scheme 1a).

With the rapid development of sustainable chemistry, developing low-cost and transition-metal-free photocatalytic methods has become a strategic priority. In 2019 [54], the groups of Fu and Shang pioneered the photocatalytic decarboxylative alkylation of silyl enol ethers and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]. Compared to previously reported radical reactions, this novel catalytic system has the key advantage of circumventing the need for external redox additives and N-heteroarenes by using a novel catalytic system based on sodium iodide (NaI) and triphenylphosphine (PPh₃), suggested to function as an electron donor–acceptor (EDA) complex [55-60]...
dichloromethane (DCM) as reaction solvent (Table 1, entries 16 and 17). Although the reaction also proceeded without NaI, only a low yield of 3aa was then obtained (Table 1, entry 18). PPh₃ and irradiation are however both essential for this decarboxylative cascade cyclization process (Table 1, entries 19 and 20).

With the optimized conditions in hand, we then explored the scope of N-arylacrylamides with different substituents. A series of acrylamides showed good compatibility under standard conditions, offering the desired oxindoles in moderate to good yields (Scheme 2). Electron-donating groups at the para-position of the phenyl ring, such as methyl or methoxy groups, decreased slightly the yield, to 68% and 66%, respectively (3ba and 3ca). When these substituents were replaced by common halogens or electron-withdrawing groups, good yields of the corresponding oxindoles (3da–ga) were achieved. A trifluoromethyl-substituted acrylamide afforded the product 3fa in very high 85% yield. In addition, ortho-substitution at the N-aryl moiety was also well tolerated, albeit with slightly decreased yields (3ha–ka, 50–63%).

Interestingly, a cyclic N-arylamide derivative was also well tolerated, furnishing polycyclic structure 3la in 67% yield. In
addition, substrates with different N-substituents, such as ethyl, benzyl, and phenyl, could be converted into the expected products 3ma–oa in good yields. It should be noted that replacing the methyl with a phenyl group at the N-arylacrylamide core significantly affected the reaction efficiency from 72% to 34% yield (3pa). Satisfyingly, substrate 1q could successfully undergo decarboxylative cascade cyclization to afford 3qa with 70% yield, which is used as a key intermediate in the synthesis of (±)-physosvenine and (±)-physostigmine alkyl analogues exhibiting inhibitory activity against acetylcholinesterase and butyrylcholinesterase [30,78-84]. Subsequently, we expanded the scope of this protocol to include a benzamide derived acrylamide 1r. The expected six-membered ring structure 3ra could be successfully isolated with a good yield (66%).

A number of alkyl radical precursors were then synthesized and evaluated in the reaction (Scheme 3). We found that redox-active esters derived from primary, secondary, and tertiary aliphatic carboxylic acids were all compatible with the method. Cyclic substrates bearing cyclobutyl, cyclopentyl, and indenyl groups could deliver the corresponding desired products with good yields (3ab–ad, 63–74%), while an adamantyl-derived substituent proved more challenging (3ae, 40%). The use of other cyclic substituents such as oxygen-containing and nitrogen-containing rings gave good yields of the target oxindoles (3af–ah, 65–76%). In addition, a symmetrically α-substituted redox-active esters furnished the corresponding quaternary oxindole 3ai with 69% yield. Moreover, an asymmetrically α-branched starting material could react with similar efficiency, affording oxindole 3aj as a 1:1.1 mixture of diastereomers. Interestingly, this method also enabled the synthesis of the highly sterically demanding oxindole 3ak in good yield when using a tert-butyl N-hydroxyphthalimide ester as the tert-butyl radical precursor. Importantly, a redox-active ester derived from methionine could be converted effectively to α-aminoalkylation product 3al in overall 70% yield, which thus
Scheme 2: Arylamide substrate scope with isolated yields of products.

provides a mild method for the functionalization and derivation of abundant natural or unnatural amino acids. Some functional groups such as a terminal alkene in 3am, a terminal alkyne in 3an, and an alkyl chloride in 3ao proved compatible, associated with encouraging yields. In order to further demonstrate the utility of our protocol, a complex scaffold derived from lithocholic acid was tested, and was found to smoothly undergo the decarboxylative cyclization towards oxindole 3ap in 63% yield.

In order to gain insight into the reaction mechanism, some control experiments were further performed. When a radical scavenger such as 2,2,6,6-tetramethyl-1-piperidinyloxyl
Scheme 3: Alkyl radical precursor scope with isolated yields of products.

(TEMPO) was added to the catalytic system under standard conditions, the reaction was fully inhibited, and a TEMPO-trapped adduct (4) was detected by HRMS (Scheme 4a). Moreover, the radical-mediated ring-opening product 3am could be obtained with 66% yield in a radical clock experiment when redox-active ester 5 was engaged to react with acrylamide 1a under standard conditions (Scheme 4b). Finally, it should be noted that benzoyl ester substrate 6a did not deliver the corresponding cyclized product 7aa (Scheme 4c). All of these outcomes indicate that a radical species should be involved in this decarboxylative cascade cyclization towards oxindoles under NaI/PPh₃ catalysis. Thus, the mechanism should run in a similar fashion to related well-documented previous reports [54,68-77], through a light-induced, phosphine-assisted, intermolecular electron transfer from sodium iodide to the redox-active ester.

Conclusion

In summary, we developed an effective photocatalytic decarboxylative radical cascade cyclization of N-arylacrylamides with various redox-active esters derived from common and/or important carboxylic acids under mild conditions. Complementary to traditional transition metal photocatalysis and organo-photocatalysis [85], the readily available and inox-
sive NaI/PPh$_3$ can operate as an efficient photoredox catalyst, providing an economical access to construct important oxindole scaffolds containing a quaternary carbon center. This synthetic method features a broad substrate scope, good functional group tolerance and operational simplicity. Mechanistic investigations revealed that this cyclization reaction proceeds via a cascade radical pathway. We expect these results to encourage the further development of NaI/PPh$_3$-catalyzed and related synthetic methods.

**Scheme 4:** Selected mechanistic experiments.

- **a)**
  - 1a + 2a \[\rightarrow\] 3aa, 0% (TEMPO (1.5 equiv)
  - MeCN, rt, 36 h
  - 456 nm blue LEDs
  - HRMS detected

- **b)**
  - 1a + 5 \[\rightarrow\] 3am, 65%
  - MeCN, rt, 36 h
  - 456 nm blue LEDs

- **c)**
  - 1a + 6a \[\rightarrow\] 7aa, not detected
  - MeCN, rt, 36 h
  - 456 nm blue LEDs

**Supporting Information**

Supporting Information File 1
Experimental section and characterization of synthesized compounds.
[https://www.beilstein-journals.org/bjoe/content/supplementary/1860-5397-19-5-S1.pdf](https://www.beilstein-journals.org/bjoe/content/supplementary/1860-5397-19-5-S1.pdf)

**Funding**
ERC project 716136: 2O2ACTIVATION is acknowledged for generous financial support. We are also thankful to the Chinese Scholarship Council (CSC) for financial support to Dan Liu (No. 202106230113), and Yue Zhao (No. 201908320377).

**ORCID® iDs**
Dan Liu - https://orcid.org/0000-0002-8517-1161
Frederic W. Patureau - https://orcid.org/0000-0002-4693-7240

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Practical synthesis of isocoumarins via Rh(III)-catalyzed C–H activation/annulation cascade

Qian-Ci Gao¹, Yi-Fei Li¹, Jun Xuan² and Xiao-Qiang Hu*¹

Abstract

Herein, we report an unprecedented Rh(III)-catalyzed C–H activation/annulation cascade of readily available enaminones with iodonium ylides towards the convenient synthesis of isocoumarins. This coupling system proceeds in useful chemical yields (up to 93%) via a cascade C–H activation, Rh-carbenoid migratory insertion and acid-promoted intramolecular annulation. The success of gram-scale reaction and diverse functionalization of isocoumarins demonstrated the synthetic utility of this protocol.

Introduction

Isocoumarins are an important structural motif in many naturally occurring lactones isolated from bacterial strains, molds, and plants, exhibiting a wide range of pharmacological properties such as antibacterial, antitumor, and anti-HIV activities (Scheme 1a) [1-5]. Fascinated by their versatile properties, researchers were prompted to develop efficient methods for the synthesis of isocoumarin scaffolds. Traditional synthetic strategies including 1) intramolecular cyclization of 2-alkenyl benzoic acids or α-alkynylbenzoates (Scheme 1b, I) [6-10], 2) oxidation of isochromans (Scheme 1b, II) [11,12], or 3) metal-catalyzed cross-coupling/cyclization of 1,2-difunctionalized arenes with alkynes or carbon monoxide (Scheme 1b, III) [13-16], have been widely applied for the assembly of isocoumarins over the past decades. Recently, the transition-metal-catalyzed ortho C–H activation/annulation of benzoic acids has emerged as an attractive approach towards
Scheme 1: Significance of isocoumarins (a), classic methods for the synthesis of isocoumarins (b) and reaction design (c).

isocoumarins [17,18]. Pioneering examples relying on the Pd, Ru, and Ir-catalyzed C–H cross coupling of benzoic acids with alkenes and alkynes were realized by the groups of Miura [19], Lee [20], Ackermann [21], Zhang [22], Jiang [23], and Jegamohan [24] et al. Despite these impressive achievements, established methods often require the use of stoichiometric oxidants or harsh conditions, thus limiting their broad applicability. Consequently, it is still highly desirable to exploit innovative and convenient methods for the rapid construction of isocoumarins.

Enaminones are bench stable and easily available, which have been established as versatile synthetic building blocks for the synthesis of cyclic scaffolds [25]. In 2016, Zhu et al. reported the first example of a Rh-catalyzed C–H functionalization of enaminones with alkynes and α-diazo-β-ketoesters to access naphthalenes [26]. Very recently, the same group developed an efficient Rh(III)-catalyzed C–H cross-coupling of enaminones with diazodicarbonyls for the divergent construction of isocoumarins and naphthalenes [27]. Moreover, Loh et al. disclosed a Rh-catalyzed formal [4 + 2] cycloaddition of enaminones with diazocarbonyls [28]. Compared with highly sensitive diazo compounds, iodonium ylides are known to show ready availability and good stability [29,30]. Our group has recently demonstrated that iodonium ylides can be used as carbene precursors in the Rh-catalyzed [4 + 2] cyclization of pyrazolidinones [31]. During the preparation of manuscript, the group of Li reported a similar Rh(III)-catalyzed [3 + 3] annulation of enaminones with iodonium ylides [32]. Inspired by the collected contributions [26-28] and based on our ongoing research in C–H activation [33-35], we recently wondered whether it might be possible to couple iodonium ylides with enaminones in a Rh(III)-catalyzed C–H activation/annulation cascade reaction for the rapid construction of isocoumarins (Scheme 1c).

Results and Discussion
Our initial experiment was performed with enaminone 1a and iodonium ylide 1b in the presence of [Cp*RhCl₂]₂ (5 mol %) as the catalyst, AgSbF₆ (10 mol %) and KOAc (50 mol %) as additives in 1,2-dichloroethane (DCE) at 100 °C for 16 hours. To our delight, the desired isocoumarin 3aa was obtained in 42% yield (Table 1, entry 1). Then, the influence of solvents has been subsequently investigated. As a result, DCE proved to be the optimal solvent, while other commonly used solvents such as toluene, dioxane, and ethanol gave inferior yields (Table 1,
Table 1: Optimization of the reaction conditions.

<table>
<thead>
<tr>
<th>entry</th>
<th>catalyst</th>
<th>additive</th>
<th>solvent</th>
<th>yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>[Cp*RhCl₂]₂</td>
<td>KOAc</td>
<td>DCE</td>
<td>42</td>
</tr>
<tr>
<td>2</td>
<td>[Cp*RhCl₂]₂</td>
<td>KOAc</td>
<td>toluene</td>
<td>39</td>
</tr>
<tr>
<td>3</td>
<td>[Cp*RhCl₂]₂</td>
<td>KOAc</td>
<td>dioxane</td>
<td>16</td>
</tr>
<tr>
<td>4</td>
<td>[Cp*RhCl₂]₂</td>
<td>KOAc</td>
<td>EtOH</td>
<td>14</td>
</tr>
<tr>
<td>5</td>
<td>[Cp*RhCl₂]₂</td>
<td>Cs(OPiV)₂</td>
<td>DCE</td>
<td>47</td>
</tr>
<tr>
<td>6</td>
<td>[Cp*RhCl₂]₂</td>
<td>Cs(OAc)₂</td>
<td>DCE</td>
<td>51</td>
</tr>
<tr>
<td>7</td>
<td>[Cp*RhCl₂]₂</td>
<td>K₂CO₃</td>
<td>DCE</td>
<td>27</td>
</tr>
<tr>
<td>8</td>
<td>[Cp*RhCl₂]₂</td>
<td>AcOH</td>
<td>DCE</td>
<td>63</td>
</tr>
<tr>
<td>9°</td>
<td>[Cp*RhCl₂]₂</td>
<td>AcOH</td>
<td>DCE</td>
<td>80</td>
</tr>
<tr>
<td>10</td>
<td>{Ru(p-cymene)Cl₂}₂</td>
<td>AcOH</td>
<td>DCE</td>
<td>trace</td>
</tr>
</tbody>
</table>

aStandard conditions: 1a (0.2 mmol), 2a (0.6 mmol), catalyst (5 mol %), AgSbF₆ (10 mol %), additive (0.5 equiv), solvent (2.0 mL) at 100 °C for 16 h. bIsolated yields. cAcOH (5.0 equiv) was used.

tables 2–4, 14–39%). Further screening of bases did not improve the outcome of the product, whereas 63% yield of 3aa was obtained when acetic acid was added into the reaction system (Table 1, entry 8). Increasing the amount of acetic acid significantly improved the reaction efficiency delivering product 3aa in 80% yield (Table 1, entry 9). The choice of a suitable catalyst was the key factor for the success of this reaction since only a trace amount of 3aa can be obtained by using [{Ru(p-cymene)Cl₂}₂] as a catalyst (Table 1, entry 10).

With the optimal conditions in hand, we then investigated the scope of this Rh-catalyzed C–H activation/annulation cascade reaction. As shown in Scheme 2, a range of functionalized enamiones were compatible with this Rh-catalytic system, furnishing the corresponding isocoumarin products in satisfying yields. For example, enamiones bearing electron-donating (Me, OEt and t-Bu), as well as electron-withdrawing groups (F, Cl, Br, I, CF₃, CN and NO₂) at ortho, meta or para-positions were well tolerated in this transformation, delivering a variety of structurally diverse isocoumarins in an efficient manner (3aa–qa, 43–82%). It is worth mentioning that the tolerance of halogen substituents (Cl, Br and I) may open up a new opportunity for further transition-metal-catalyzed cross-coupling reactions. Sensitive groups, such as ester, trifluoromethyl and nitro substituents, were retained unchanged in the final products (3ja, 3ka, 3ma, and 3na). Also the strongly coordinating thioether substituent proved to be suitable for this protocol, providing the desired product 3ea in 76% yield. Moreover, under the standard conditions, 3-thienyl and 2-naphthyl-substituted enamiones were smoothly coupled with iodonium ylide 1b to give the expected isocoumarins 3ra and 3sa in 60% and 78%, respectively.

Next, we sought to test the generality of this reaction with respect to iodonium ylides. As outlined in Scheme 3, iodonium ylides featuring dimethyl, methyl, and phenyl groups underwent the current reaction efficiently, delivering the desired products 3ab–ae in moderate to good yields (43–93%).

To demonstrate the synthetic utility of this methodology, a gram-scale synthesis of isocoumarin 3ia was firstly performed. Under the optimal conditions, the desired product 3ia was successfully obtained in 84% yield (1.1 g) via a simple recrystallization from the reaction mixture (Scheme 4a). In the presence of hydroxylamine hydrochloride, the carbonyl group of the ketone can be selectively converted into an oxime product 4 (Scheme 4b, 71% yield). In addition, the reaction of the isocoumarin 3ia with p-toluenesulfonyl hydrazide proceeded smoothly to deliver hydrazone 5 in 66% yield (Scheme 4b, right). Of note, oxime and hydrazone compounds are versatile synthetic building blocks, which have been widely applied in transition-metal-catalyzed cross-coupling reactions and radical transformations [36-38].
Scheme 2: Scope of enaminones.

Scheme 3: Scope of iodonium ylides.
Based on the literature precedents [27] and our previous work [33-35], a mechanism for this Rh-catalyzed C–H activation/annulation reaction was proposed and depicted in Scheme 5. In the presence of AgSbF$_6$, dimeric [Cp*RhCl$_2$]$_2$ transforms into the active Rh catalyst. Subsequently, the oxygen atom of the enaminone is coordinated to the Rh catalyst, following by a
Rh(III)-promoted ortho C–H activation to form a five-membered ruthenate cycle 1–A. Then, the reaction of the iodonium ylide with intermediate 1–A generates a Rh-carbenoid intermediate 1–B, which undergoes a rapid migratory insertion to give intermediate 1–C. The protonation of 1–C produces the intermediate 1–D with the regeneration of the active Rh catalyst for the next catalytic cycle. Under acidic conditions, the further protonation of compound 1–D delivers an imine intermediate 1–E, which undergoes an intramolecular annulation to give 1–F. The final isocoumarin product 3ba can be generated from 1–F by elimination of imine 1–G [39]. Finally, the rapid hydrolysis of the resulting 1–G gives rise to acetaldehyde and dimethylamine as byproducts.

**Conclusion**

In summary, an efficient Rh-catalyzed C–H activation/annulation reaction of enaminoles with iodonium ylides has been developed. This reaction features simple operation and readily available substrates, enabling the rapid construction of biologically attractive isocoumarins in useful to good yields. The success of a gram-scale reaction and diverse functionalization of the isocoumarin products highlight the tremendous synthetic potential of this methodology in chemical synthesis and drug discovery.

**Experimental**

A 10 mL screw-cap vial was charged with enaminoles 1 (0.2 mmol), iodonium ylide 2 (0.6 mmol), [Cp*RhCl₂]₂ (6.2 mg, 5 mol %), AgSbF₆ (6.9 mg, 10 mol %), HOAc (6.2 mg, 5 mol %), AgSbF₆ (0.2 mmol), iodonium ylide (0.6 mmol), [Cp*RhCl₂]₂ (0.2 mmol) and DCE (2 mL) under N₂ atmosphere. Then, the reaction mixture was stirred at 100 °C for 16 h. The crude product was purified by flash chromatography on silica gel (petroleum ether/ethyl acetate 5:1) directly to give the desired products 3. (Note: a heating module was used for the preparation of isocoumarin products 3.)

**Supporting Information**

Supporting Information File 1

Experimental and copies of spectra.

[https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-19-10-S1.pdf](https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-19-10-S1.pdf)

**Funding**

We are grateful to the National Natural Science Foundation of China (No. 21901258, 22271314, 21971001 and 21702001), the Foundation of South-Central MinZu University (YZZ19003), Hubei Provincial Natural Science Foundation, China (2021CFA022) for support of this research.

**ORCID® iDs**

Jun Xuan - https://orcid.org/0000-0003-0578-9330

Xiao-Qiang Hu - https://orcid.org/0000-0001-9094-2357

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Transition-metal-catalyzed C–H bond activation as a sustainable strategy for the synthesis of fluorinated molecules: an overview

Louis Monsigny‡, Floriane Doche‡ and Tatiana Besset*
The goal of this review is to highlight and discuss the recent approaches for the synthesis of fluorinated derivatives by the direct incorporation of a fluorinated group of XR_F type (e.g., SCF_3, SeCF_3, SCF_2CO_2Et, OCH_2CF_3) by transition-metal-catalyzed C–H bond activation (Scheme 1). The review will be organized in two main parts, dedicated to the construction of a C–SCF_2R/SeCF_3 and C–OCH_2CF_3 bond. This review does not aim to be exhaustive and key examples were carefully chosen to provide the reader a nice overview. Since reviews dealing with transition-metal-catalyzed functionalization of compounds by C–H bond activation with fluorinated reagents [60-77] and vinylic, allylic or propargylic fluorinated building blocks [49] have been already reported, these reactions will not be included.

**Scheme 1:** Transition-metal-catalyzed C–XR_F bond formation by C–H bond activation: an overview.

**Review**

I. Transition-metal-catalyzed directed C–chalcogen bond formation (C–S, C–Se) by C–H bond activation

In the past decade, particular attention has been paid to the development of new methodologies for the incorporation of sulfur-containing fluorinated groups. Although the transition-metal-catalyzed direct C–H bond functionalization appeared to be a powerful tool for the installation of C–C, C–N, or C–O bonds, the direct formation of a C(sp^2)–SR or a C(sp^3)–SR_F bond remains a challenging task. In this context, key players in the field have been interested in the design of original methodologies for the trifluoromethylthiolation and more recently the difluoromethylthiolation of various compounds by transition-metal catalysis [78]. Moreover, a recent interest was devoted to the trifluoromethylselenolation reaction as depicted in this section.

I.1) Transition-metal-catalyzed C–H trifluoromethylthiolation of aromatic C(sp^2) centers

Thanks to its unique features such as an interesting lipophilicity (Hansch parameter = 1.44) [79,80] and a strong withdrawing character, the development of new methodologies for the incorporation of the SCF_3 residue on various molecules has known a tremendous expansion [81-113].

**Copper catalysis:** In 2012, Daugulis and co-workers reported the copper-promoted trifluoromethylthiolation of benzamide derivatives 1 at the ortho-position by C–H bond activation [114]. Indeed, using a bidentate directing group (amide derived from the 8-aminoquinoline), the mono- and difunctionalized compounds were obtained when Cu(OAc)_2 (0.5 equiv) and the toxic and volatile disulfide F_3CS–SCF_3 were employed (Scheme 2, 10 examples, up to 76% yield). With this approach, derivatives bearing an aromatic part substituted at the para-position with electron-donating substituents (1a,b), halogens (1c) as well as

**Scheme 2:** Cu(OAc)_2-promoted mono- and difluoromethylthiolation of benzamide derivatives derived from 8-aminoquinoline reported by the group of Daugulis [114].
electron-withdrawing groups (1d) were difunctionalized in good yields. The substitution pattern on the aromatic ring did not affect the reaction efficiency, the meta-substituted derivative 2e as well as the ortho-substituted derivative 2f were obtained in high yields (70% and 63% yields, respectively). It should be noted that the presence of ortho-substituents on the aryl residue allowed the monofunctionalization to occur selectively. Also, amide 1g bearing a disubstituted arene was successfully functionalized in 59% yield. Finally, the difunctionalized thiophene derivative 2h was obtained in 56% yield.

In 2016, Wang’s group developed another methodology for the trifluoromethylthiolation of azacalix[1]arene[3]pyridines by C–H bond activation using a complex of Cu(ClO$_4$)$_2$·6H$_2$O and the shelf-stable Me$_4$NSCF$_3$ [115,116] as a nucleophilic source of SCF$_3$ (Scheme 3) [100]. Within these conditions, a set of six azacalix[1]arene[3]pyridines bearing electron-donating groups, halogens or electron-withdrawing groups were functionalized and the expected products were isolated in moderate to high yields (4a–f, 58–91%).

**Palladium catalysis:** Several works have been reported for the palladium-catalyzed trifluoromethylthiolation reaction of various aromatic compounds 5 by C–H bond activation and involved in most cases an electrophilic SCF$_3$ source (R$_1$R$_2$NSCF$_3$). For these transformations, the following working hypothesis was generally suggested (Scheme 4). After coordination of the palladium catalyst to a directing group, the metallacycle A is formed. This latter undergoes an oxidative addition in the presence of an electrophilic source or an oxidation/ligand exchange in the presence of a nucleophilic source (i.e., AgSCF$_3$) and an oxidant (B in Scheme 4). Finally, after a reductive elimination step, the expected functionalized product 6 is obtained and the palladium catalyst is regenerated.
In 2014, Shen and Xu [117] developed a new methodology for the selective functionalization of 2-arylpyridine derivatives using an electrophilic SC\textsubscript{F}\textsubscript{3} reagent, the Haas reagent I (Scheme 5) [118]. A broad range of 2-arylpyridine derivatives were trifluoromethylthiolated in good to high yields (18 examples, from 52 to 91% yields). The substitution pattern of the aromatic ring had no impact on the outcome of the reaction as illustrated with substrates substituted by a methyl group (7b, 7d, and 7f) at the para-, meta- and ortho-positions, which were readily functionalized in 71%, 84% and 78% yields, respectively. This reaction was also tolerant of a 2-naphthyl group, the palladium-catalyzed trifluoromethylthiolation afforded the corresponding product 8h in 76% yield. Also, the 2,4-dimethoxylated substrate 7g and the benzothiophene derivative 7i were successfully trifluoromethylthiolated in 76% and 63% yields, respectively. This reaction proved to be compatible with the presence of an ester (8c) or a halogen (8e). Other directing groups, such as substituted pyridines (9a and 9b) and pyrimidine (9e) turned out to be also efficient in this transformation (Scheme 5, 4 examples, up to 84% yield).

The same year, the group of Huang reported an elegant and straightforward palladium(II)-catalyzed ortho-selective trifluoromethylthiolation of arenes bearing various directing groups using the nucleophilic trifluoromethylthiolating source AgSC\textsubscript{F}\textsubscript{3} in combination with Selectfluor\textsuperscript{®} as oxidant (Scheme 6, 29 examples, up to 91% yield) [119]. 2-Arylpyridine derivatives bearing electron-donating groups, electron-
Scheme 6: C(sp^2)–SCF₃ bond formation by Pd-catalyzed C–H bond activation using AgSCF₃ and Selectfluor® as reported by the group of Huang [119]. ²⁰ equiv of C₂HCOOH were used instead of AcOH.
withdrawing groups or halogen at the para- and meta-positions of the aromatic ring were readily functionalized (11a-g, 58–85% yields). Also 2-(2-methoxyphenyl)pyridine (11h) and 2-(2-naphthyl)pyridine (11i) were found to be suitable substrates leading to the corresponding products 12h and 12i in 91% and 83% yields, respectively. The use of other directing groups was also suitable for this transformation such as methyl and cyano-substituted pyridines 13a,b, pyrimidine (13c), pyrazole (13d), as well as the amide derived from 8-aminoquinoline 13e (Scheme 6, 13 examples, up to 75% yield).

In this study, two mechanisms were reported. The first one suggested that a palladacycle C is formed after the irreversible chelation of the 2-phenylpyridine substrate with palladium, which is the rate-determining step (KIE = 2.7). Subsequently palladacycle C is oxidized by Selectfluor® to form a palladium(IV) complex D. After a ligand exchange with AgSCF₃, the intermediate E is obtained, which, after reductive elimination, releases the desired product 12 and regenerates the catalyst. Alternatively, a ligand exchange with AgSCF₃ occurs before the oxidation step, generating the palladium(II) complex F. After an oxidative addition in the presence of Selectfluor®, the palladium(IV) intermediate E is generated. Finally, after reductive elimination step, the desired product 12 is released and the catalyst regenerated. Note that, in this process, Selectfluor® is playing a key role. Indeed, using this electrophilic fluorinating source as oxidant generates a Pd(IV)(ppy)F(OAc)₂ (ppy = 2-phenylpyridine) complex as intermediate. As the competitive C–F bond formation was disfavored (slow reductive elimination step), the desired trifluoromethylthiolated product 12 is selectively afforded after a F/SCF₃ ligand exchange.

In 2015, Ye and Liu reported the palladium-catalyzed trifluoromethylthiolation of 2-arylpyridine derivatives using the Billard reagent II (Scheme 7) [120]. Unlike Shen’s methodology (Scheme 5), the use of benzoyl chloride was necessary to activate the trifluoromethylthiolated reagent [120]. Unsubstituted and pyridines substituted derivatives 15 were very efficiently ortho-trifluoromethylthiolated (17 examples, up to 91% yield). This methodology was tolerant to electron-donating and electron-withdrawing groups as well as halogens (16b-f, 62–91% yields). The meta-substituted (16f) and disubstituted (16j) products were obtained in high yields (73% and 79%, respectively).

In 2018, Anbarasan and co-workers described a palladium-catalyzed trifluoromethylthiolation of arenes by C–H bond activation bearing several directing groups (Scheme 8) [121]. With
Scheme 8: Palladium-catalyzed ortho-trifluoromethylthiolation of 2-arylpyridine and analogs reported by Anbarasan [121].

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this methodology, the functionalization of 2-phenylpyridine derivatives was possible (10 examples, up to 81% yield) using the Billard reagents III or IV [122]. The trifluoromethylthiolation of 2-arylpirdines substituted by electron-donating groups such as methyl, methoxy or methylthio groups (17b–d) or by halogen (17e) was achieved (Scheme 8, up to 77% yield). Note that in case of disubstituted 2-(4-ethoxy-3-fluorophenyl)pyridine (17h), the expected product 18h was isolated in 31% yield. Moreover, selective oxidation of the SCF₃ residue into the corresponding sulfoxide and the sulfone was possible and the corresponding products were obtained in 98% and 95% yields, respectively.

This reaction was successfully expanded to the trifluoromethylthiolation of derivatives bearing various directing groups such as substituted pyridines (4 examples, up to 84% yield, 19a, 84% yield), substituted pyrazole derivatives (6 examples, up to 55% yield, 19b, 55% yield), pyrimidine (19c,
31% yield) as well as quinoline and isoquinoline (19d and 19e, 65% and 66% yields, respectively). In addition, the trifluoromethylthiolated benzo[\(h\)]quinoline 20f was obtained in good yield (63%).

The same year, Besset and co-workers reported a palladium-catalyzed C(sp\(^2\))–SC\(_{\text{F}}\(_3\) bond formation on amides derived from 8-aminooquinoline as a cleavable directing group in the presence of the Munavalli reagent V (Scheme 9, 12 examples, up to 71% yield) [106]. Depending on the substitution pattern on the aromatic ring, the amides were mono- or difunctionalized. Indeed, meta- and ortho-substituted derivatives (21a–d) were selectively trifluoromethylthiolated while para-substituted substrates led to the difunctionalized products 22e and 22f. Within these reaction conditions, the polysubstituted derivative 21g was also functionalized in high yield (71%).

Pleasingly, other metals have been also successfully applied for the trifluoromethylthiolation of aromatic derivatives by C(sp\(^2\))–H bond activation such as Rh(III) and Co(III)-based catalysts as depicted below.

**Rhodium catalysis:** In 2015, the group of Li disclosed the Cp*Rh(III)-catalyzed regioselective trifluoromethylthiolation of N-substituted indoles with (substituted) pyridines or pyrimidine as the directing groups (Scheme 10) [123]. The selective trifluoromethylthiolation of indoles at the C2 position was achieved in the presence of N-(trifluoromethylthio)saccharine (VI, Shen’s reagent) as both oxidant and electrophilic source (18 examples, up to 91% yield). Indoles bearing various electron-donating and electron-withdrawing groups as well as halogens at the C5-position and at the C6-position were functionalized in high yields (24a–f, 82–87% yields). The substitution of the indole at the C3-position did not impact the reaction and the product 24g was obtained in 91% yield. Substituted pyridines and pyrimidine (24h and 24i) were also used as directing groups (7 examples, up to 86% yield). This methodology was extended to the functionalization of other heteroaromatic derivatives (24j, 87% yield). It should be noted that the presence of zinc triflate, a Lewis acid, was used for the activation of the electrophilic source VI.

**Cobalt catalysis:** In 2017, Wang described the Cp*Co(III)-catalyzed trifluoromethylthiolation of 2-phenylpyridine derivatives using AgSCF\(_3\) (Scheme 11) [124]. This methodology allowed the functionalization of several aromatic compounds bearing a pyridine or a pyrimidine as a directing group (20 examples, up to 65% yield). The reaction proceeded smoothly with substrates bearing an electron-donating group (25b,c), halogen (25d) or withdrawing group (25e) and the desired SCF\(_3\)-containing products were obtained in moderate to good yields. The functionalization of trisubstituted arene 25g and

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**Scheme 9:** Mono- and ditrifluoromethylthiolation of benzamide derivatives derived from 8-aminooquinoline using PdCl\(_2\) as catalyst reported by Besset and co-workers [106]. a2.2 equiv of the fluorinated source V were used.
Scheme 10: Regioselective Cp*Rh(III)-catalyzed directed trifluoromethylthiolation reported by the group of Li [123]. Cp* = pentamethylcyclopentadienyl.

heteroarene 25h was also possible leading to the corresponding products 26g and 26h in moderate yields (23% and 42% yields, respectively). Regarding the reaction mechanism, the active Co(III) complex G was obtained from the dimeric catalyst [Cp*CoI2]2 in the presence AgSCF3 and/or NaOPiv·H2O. Then, the reversible formation of the metallacycle H occurs, which after a ligand exchange in the presence of AgSCF3 leads to the formation of adduct J. The product 26 is released via a reductive elimination step, generating at the same time the reduced cobalt Cp*Co(I), which is converted to the active catalyst after oxidation.

The same year, Yoshino and Matsunaga described a similar methodology, using the cobalt(III) complex [Cp*Co(CH3CN)3](SbF6)2 and N-trifluoromethylthiodibenzenesulfonimide VII as electrophilic SCF3 source (Scheme 12) [125]. Under these reaction conditions, 2-arylpyridines (9 examples, up to 94% yield) or 6-arylpurines (10 examples, up to 72% yield) were ortho-trifluoromethylthiolated. The substitution pattern of the aryl part did not impact the outcome of the reaction. This methodology was also tolerant to a large range of functional groups (ester, halogen) as illustrated by the products 28c, 28j, 28d, 28k, and 28g.

I.2) Transition-metal-catalyzed C–H trifluoromethylthiolation of vinyllic C(sp2) centers

Several research groups have been interested in the development of strategies for the formation of vinyllic C(sp2)–SCF3 bonds, offering an efficient tool towards the synthesis of challenging Z-isomers.

Palladium catalysis: In 2018, Bouillon and Besset described the first example of a selective palladium-catalyzed trifluoromethylthiolation of α-arylacrylamides derived from 8-aminoquinolines 29 by C–H bond activation (Scheme 13) [104]. Using the Munavalli reagent V as an electrophilic SCF3 source, this diastereoselective method selectively led to the formation of Z-isomers and turned out to be robust (not air or moisture sensitive). Under these mild reaction conditions, a panel of α-(hetero)arylacrylamides were trifluoromethylthiolated in good to high yields. Acrylamides substituted at the α-position by an aryl bearing an electron-donating group (OMe) or halogen (Cl) at the para-position were readily functionalized (30b and 30c, 70% and 75% yields, respectively). The substitution of the arene with a CF3 residue at the meta-position or with a methoxy group at the ortho-position did not have any impact to the outcome of the reaction (30e and 30f, 62% and 78% yields, re-
spectively). It should be noted that acrylamides bearing a disubstituted arene (29g,h) and an heteroaryl (29i) at the α-position were also suitable substrates for this reaction. Finally, acrylamides bearing a methyl group at the α-position (29j) or the α,β-dimethylated acrylamide (29k) were suitable substrates albeit the corresponding products were obtained in 30% and 20% yields, respectively. Several mechanistic experiments revealed that the C(sp²)–H bond activation step was reversible and represented the rate-determining step (KIE = 2.4). First, the chelation of the palladium(II) catalyst with the bidentate
directing group, followed by the C(sp³)–H bond activation involving a concerted metalation–deprotonation pathway affords the metallacycle K. After an oxidative addition in the N–SCF₃ bond of the Munavalli reagent V, the palladium(IV) species L is obtained. Finally, the reductive elimination affords the product and regenerates the catalyst. The same year, Besset and co-workers extended this methodology to a larger class of acrylamides [106].

I.3) Transition-metal-catalyzed trifluoromethylthiolation of aliphatic C(sp³)–H bonds

Despite the important progresses presented in the previous section, some limitations of the trifluoromethylthiolation persist. In particular, the functionalization of a C(sp³)–H bond with a trifluoromethylthiolated moiety by transition-metal-catalyzed C–H activation remains a challenging task both in terms of reactivity and selectivity.

In 2015, Besset reported the first C(sp³)–SCF₃ bond formation of unactivated primary C(sp³) centers by transition-metal-catalyzed C–H activation with the Munavalli or the Billard reagents as the trifluoromethylthiolation source (Scheme 14) [126]. Using a bidentate directing group, this methodology allowed the functionalization of a large range of aliphatic amides with a primary β-C(sp³)–H bond (21 examples, up to 53% yield). The methodology was applied to the functionalization of a series of amides having an α-quaternary center (α,α-dialkyl (31a), α-alkyl,α-benzyl derivatives 31c–f) as well as to an amide with an α-tertiary center (31b) and pleasingly, the presence of α-C–H bonds did not have a significant impact on the outcome of the reaction. It should be noted that this methodology afforded the products with a high regioselectivity, and no incorporation of the SCF₃ moiety on the benzylic or at the C5 position of the quinoline part of the directing group was observed. Note that in 2018, Besset and Lebel developed a more efficient process for the palladium-catalyzed trifluoromethylthiolation by C–H bond activation under continuous flow conditions [127].

I.4) Difluoromethylthiolation of aromatic and vinylic C(sp²)–H bonds (C–SCF₂H and C–SCF₂CO₂Et bonds)

More recently, researchers became interested in the synthesis of molecules substituted with original and functionalized fluorinated moieties such as SCF₂FG [128-136] (FG = functional group). In sharp contrast with the trifluoromethylthiolation reaction, only a handful of reports dealt with the incorporation of such high value-added fluorinated moieties onto C(sp²) centers by transition-metal-catalyzed C–H bond activation.
Scheme 13: Diastereoselective trifluoromethylthiolation of acrylamide derivatives derived from 8-aminoquinoline using PdCl₂ reported by Bouillon and Besset [104,106].

a 20 mol % of PdCl₂ and 2.0 equiv of SCF₃ source V were used for 36 h.
b The reaction was conducted with 30 mol % of PdCl₂ and 2.0 equiv of reagent V were used.
In 2022, He and Pan reported the first example of a difluoromethylthiolation achieved by transition-metal-catalyzed C–H bond activation (Scheme 15) [137]. Using the reagent IX and a catalytic amount of PdCl₂, they succeeded in the functionalization of acrylamides 33 derived from 8-aminoquinoline. Within these mild conditions, α-arylacrylamides substituted at para-, meta-, and ortho-positions were readily difluoromethylthiolated (34a–g, 81–95% yields). This reaction was also tolerant to a large class of functional groups (halogens, cyano, trifluoromethyl), affording products 34c, d and 34e in high yields (86%, 89% and 81%, respectively). The α-methyl-substituted acrylamide 33j also underwent difluoromethylthiolation to give product 34j in 81% isolated yield. The α,β- and β-substituted acrylamides were functionalized in high yields (34h, i and 34k–m, 71–86%). The plausible mechanism is similar as the one reported by Besset for the trifluoromethylthiolation of acrylamides derived from 8-aminoquinoline (Scheme 13).

The same year, Besset and co-workers reported the first palladium-catalyzed C(sp²)–SCF₂ bond formation by C–H bond activation (Scheme 16) [138]. In the presence of the electrophilic SCF₂CO₂Et source X, the methodology was successfully applied for the functionalization of 2-arylpyridine derivatives 35a–i as well as 2-vinylpyridine derivatives 35j–m (35 examples, up to 87% yield). The substitution pattern of the aryl substituent of the 2-phenylpyridine derivatives did not influence the reaction as for instance products 36b, 36h, and 36i were obtained in good yields (70%, 63% and 61% yield, respectively). This reaction was tolerant to a broad range of functional groups such as halogens, ester, aldehyde, cyano, and nitro (36c–g, 36–74% yield). It is noteworthy that a disubstituted compound 35j and a thiophene derivative 35k were also efficiently difluoromethylthiolated (36j and 36k, 72% and 65%, respectively). α-Substituted vinylpyridines with electron-donating or electron-withdrawing groups on the aromatic ring were functionalized and 36l and 36m were easily isolated in 73% and 81% yields, respectively. Even an α,β-disubstituted vinylpyridine 35a and the benzoquinoline 35o were smoothly functionalized showing the efficiency of the approach. Of high interest, the modularity of the SCF₂CO₂Et was highlighted by its conversion into various other fluorinated residues (amide, carboxylic acid) and its selective oxidation into the corresponding sulfoxide and sulfone.

I.5) Trifluoromethylselenolation of aromatic and vinylic C(sp²)–H bonds by palladium catalysis

Very recently, the palladium-catalyzed trifluoromethylselenolation of (hetero)aromatic and olefinic derivatives has been investigated by the group of Billard using similar catalytic systems as those depicted for the trifluoromethylthiolation reactions. Indeed, using amides 37 derived from 5-methoxy-8-aminoquinoline, the functionalization of (hetero)aromatic compounds was achieved using 20 mol % of Pd(CH₃CN)₂Cl₂ in the presence of TolSO₂SeCF₃ as the fluorinating source (14 examples, Scheme 17) [139]. Of note, when the reaction was carried out on derivatives bearing substituents at the meta- (37e and 37d) as well as at the para- (37a and 37b) positions, the corresponding products 38a–d were obtained as a mixture of mono- and disubstituted trifluoromethylselenolated derivatives with global yields ranging from 48% to 70%. When one of the ortho-posi-
tions of the aromatic ring was substituted with a Me (37e), a OMe (37f), a Cl (37g) or a CF$_3$ (37h) group, the corresponding compounds 38e, 38f, 38g, and 38h were isolated in moderate to high yields (43% to 80%). The methodology also allowed the trifluoromethylselenolation of the furan derivative 37i, which led to the desired product 38i in 30% yield. A careful monitoring of the reaction unveiled the rapid formation of the CF$_3$SeSeCF$_3$ dimer, which could be the active trifluoromethylselenating reagent in this transformation.

In 2022, Magnier, Billard and co-workers applied the previous methodology to the trifluoromethylselenolation of acrylamide derivatives [140]. Using the same directing group, a panel of α-arylacrylamide derivatives 39a–f was successfully functional-
Scheme 16: Palladium-catalyzed (ethoxycarbonyl)difluoromethylthiolation reaction of 2-(hetero)aryl and 2-(α-aryl-vinyl)pyridine derivatives reported by Besset [138].

Selected examples:
- 36a, R = H, 74%
- 36b, R = Ph, 70%
- 36c, R = Cl, 74%
- 36d, R = CO₂Me, 57%
- 36e, R = CHO, 36%
- 36f, R = NO₂, 58%
- 36g, R = CN, 55%
- 36h, R¹ = H, R² = Me, 63%
- 36i, R¹ = F, R² = H, 61%
- 36j, R¹ = F, R² = OMe, 72%
- 36k, 65%
- 36l, R = Me, 73%
- 36m, R = CF₃, 81%

Scheme 17: Pd(II)-catalyzed trifluoromethylselenolation of benzamides derived from 5-methoxy-8-aminoquinoline reported by the group of Billard [139]. The yields given are the sum of the yields of mono- and ditrifluoromethylselenolated products.

Selected examples:
- 38a, R = 4-H, 55%
- 38b, R = 4-Me, 48%
- 38c, R = 3-Me, 69%
- 38d, R = 3-OMe, 70%
- 38e, R = Me, 71%
- 38f, R = OMe, 43%
- 38g, R = Cl, 80%
- 38h, R = CF₃, 66%
- 38i, 30%

Furthermore, a series of β-substituted acrylamides 39i-m with various substituents readily underwent the trifluoromethylselenolation reaction with high selectivity in moderate yields. Finally, the SeCF₃-containing β-methylacrylamide 40n was also obtained in 37% yield.
II. Transition-metal-catalyzed fluoroalkoxylation of (hetero)arenes by C–H bond activation

II.1) Fluoroalkoxylation of aromatic C(sp²)–H bonds by transition-metal catalysis

Over the last years, key advances have been made for the formation of a C(sp²)–OCHRCF₃ bond by transition-metal-catalyzed C–H bond activation. Indeed, fluorinated ethers [71,141-153] are key compounds, with especially molecules substituted with the 2,2,2-trifluoroethoxy moiety (OCH₂CF₃), an important fluorinated group found in several bioactive compounds such as flecanide [154,155] and lansoprazole [156], as flagship molecules. Although the transition-metal-catalyzed hydroxylation and alkoxylation have been studied especially under palladium catalysis [157,158], the direct dehydrogenative 2,2,2-trifluoroethoxylation of (hetero)arenes, often using 2,2,2-trifluoroethanol as a readily available, inexpensive, and green fluorination source [159,160], is still underexplored.

In 2004, Sanford and co-workers reported the dehydrogenative 2,2,2-trifluoroethoxylation of benzo[h]quinoline under palladium catalysis [161]. Since this seminal work and in the course of their investigation towards the development of new methods for the alkoxylation of C(sp²) centers by transition-metal catalysis, few examples of transition-metal-catalyzed dehydrogenative 2,2,2-trifluoroethoxylation reactions have been reported. In 2021, the palladium-catalyzed ortho-2,2,2-trifluoroethoxylation of 3-arylcoumarins was depicted by the group of Kumar (6 examples, up to 69% yield) [162]. Further developments unveiled the use of copper catalysts for such functionalization. In 2013, the group of Daugulis described the copper-catalyzed ortho-2,2,2-trifluoroethoxylation of a 3-trifluoromethylated benzamide derived from 8-aminoquinoline, giving the corresponding product in 73% yield [149]. The group of Baidya showed that the dehydrogenative 2,2,2-trifluoroethoxylation of benzamide with another bidentate directing group was also possible in the presence of Cu(OAc)₂ and hexamethyldisilane [163]. Using N,N- and N,O-bidentate directing groups, the construction of C(sp²)–OCH₂CF₃ bonds by C–H bond activation was also reported using Ni [164] and Co catalysis [165-167]. In 2022, Volla and co-workers reported the ortho-2,2,2-trifluoroethoxylation of benzamide using an N,O-bidentate directing group by merging Co- and visible light organophotocatalysis [168].

Palladium catalysis: In 2017, Ji, Li and co-workers reported a thorough study on the Pd-catalyzed 2,2,2-trifluoroethoxylation of N-sulfonylbenzamides in the presence of PhI(OAc)₂ as oxidant and an excess of TFA (Scheme 20) [150]. The functionalization...
Scheme 19: Transition-metal-catalyzed dehydrogenative 2,2,2-trifluoroethoxylation of (hetero)aromatic derivatives by C–H bond activation [149].

Sanford, 2004 [161]a
Pd cat. b (0.7 mol %)
Phl(OAc)2 (2.0 equiv)
100 °C, 21 h, air
1 example
71% yield

Kumar, 2021 [162]a
Pd(OAc)2 (10 mol %)
K2S2O4 (2.0 equiv)
TFA (2 equiv)
80 °C, 12 h, air
6 examples
54–69% yield

Daugulis, 2013 [149]c
(CuOH)2CO3 (11 mol %)
pyridine, 110 °C, 16 h, O2
1 example
73% yield

Baidya, 2016 [163]a
Cu(OAc)2 (1.0 equiv)
SiMe4 (1.0 equiv)
K2CO3 (3.0 equiv)
DMSO, 80 °C, 3 h, air
1 example
69% yield

Sundararaju, 2018 [164]a
Ni(OAc)2·4H2O (10 mol %)
AdCOOH (20 mol %)
Ag2SO4 (2.0 equiv)
Na2CO3 (2.0 equiv)
DMF, 130 °C, 16 h, air
1 example
31% yield

Niu and Song, 2015 [165]a
Co(OAc)2·4H2O (20 mol %)
Ag2O (1.0 equiv)
NaOAc (2.0 equiv)
60 °C, 12 h, air
1 example
67% yield

Wei and Niu, 2015 [167]a
Cp*Co(CO)2 (20 mol %)
Ag2O (2.0 equiv)
Na2CO3 (1.0 equiv)
70 °C, 12 h, air
1 example
85% yield

Yang, Wu and Wu, 2019 [166]a
Co(OAc)2·4H2O (20 mol %)
DMAP (2.0 equiv)
Ag2CO3 (2.0 equiv)
o-chlorotoluene, 100 °C, 20 h, air
1 example
51% yield
1 example with (CF3)2CHOH
45% yield

Volla, 2022 [168]a
Co(OAc)2·4H2O (10 mol %)
eosin Y (5 mol %)
NaOAc (1.0 equiv)
rt, 12 h, O2
blue LED
1 example
65% yield

of diversely substituted derivatives bearing either electron-donating groups (41b, 41d, and 41f) and electron-withdrawing groups (41c, 41e, and 41g) was achieved (19 examples, up to 93% yield). Of note, the transformation was also efficient with disubstituted substrates such as 41h–j. The authors suggested the following mechanism. After formation of the metal-lacyle O, the latter is oxidized leading to the Pd(IV) species P. After a ligand exchange, the intermediate Q is generated.
Scheme 20: Pd(II)-catalyzed ortho-2,2,2-trifluoroethoxylation of N-sulfonylbenzamides reported by the group of Ji and Li [150].

Ji and Li, 2017 [150]

\[
\begin{align*}
\text{R} &= \text{Me or NO}_2 \\
\text{41} &\quad \text{Pd(OAc)}_2 (10 \text{ mol } \% ) \quad \text{Phi(OAc)}_2 (2.0 \text{ equiv} ) \quad \text{TFA (20.0 \text{ equiv})} \\
\text{CF}_3\text{CH}_2\text{OH} &\quad 24 \text{ h, rt, air} \\
\text{42} &\quad 19 \text{ examples up to } 93\% \text{ yield}
\end{align*}
\]

selected examples

\[
\begin{align*}
\text{42a}, \text{R} &= \text{H}, 62\% \\
\text{42b}, \text{R} &= 4-\text{OMe}, 55\%^{a,b} \\
\text{42c}, \text{R} &= 4-\text{F}, 55\%^a \\
\text{42d}, \text{R} &= 3-\text{Me}, 61\%^c \\
\text{42e}, \text{R} &= 3-\text{Br}, 52\%^a \\
\text{42f}, \text{R} &= 2-\text{Me}, 93\% \\
\text{42g}, \text{R} &= 2-\text{Cl}, 62\%
\end{align*}
\]

Finally, a reductive elimination step affords the expected functionalized product 42 and regenerates the catalyst.

The 2,2,2-trifluoroethoxylation reaction is not restricted to amides. In 2020, Yorimitsu and co-workers developed a methodology allowing the formation of a C(sp^2)-OR bond through palladium-catalyzed C–H bond activation [169]. Using Pd(OPiv)\(_2\) as the catalyst in the presence of Phi(OAc)\(_2\) (PIDA), the naphthalene sulfoxide 43a was 2,2,2-trifluoroethoxylated in 82% yield (Scheme 21). The substituent of the sulfoxide part does not impact the efficiency of the reaction as illustrated by the synthesis of compounds 44b and 44c. The presence of an electron-donating substituent in the \textit{para}-position of the directing group was found to be deleterious for the reaction since 44d was obtained in 31% yield while its brominated analog 44e was isolated in 70% yield. Mechanistic studies indicated that the C–H bond activation event was the rate-limiting step and the authors suggested a similar mechanism to the one...
depicted in Scheme 20: formation of a palladacycle thanks to a concerted metalation deprotonation (CMD) process followed by oxidation, ligand exchange with CF$_3$CH$_2$OH, and finally, reductive elimination affording the expected product and regenerating the catalyst. Gratifyingly, the approach was applied to the incorporation of other fluorinated moieties such as OCH$_2$CF$_2$H and OCH$_2$(CF$_2$)$_n$H ($n = 2$ or 4) and gave compounds 46a–c.

Thanks to the transient directing group strategy [35,170-182], efficient methodologies for the functionalization of previously reluctant compounds such as benzaldehyde derivatives were developed, and key advances are depicted below.

In 2021, Wang and co-workers described the selective palladium-catalyzed ortho-2,2,2-trifluoroethoxylation of a series of benzaldehydes (Scheme 22, 35 examples) using the amino acid $l$-valine in the presence of K$_2$S$_2$O$_8$ and TFA at 80 °C [153]. This reaction proved to be highly tolerant to various substituents including a CF$_3$ group at the ortho-, meta- and para-positions (48m, 48j, and 48f, respectively), halogens (48d, 48h, and 48l), an ester moiety (48e and 48i), and a methoxy group (48c). Note that even di- and trisubstituted benzaldehydes 47n–q were smoothly functionalized under these conditions. The authors also suggested a plausible mechanism. The amino acid acts as an organocatalyst and first reacts with the benzaldehyde 47 to generate the transient directing group (47'). Then, formation of the palladacycle (species R) followed by its oxidation to a Pd(IV) intermediate and a ligand exchange with 2,2,2-trifluoroethanol leads to the formation of the species S. The latter complex S undergoes a reductive elimination leading to the compound 48' along with the regeneration of the palladium catalyst. Finally, after hydrolysis of 48', the expected product 48 is afforded together with the organocatalyst.
Scheme 22: Pd(II)-catalyzed selective ortho-2,2,2-trifluoroethoxylation of benzaldehyde derivatives by means of the transient directing group strategy as reported by the group of Wang [153].

Then, the group of Sun and Wang used a similar approach for the 2,2,2-trifluoroethoxylation of benzaldehydes under palladium catalysis using the amino acid 51 as organic catalyst in the presence of the fluoropyridinium salt 52 (19 examples, up to 88% yield, Scheme 23) [183]. Pleasingly, the methodology was extended to the formation of C(sp²)–OR_F bonds starting from...
be benzaldehyde (OR_F = 2,2-difluoroethoxy 50g, 2,2-difluoropropoxy 50h, and 1,1,1,3,3,3-hexafluoroisopropoxy 50i).

II.2) Fluoroalkoxylation of aliphatic C(sp^3)–H bonds by transition-metal-catalysis

The functionalization of C(sp^3) centers by transition-metal-catalyzed C–H bond activation remains highly challenging [24,184-191]. In particular, the 2,2,2-trifluoroethoxylation of aliphatic derivatives is still limited to a handful of examples as illustrated by the two examples depicted in Scheme 24 [71,192]. Using a bidentate directing group (namely NHPA and CONHPIP for 53 and 55, respectively), the groups of Chen [71] and Shi [192] independently reported the palladium-catalyzed selective 2,2,2-trifluoroethoxylation of aliphatic amines and amides at the γ and β positions, respectively, using trifluoroethanol as fluorination source in the presence of PIDA. Hence, an efficient access to the corresponding monoether derivatives 54 and 56 in 71% and 65%, respectively, were obtained.

Conclusion

In summary, this review provides an overview of the major developments made over the last years for the synthesis of fluorinated compounds by transition-metal-catalyzed C–H bond activation. This review focused on the construction of C(sp^2)–XR_F bonds and C(sp^3)–XR_F bonds with an emphasis on the trifluoromethylation reaction and transformations using emergent fluorinated residues (SCF_3, SeCF_3, SCF_2H, SCF_2CO_2Et or OCH_2CF_3 groups). Well-designed catalytic systems and suitable fluorinating sources were the key to success for these major developments.

Despite these advances, synthetic challenges still need to be overcome. These synthetic tools are so far still restricted to some fluorinated moieties and extension to other high value-added fluorinated residues [131,136,193-208] is of high importance. Besides, in comparison with the functionalization of C(sp^2) centers on aromatic and vinylic derivatives, transition-metal-catalyzed functionalization of C(sp^3)–H bonds remains largely underexplored to date. Furthermore, the development of enantioselective transformations allowing the synthesis of enantioenriched fluorine-containing compounds by transition-metal-catalyzed C–H bond activation will have a significant impact as for instance an access to pharmacologically relevant derivatives. Finally, the use of abundant non-noble transition metals [209-211] in such reactions combined or not with modern technologies (photocatalysis and electrocatalysis) is still underexplored and any advances will be of high importance especially from a sustainability point of view aiming at developing greener synthetic routes towards fluorinated molecules.
Scheme 24: Pd(II)-catalyzed selective 2,2,2-trifluoroethoxylation of aliphatic amides using a bidentate directing group reported by the groups of Chen [71] and Shi [192].

Funding
This work has been partially supported by University of Rouen Normandy, INSA Rouen Normandy, the Centre National de la Recherche Scientifique (CNRS), European Regional Development Fund (ERDF), Labex SynOrg (ANR-11-LABX-0029), Carnot Institute I2C, the graduate school for research XL-Chem (ANR-18-EURE-0020 XL CHEM), Carnot Institute I2C, the graduate school for research XL-Chem (ANR-18-EURE-0020 XL CHEM), and Region Normandie. L.M. and T.B. thank the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation program (grant agreement no. 758710). F.D. thanks Labex SynOrg (ANR-11-LABX-0029) and the Region Normandy (RIN 50% program) for a doctoral fellowship.

ORCID iDs
Louis Monsigny - https://orcid.org/0000-0001-9325-1316
Floriane Doche - https://orcid.org/0000-0001-9317-3274
Tatiana Besset - https://orcid.org/0000-0003-4877-5270

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Direct C2–H alkylation of indoles driven by the photochemical activity of halogen-bonded complexes

Martina Mamone¹, Giuseppe Gentile¹, Jacopo Dosso¹, Maurizio Prato¹,²,³ and Giacomo Filippini*¹

Abstract
A light-driven metal-free protocol for the synthesis of sulfone-containing indoles under mild conditions is reported. Specifically, the process is driven by the photochemical activity of halogen-bonded complexes formed upon complexation of a sacrificial donor, namely 1,4-diazabicyclo[2.2.2]octane (DABCO), with α-iodosulfones. The reaction provides a variety of densely functionalized products in good yields (up to 96% yield). Mechanistic investigations are reported. These studies provide convincing evidences for the photochemical formation of reactive open-shell species.

Findings
Direct replacement of carbon–hydrogen (C–H) bonds with new carbon–carbon (C–C) and carbon–heteroatom (C–X) bonds has been and still is a central topic in organic synthesis [1,2]. Historically, organic chemists have extensively relied on the use of noble-metal-based catalysts (e.g., Pd, Rh, Ir, among others) to achieve such type of functionalization [3-5]. However, reliance on noble metal complexes has been constantly declined over recent years due to cost, availability, and toxicity, therefore discouraged by the modern guidelines towards implementation of sustainable chemical production schemes [6]. In the last decades, organic photochemistry has become a prominent tool to guide the development of greener and more convenient synthetic protocols [7-12]. In this context, photochemical approaches based on electron donor–acceptor (EDA) complexes have been successfully exploited to drive the direct C–H functionalization of a large number of organic substrates [13-18]. In this approach, an electron acceptor substrate (“A”) and a donor molecule (“D”) interact to form a new aggregate defined as
EDA complex (Figure 1a). Although the two molecular entities might not directly absorb visible light, the newly formed complex usually presents a charge transfer state which results in a bathochromic shift of the absorption towards the visible range [19,20]. Upon light irradiation, the EDA complex may undergo an intramolecular single-electron-transfer (SET) process to produce a radical ion pair (D\(^{+}\), A\(^{-}\)). To avoid the occurrence of a back-electron-transfer (BET), a suitable leaving group (LG) needs to be included in one of the precursors. In this manner, reactive intermediates (e.g., radical species) may be generated in solution through the irreversible fragmentation of the substrates [15,21,22]. These intermediates eventually react to yield the final products “A–D.” This approach is not limited to reagents with appropriate donor–acceptor characteristics [13,19].

Indeed, sacrificial electron donors and electron-deficient radical precursors can be used to form photoactive EDA complexes. Specifically, these aggregates can be employed to photochemically generate electrophilic radicals that can drive the functionalization of suitable electron-rich substrates [23].

Exploiting this strategy, here we report a novel metal-free methodology for the direct homolytic aromatic substitution (HAS) reaction of indoles 1 with \(\alpha\)-iodosulfones 2 to yield the alkylated derivatives 3 (Figure 1b). Indoles play a crucial role in many natural and industrial processes. Therefore, the direct chemical manipulation of the indole system is a matter of paramount importance [24-27]. Moreover, the sulfonyl group is an extremely versatile chemical moiety which may be easily transformed into different functionalities employing conventional synthetic methods. As an example, the sulfonyl group removal under simple reductive treatment may give access to important methylated compounds [12,21]. This operationally simple approach occurs at ambient temperature and under visible-light irradiation. Interestingly, this method employs 1,4-diazabicyclo[2.2.2]octane (DABCO) as sacrificial donor in the EDA complex formation with 2. To test the feasibility of our design plan, we focused on the reaction between 3-methylindole (1a, 2 equiv) and \(\alpha\)-iodosulfone 2a (Table 1).

The experiments were conducted at ambient temperature in acetonitrile (0.5 M) and under irradiation by a Kessil lamp at 456 nm. When adding 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) as sacrificial donor (1 equiv), the desired product 3a was formed in good chemical yield (entry 1, Table 1). Control experiments were conducted to obtain more mechanistic clues (entries 2–5, Table 1). An experiment revealed how the exclusion of light completely suppressed the process, therefore establishing the photochemical nature of the transformation (entry 2, Table 1). In addition, we confirmed that DBU was essential for the reactivity, since no reaction occurred in its absence (entry 3, Table 1). Reactivity was also inhibited under an aerobic atmosphere and in the presence of 2,2,6,6-tetramethylpiperidinyloxyl (TEMPO). These experiments are consonant with the occurrence of a radical mechanism (entries 4 and 5, Table 1) [28]. Afterwards, the effect of the chemical nature of the sacrificial donor on the reaction was investigated (entries 6–9, Table 1). In particular, we employed 2,6-lutidine, 1,1,3,3-tetramethylguanidine (TMG), triethylamine (NEt\(_3\)), and DABCO. Interest-
Table 1: Optimization of the reaction conditions and control experiments.

<table>
<thead>
<tr>
<th>entry</th>
<th>donor</th>
<th>[M]</th>
<th>1a:2a donor (equiv)</th>
<th>light source (nm)</th>
<th>yield (%)</th>
<th>note</th>
</tr>
</thead>
<tbody>
<tr>
<td>1b</td>
<td>DBU</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>56</td>
<td>a</td>
</tr>
<tr>
<td>2</td>
<td>DBU</td>
<td>0.5</td>
<td>1:2:1</td>
<td>light off</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>–</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>4c</td>
<td>DBU</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>5d</td>
<td>DBU</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>2,6-lutidine</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>0</td>
<td></td>
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<tr>
<td>7</td>
<td>TMG</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>64</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>NEt₃</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
<td>64</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>DABCO</td>
<td>0.5</td>
<td>1:2:1</td>
<td>456</td>
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</tr>
<tr>
<td>10</td>
<td>DABCO</td>
<td>0.25</td>
<td>1:2:1</td>
<td>456</td>
<td>64</td>
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<tr>
<td>11</td>
<td>DABCO</td>
<td>1.0</td>
<td>1:2:1</td>
<td>456</td>
<td>69</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>DABCO</td>
<td>0.5</td>
<td>1:1:1</td>
<td>456</td>
<td>65</td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>DABCO</td>
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<td>2:1:1</td>
<td>456</td>
<td>73</td>
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<tr>
<td>14</td>
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<td>0.5</td>
<td>2:1:1.5</td>
<td>456</td>
<td>95</td>
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<tr>
<td>15e</td>
<td>DABCO</td>
<td>0.5</td>
<td>2:1:1.5</td>
<td>456</td>
<td>18</td>
<td></td>
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<tr>
<td>16f</td>
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<td>2:1:1.5</td>
<td>456</td>
<td>60</td>
<td></td>
</tr>
</tbody>
</table>

a Yield determined by 1H NMR spectroscopy using 1,3,5-trimethoxybenzene as the internal standard. b Conditions: indole 1a (0.1 mmol), α-iodosulfone 2a (0.2 mmol), acetonitrile (MeCN, 200 μL), donor (0.1 mmol), ambient temperature. c Reaction in air. d Reaction performed in the presence of 2 equiv of TEMPO. e Reaction performed in hexane as solvent. f Reaction performed in methanol as solvent.

ingly, the use of DABCO provided the best result in terms of reactivity, yielding compound 3a in 77% yield. We also observed that either increasing or decreasing the concentration of the reaction mixture did not bring any improvement (entries 10 and 11, Table 1). Then, the ratio between the reagents was optimized. In particular, the use of 2a:1a in a 1:1 ratio resulted in the formation of 3a in 65% yield (entry 12, Table 1). Moreover, we found that employing an excess of 1a (2 equiv) led to the formation of 3a in a 73% yield (entry 13, Table 1). Due to an easier purification of product 3a from the reaction crude by flash column chromatography, we decided to keep optimizing the transformation using the stoichiometric ratio indicated in entry 13 of Table 1. Importantly, product 3a was obtained in excellent yield (95%) using 1.5 equivalents of DABCO (entry 14, Table 1). In addition, the use of hexane as solvent provided the desired product 3a in low chemical yield (entry 15, Table 1). On the other hand, 3a was obtained in moderate yield (60%) using methanol as solvent (entry 16, Table 1). To shed light on the reaction mechanism, the formation of an EDA complex between the α-iodosulfone 2a and DABCO was investigated using both UV–vis and nuclear magnetic resonance (NMR) spectroscopy [29].

In particular, the optical absorption spectra of substrate 2a (green dotted line), DABCO (red dotted line), and the solution containing both 2a and DABCO (blue line) were recorded in acetonitrile (Figure 2). Specifically, it was observed that the addition of DABCO to the solution of 2a induced a bathochromic shift of the absorption spectrum towards the visible region, thus indicating the formation of an EDA complex between these chemical species. Importantly, we also confirmed that indole 1a and 2a do not form a photoactive EDA complex when mixed in solution (see Figure S1 in Supporting Information File 1). To further corroborate the hypothesis of an EDA complex being at the roots of the observed reactivity, NMR studies were also performed on samples containing the α-iodosulfone 2a and different concentrations of DABCO in deuterated acetonitrile (Figure 3).

Interestingly, a change in chemical shift of the diagnostic α-protons of 2a was displayed upon addition of increasing amounts of DABCO, suggesting the presence of the halogen-bonding interaction [30]. More precisely, the 1H NMR signal of the α-hydrogens (Hₐ) within 2a was found to shift to lower ppm values because the Hₐ nuclei have been affected by higher elec-
Figure 2: Optical absorption spectra recorded in acetonitrile in 1 cm path quartz cuvettes. [DABCO]: 0.5 M; [2a]: 0.5 M.

Figure 3: ¹H NMR titration of DABCO in a solution of 2a in ACN-d₅ to detect their halogen-bonding association through the shift of the signal of Hα.

The reaction could satisfactorily tolerate a diverse set of α-iodosulfones 2 to deliver the corresponding products 3a–d from moderate to excellent yields (up to 73% yield). We found that different indoles actively participated in the photochemical alkylation, leading to the products 3e–i (up to 96% yield). It is worth noting that derivatives 3e–g were isolated in moderate yields as single regioisomer since the alkylation step took place exclusively in position 2 of the starting indoles. As limitation,
Figure 4: Proposed reaction mechanism for the photochemical alkylation of 1a with the α-iodosulfone 2a in the presence of DABCO.

Scheme 1: Study of scope of the HAS reaction between indoles 1 and α-iodosulfones 2. Yields in parentheses were determined by $^1$H NMR analyses, using 1,3,5-trimethoxybenzene as an internal standard.
we observed that indole-3-carboxaldehyde (1g) was not a suitable substrate for this transformation.

Conclusion
In conclusion, we reported a novel photochemical method for the direct C–H alkylation of indoles with α-iodosulfones. This approach exploits the photochemical activity of halogen-bonded EDA complexes, formed between α-iodosulfones and DABCO, that are able to produce reactive C-centered radicals under mild reaction conditions.

Supporting Information
Supporting Information File 1
General procedures and products characterization. [https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-19-42-S1.pdf]

Acknowledgements
M.P. is the AXA Chair for Bionanotechnology (2016–2023).

Funding
The authors gratefully acknowledge the University of Trieste, INSTM and the Maria de Maeztu Units of Excellence Program from the Spanish State Research Agency (Grant No. MDM-2017–0720). G.F. and J.D. kindly acknowledge FRA2022 INSTM and the Maria de Maeztu Units of Excellence Program. The authors gratefully acknowledge the University of Trieste, F. Glorius, 2016–2023.

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The definitive version of this article is the electronic one which can be found at: https://doi.org/10.3762/bjoc.19.42
C3-Alkylation of furfural derivatives by continuous flow homogeneous catalysis

Grédy Kiala Kinkutu1,2, Catherine Louis2, Myriam Roy1, Juliette Blanchard*2 and Julie Oble*1

Abstract

The C3-functionalization of furfural using homogeneous ruthenium catalysts requires the preinstallation of an ortho-directing imine group, as well as high temperatures, which did not allow scaling up, at least under batch conditions. In order to design a safer process, we set out to develop a continuous flow process specifically for the C3-alkylation of furfural (Murai reaction). The transposition of a batch process to a continuous flow process is often costly in terms of time and reagents. Therefore, we chose to proceed in two steps: the reaction conditions were first optimized using a laboratory-built pulsed-flow system to save reagents. The optimized conditions in this pulsed-flow mode were then successfully transferred to a continuous flow reactor. In addition, the versatility of this continuous flow device allowed both steps of the reaction to be carried out, namely the formation of the imine directing group and the C3-functionalization with some vinylsilanes and norbornene.

Introduction

The conversion of biomass derivatives into value-added products is one of the key branches of green chemistry and of the development of a sustainable chemical industry [1-4]. Furfurals, which are versatile platform molecules derived from renewable lignocellulose present in agricultural wastes [5-8], have proven to be of great importance for the preparation of value-added chemicals, biofuels, as well as monomers for materials science [9-15]. In this context, their functionalization is fundamental to further improve their inclusion in fine organic synthesis and industrial processes. For this reason, in recent years, innovative protocols for the formation of new bonds on furfural derivatives have been developed. In particular, their direct functionalization by transition-metal-catalyzed C–H activation processes [16-18] has become a major area of interest where only a few
methods have been reported so far. Most examples concern functionalization at C5, which is the most reactive site. In contrast, C3-functionalizations of the formyl-furan unit via directing groups, as well as C4-functionalizations, have been much less studied [19,20].

Within the framework of a large project oriented towards the selective formation of new bonds from furfural derivatives without changing the redox state of the aldehyde function, we have developed a number of directed Ru(0)-catalyzed C3-functionalizations of furfurylimines, such as alkylation [21], arylation [22], alkenylation [23] and acylation [24], as well as an Ir-catalyzed directed C3-silylation (Scheme 1a) [25]. These batch processes rely on the use of a homogeneous metal catalyst at elevated temperatures necessary to cleave the C3–H bond by oxidative addition. These experimental conditions, easily used in the laboratory, are potentially problematic for scale-up due to efficiency and safety issues (related to the high temperature). Thus, despite the synthetic interest of the molecules that can be obtained, transfers to industry are difficult. In order to circumvent this drawback, we considered transposing these batch reactions to a flow chemistry process.

In recent years, the use of continuous flow chemistry in synthetic organic synthesis has increased dramatically and has rapidly become a routine tool for classical synthesis [26-29]. In particular, many efforts have been devoted to the development of flow alternatives for transition-metal-catalyzed cross-couplings [30] and for some C–H functionalizations [31]. Nevertheless, there are very few flow processes that have been implemented to functionalize furfurals, the scarce examples being only based on photochemical processes [32-34]. The current strong interest in continuous flow strategies is related to the modernization of flow equipment providing chemists, not only a unique control of reaction parameters, such as improved mass and heat transfer, but also reduced safety risks and increased reproducibility of the results [29,35,36]. These features should therefore allow us to scale up our directed C3-functionalizations of furfurylimines.
under safe reaction conditions while providing products in shorter reaction times. In addition, the ability to couple multiple reactors with a flow apparatus could also enable us to perform these functionalizations directly from furfural by forming the imine in a first reactor. It should be noted that, in batch, in-situ imine formation is currently impossible with catalytic or stoichiometric amounts of amine due to decarbonylation of furfural under the reaction conditions [21]. We thus present here an adaptation of our Ru(0)-catalyzed C3-alkylation strategy of furfural derivatives to a continuous flow system (Scheme 1b).

Results and Discussion
First optimization with a home-made pulsed-flow setup
We undertook the optimization of this flow strategy for the C3-alkylation reaction (Murai reaction) [37,38] of the furfurylimine 1 bearing a removable N,N'-bidentate directing group. In a previous study, this starting material had proved to be the most reactive imine in batch, leading, in the presence of 5 mol % of [Ru3(CO)12] and 3 equivalents of triethoxyvinylsilane in toluene at 150 °C after 5 h, to the alkylated aldehyde 2a with 62% yield, after purification on silica gel (Scheme 2) [21,39].

The flow reactions for this first optimization were performed using a home-made setup based on an HPLC apparatus (Jasco) equipped with an injection valve (Rheodyne) comprising a 105 μL loop into which the reaction mixture is loaded and then pushed by the solvent delivered by the HPLC pump (see Supporting Information File 1, p. S8). All the content of the loop is thus sent into the reactor. This system is coupled to a gas chromatography oven, in which the stainless-steel tubular reactor (length: 4.6 m, internal diameter of 0.8 millimeter, corresponding to a volume of 2.31 mL) is placed. The system pressure is controlled by a back-pressure regulator (BPR) to keep a pressure of about 130 bar, i.e., at a pressure much higher than that which causes the solvent (toluene) to boil in the reaction temperature range (150–200 °C). This homemade, pulsed-flow setup was used for optimizing the protocol while saving on reactants and catalyst.

Initial tests with the commercial complex [Ru3(CO)12] at high temperature with different residence times provided the desired C3-alkylated imine 12a in NMR yields ranging from 30% to 65% (Table 1, entries 1–3 and Table S1 in Supporting Information File 1, p. S10). A continuous flow system was thus found to be compatible with the realization of this type of C–H functionalization. This process led to a significant reduction of the reaction time compared to the batch, in particular by increasing the temperature to 200–250 °C, without significant losses of activity and selectivity. Unfortunately, with this catalyst, repeatability problems were detected (yield fluctuation of approximately 20%) which could be assigned to the low solubility of this catalyst in toluene. In order to overcome these problems, we synthesized tri ruthenium carbonyl complexes with phosphine ligand(s), namely (triethoxysilyl)ethyl)phosphine L1 or triphenylphosphine [40-42]. Their synthesis, well-described in the literature, is detailed in Supporting Information File 1 (pp. S3–S6). Moreover, a kinetic study carried out in batch in the presence of the [Ru3(CO)11(L1)] (comp1), [Ru3(CO)10(L1)2] (comp2) or [Ru3(CO)9(L1)3] (comp3) catalysts allowed to show, on the one hand, the absence of solubility problems, and to discover, on the other hand, that the presence of three L1 ligands (comp3) leads to a reaction rate clearly lower than that of a catalyst carrying one or two ligands (see p. S7 of Supporting Information File 1 for the reaction kinetic curves of catalysts). In addition, the catalyst with a single L1 ligand (comp1) was found to be more reactive than the one with two ligands (comp2), and was therefore selected for further optimization. In contrast, comparison of its reaction kinetic curve with that of [Ru3(CO)12] indicates that comp1 is slightly less active than [Ru3(CO)12]. Beside these three catalysts, a fourth one [Ru3(CO)11(PPh3)] comp4, was also used for this study.

For the continuous flow reaction, we observed, for the same residence time, a slight decrease in performance with comp1.
Table 1: Optimization of the catalyst for the alkylation reaction on the homemade pulsed-flow setup.

<table>
<thead>
<tr>
<th>Entry</th>
<th>[Ru₃]</th>
<th>T [°C]</th>
<th>t_r [min]</th>
<th>Conv [%]³</th>
<th>Yield [%]⁴,⁵</th>
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</thead>
<tbody>
<tr>
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<td>Ru₃(CO)₁₂</td>
<td>165</td>
<td>90</td>
<td>58</td>
<td>45</td>
</tr>
<tr>
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<td>Ru₃(CO)₁₂</td>
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<td>30</td>
<td>90</td>
<td>65</td>
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<tr>
<td>3</td>
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<td>6</td>
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<tr>
<td>4</td>
<td>Ru₃(CO)₁₁(L₁)</td>
<td>comp₁</td>
<td>200</td>
<td>30</td>
<td>66</td>
</tr>
<tr>
<td>5</td>
<td>comp₁</td>
<td>200</td>
<td>46</td>
<td>79</td>
<td>63</td>
</tr>
<tr>
<td>6</td>
<td>comp₁</td>
<td>200</td>
<td>77</td>
<td>95</td>
<td>55</td>
</tr>
</tbody>
</table>

³Yields and conversions were calculated by ¹H NMR using p-dinitrobenzene as internal standard; ⁴Non-repeatable results; ⁵repeated four times with four different results ranging from 75% to 53%.

This can be attributed, as mentioned above, to the slightly faster reaction kinetics of the [Ru₃(CO)₁₂] catalyst compared to that of comp₁. Nevertheless, the better solubility of comp₁ in toluene allows to get around the problems of reproducibility. Moreover, increasing the residence time to 46 min resulted in 63% NMR yield of I₂a (Table 1, entry 5), which was very similar to the results obtained with [Ru₃(CO)₁₂] in 30 min. A further increase in residence time to 77 min led to a lower yield (Table 1, entry 6), probably due to products degradation under longer heating.

In addition, we found that the comp₁ [Ru₃(CO)₁₁(L₁)] was more efficient when the reaction mixture was preheated before being introduced into the reactor at 200 °C. The setup was thus modified (Scheme 3) to include a 0.8-millimeter-diameter stainless-steel preheating loop (outside the oven). An improvement in efficiency was then observed when the reaction mixture was preheated to 130 °C for 5 min. Interestingly, only traces of product I₂a were observed after 5 min at 130 °C, implying that the Murai reaction was indeed taking place only when passing through the second reactor. Finally, after some optimizations, the temperature in the second reactor could be lowered to 180 °C leading to an NMR yield of C₃-alkylated imine I₂a of 79% (Scheme 3).

Heat transfer calculations showed us that the reaction mixture rises to the set temperatures in two seconds, and more importantly, that the inlet of reactor 2 is at room temperature after passing through the tube of 50 cm that connects the two reactors. Hence, we rationalized such a performance improvement from a chemical point of view: the [Ru₃(CO)₁₂] complex is known to thermally degrade by deligation, resulting in the formation of ruthenium aggregates [43]. We therefore propose that the active species is a mononuclear carbonyl complex in which the ruthenium is coordinated to the two nitrogen atoms of the
directing group (amino-imine). Preheating for 5 minutes at 130 °C would generate it from \([\text{Ru}_3(\text{CO})_{11}(\text{L})]\), which would therefore be more accurate to consider as a precatalyst (Scheme 4, path b). The mononuclear complex would then initiate the alkylation reaction at 180 °C following elementary steps previously determined by DFT [21]. Conversely, a high
starting temperature would favor the formation of ruthenium aggregates, which could also generate, but less efficiently, the active catalyst (the mononuclear ruthenium(0) species) by leaching (Scheme 4, path a).

In order to detect the postulated reaction intermediate (Scheme 4) and the formation of ruthenium aggregates under reaction conditions, imine 1 was treated at 150 °C in toluene for 1 h with 0.33 equiv of comp4 [Ru3(CO)11(PPh3)], a catalyst analogue to comp1 but bearing a less expensive phosphine ligand (Scheme 5A). The chosen ratio of imine to catalyst was consistent with the stoichiometric amounts needed to form the postulated intermediate. The temperature of 150 °C was chosen taking into account the efficiency of the batch reaction between imine 1 and triethoxyvinylsilane (3 equiv) in the presence of 5 mol % of this catalyst at 150 °C, which leads in 5 h to the alkylated imine I2a with a NMR yield of 77% (conv. 100%).

Even though the reaction intermediate we postulated on Scheme 4 could not be detected, a solid was recovered after evaporation of the solvent and precipitation in pentane. This solid displayed a 31P NMR signal at 55.1 ppm (see Supporting Information File 1, p. S24), a value completely different from comp4 (singlet at 35.06 ppm, see Supporting Information File 1, p. S20), meaning that the ruthenium trimer was no longer present. The TEM analysis of the recovered solid phase (Scheme 5B) showed the formation of large aggregates with high electron density. Moreover, ruthenium was detected by XPS analysis (Scheme 5B); the binding energy of the 3d_{5/2} orbital was 280.94 eV, which corresponds to Ru(0). Double bonds π C=C were also detected in the sample at 285.49 eV, reflecting the presence of the PPh3 groups, but no C=O double bonds could be observed (while the presence of a C=O bond was clearly observed on the XPS spectrum of comp4 (see Supporting Information File 1, p. S6)).

These Ru aggregates were also used in the reaction with furfurylimine 1 and triethoxyvinylsilane in toluene at 150 °C for 5 h (batch conditions). In this case, only 24% of I2a were obtained (Scheme 6), a significant decrease compared to the NMR yield of 77% with comp4 as (pre)catalyst. These Ru(0) aggregates are therefore active, but the reaction kinetics are slower. While this observation is not a strict confirmation of our hypothesis regarding the formation of a monometallic complex, it is still consistent with it.

Second optimization with a continuous flow chemistry system

Following these encouraging preliminary results obtained with our home-made, pulsed-flow setup, we decided to run continuous flow experiments using a commercial setup (Vapourtec E-series flow device). This equipment offers the added advan-
Scheme 6: Ruthenium aggregate-catalyzed alkylation reaction.

With comp4 (5 mol %): 77% NMR yield, 100% conv.

588

With the optimized conditions in hand (Table 2, entry 6), we were interested in extending the scope of this furfural alkylation reaction using a flow chemistry process to other reactants. By also playing on the flow rate, this allowed us to have conditions close to the best ones observed during optimization on the pulsed-flow device, i.e., \( t_1 = 5 \text{ min} \) and \( t_2 = 50 \text{ min} \) (see Table S2 in Supporting Information File 1, p. S11). Product recovery was initiated when the system reached a steady state, based on the dispersion curves provided with the apparatus (see Supporting Information File 1, p. S13). This equipment is a medium pressure system that cannot withstand pressures exceeding 10 bar. Hence, we worked at about 7.5 bar to stay below the boiling curve of toluene in the temperature range used (180–200 °C).

Finally, it is interesting to note that when the preheating was removed, the same NMR yield was measured with 1 mol % of comp4 as catalyst (Table 2, entries 5 and 6). On the contrary, when preheating was removed with 5 mol % comp4 (entry 1 in Table 2), a drastic decrease in yield was observed, from 62% to 44% (entry 1 vs entry 7). This allowed us to assume that such preactivation is no longer necessary with 1 mol % of comp4. Thus, a lower catalyst loading, i.e., a lower concentration of the catalyst in the solution, appears to prevent, or at least greatly reduce, the formation of ruthenium aggregates as observed previously, probably by simple dilution effect. As such, the preheating was suppressed for the continuation of our investigations.

Extending the scope of the C3-alkylation of furfural in continuous flow

With the optimized conditions in hand (Table 2, entry 6), we were interested in extending the scope of this furfural alkylation reaction using a flow chemistry process to other reactants. For this, after each reaction, an aliquot of the resulting product was recovered for analysis and purification. The NMR yields were calculated on the alkylated imine before purification (based on a starting concentration of furfural of 0.35 M), and the isolated yields corresponded to the C3-alkylated aldehydes after the hydrolysis step that took place during purification. The productivity of each system is given in grams per hour (Scheme 7).
The optimized flow process conditions could be applied to a variety of vinylsilanes: trialkoxy-, triaryl-, and trialkylvinylsilanes, already used in the batch study [21]. The products 2a–e were obtained in good yields and thus with good productivity. Alkenes without silicon in the vinyl position seemed much less reactive, such as a vinylacetal, a hindered olefin (3,3-dimethyl-1-butene), or styrene. In these cases, functionalized furfurals were not isolated. In contrast, norbornene, which has a more reactive double bond due to ring tension, gave endo product 2f with an isolated yield of 49%. On the other side, disubstituted vinylsilanes proved to be ineffective, certainly because of the steric hindrance of the double bond decreasing the kinetics of the hydroruthenation step.

We also wanted to extend this alkylation reaction to a tert-butyldimethylsilyl (TBS)-protected 5-HMF derivative. Unfortunately, the yields obtained were very moderate. This reaction having slower kinetics could benefit from being performed at lower temperatures and longer residence time to reduce catalyst degradation. This is unfortunately not possible to implement at the moment with our reactors.

**Conclusion**

In conclusion, we have developed a method for the direct 2-step Ru-catalyzed alkylation of the C3–H bond of furfural by flow chemistry, via the preinstallation in a fixed bed reactor of an ortho-directing imine group that can be easily removed upon purification on silica. The reaction was found to be very efficient, with a Ru3(CO)11(PPh3) catalyst loading that could be lowered to 1 mol %, allowing for higher yields than batch conditions while requiring 5 times less catalyst. Furthermore, the interest of this flow chemistry approach lays in the scaling up of our reactions. To our great satisfaction, we could show that the productivity of the flow chemistry approach is better than the batch approach with the same catalyst (Scheme 8). This strategy represents a novel method to produce functionalized...
Scheme 7: Scope of continuous flow furfural derivative alkylation reaction.

Scheme 8: Scaling up comparison: batch and continuous flow conditions.

**Experimental**

**Triphenylphosphine triruthenium undecacarbonyl (comp4)**

Following a slightly modified procedure compared to the one reported [41], triruthenium dodecacarbonyl (1.4 g, 2.19 mmol, 1 equiv) was dissolved in freshly distilled and degassed THF (0.036 M) at 40 °C. The phosphine ligand (574.40 mg, 2.19 mmol, 1 equiv) dissolved in THF (0.11 M) was then added to the middle. The mixture was stirred at room temperature and treated dropwise with a solution of sodium benzophenone ketyl (about 0.05 equiv added) in THF (0.027 M) via a syringe until the phosphine ligand was completely consumed (monitored by TLC, ≈10 min). The solvent was then evaporated under reduced pressure. The remaining crude was purified by silica gel column chromatography using pentane as eluent, leading to 1.3 g of the desired complex as an orange solid (68% yield). 

**furfurals, providing synthetically relevant building blocks on a large scale.**
grown from a solution in Et₂O and identified by X-ray diffraction as a known phase of comp4 [41].

**General procedure for C3-alkylation of furfural in continuous flow (vapourtec)**

**Mixture A**: An oven-dried sealed tube equipped with a magnetic stirrer under argon, was loaded with furfural (240.20 mg, 2.50 mmol, 1 equiv), 2-(piperidin-1-yl)ethanamine (320.55 mg, 2.50 mmol, 1 equiv) and filled with dried toluene to a total volume of 3.5 mL.

**Mixture B**: An oven-dried sealed tube equipped with a magnetic stirrer, was loaded with triphenylphosphine triruthenium undecacarbonyl (1 mol % with regards to furfural) and degassed with argon. Vinyltriethoxysilane (3 equiv with regards to furfural) was then added to the middle, and the mixture was filled with dried toluene to a total volume of 3.5 mL. The mixture was stirred at room temperature to completely dissolve the catalyst.

The solution A is pumped into pump 1 (0.1 mL·min⁻¹) and passed through the packed bed reactor which is set at 130 °C containing MgSO₄. The residence time depends on the intrinsic volume (Vi) of this reactor, and is kept constant at ≈18 min. The solution B is pumped through pump B (0.1 mL·min⁻¹). The mixture of the two solutions A and B passed first through the coil reactor at 130 °C and then into a second coil reactor at the desired temperature. Product recovery is initiated when the system reaches a steady state, based on the dispersion curves given by the apparatus. After reaching the steady state an aliquot of the product was taken for ¹H NMR analysis using p-dinitrobenzene as an internal standard.

**3-(2-(Triethoxysilyl)ethyl)furan-2-carbaldehyde (2a)**

The reaction of mixture A containing furfural (240.20 mg, 2.50 mmol, 0.7 M) and 2-(piperidin-1-yl)ethanamine (320.55 mg, 2.50 mmol, 0.7 M), with mixture B containing triruthenium undecacarbonyl (22 mg, 0.025 mmol, 0.007 M) and vinyltriethoxysilane (1.43 g, 7.50 mmol, 1.07 M) was conducted by continuous flow chemistry, residence time ¹ = 18 min, residence time ² = 50 min. An aliquot of 0.5 mL of the product mixture was evaporated (93% conv., 77% NMR yield), and the crude was purified by silica gel column chromatography eluting with a mixture of cyclohexane/EtOAc 9:1 to give 38 mg of the desired product as an orange oil (75% yield). ¹H NMR (300 MHz, CDCl₃) δ 9.76 (s, 1H), 7.54 (d, J = 1.7 Hz, 1H), 6.49 (d, J = 1.7 Hz, 1H), 3.81 (q, J = 7.0 Hz, 6H), 2.95–2.84 (m, 2H), 1.22 (t, J = 7.0 Hz, 9H), 1.01–0.90 (m, 2H). These data are in good agreement with those reported in literature [21].

**Supporting Information**

**Supporting Information File 1**

Experimental and copies of spectra. [https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-19-43-S1.pdf]

**Acknowledgements**

The authors acknowledge MS3U of Sorbonne Université for HRMS analysis and CNRS. We also thank J. Forté for the XRD analyses (IPCM) and A. Miche for the XPS analyses (LRS).

**Funding**

The authors acknowledge H2020-WIDESPREAD-05-2020-Twinning project Biomass4Synthons (B4S: grant agreement 951996) for financial support.

**ORCID® iDs**

Myriam Roy - https://orcid.org/0000-0001-9903-5067
Juliette Blanchard - https://orcid.org/0000-0003-1935-4207
Julie Oble - https://orcid.org/0000-0002-4002-255X

**Preprint**

A non-peer-reviewed version of this article has been previously published as a preprint: doi:10.26434/chemrxiv-2023-0s98h

**References**

which can be found at:

The definitive version of this article is the electronic one available online.

It should be noted that the hydrolysis of the imine occurs during the purification on silica gel.

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Photocatalytic sequential C–H functionalization expediting acetoxymalonylation of imidazo heterocycles

Deepak Singh, Shyamal Pramanik and Soumitra Maity*

Abstract

The importance of functionalized imidazo heterocycles has often been featured in several impactful research both from academia and industry. Herein, we report a direct C-3 acetoxymalonylation of imidazo heterocycles using relay C–H functionalization enabled by organophotocatalysis starring zinc acetate in the triple role of an activator, ion scavenger as well as an acetylating reagent. The mechanistic investigation revealed a sequential sp² and sp³ C–H activation, followed by functionalization driven by zinc acetate coupled with the photocatalyst PTH. A variety of imidazo[1,2-a]pyridines and related heterocycles were explored as substrates along with several active methylene reagents, all generating the products with excellent yields and regioselectivity, thus confirming excellent functional group tolerability.

Introduction

Among all N-fused heterocycles, imidazo[1,2-a]pyridines (IPs) are the prevalent moieties in several bioactive pharmaceuticals and natural products [1-4]. Moreover, due to their susceptibility towards ‘exited-state intramolecular proton transfer’ phenomena, IPs are also effective in optoelectronics and materials sciences [5,6]. C-3-functionalized imidazo[1,2-a]pyridines are particularly familiar due to their biological and medicinal attributes [7-11]. Not surprisingly, the C-3 functionalization of IPs is a continuing interest of research in the synthetic community [12-16].

Despite many successful strategies in this field, the regioselective C–H functionalization is still challenging for chemists to combine a C(sp³) carbon of incoming functionalities and C(sp²) carbon of the IP core. The direct C-3 alkylation of imidazopyridines using active malonates and related moieties has been achieved by different routes [17-20]. However, these reactions rely either on harsh reaction conditions or require the preactivation of substrates, which limits their synthetic efficiency. A photocatalytic quaternary C-3 alkylation has also been reported recently (Scheme 1A) [21,22]. During the course of our study,
the Wu group reported a solvent-controlled chemodivergent formation of C-3 ethoxycarbonylmethylated and hydroxyalkylated IPs under visible light using water or alcohol as the source of the oxygenated group under degassed conditions [22]. However, all these photochemical methods require the usage of a substantial amount of base, the preactivation with a boron complex (B$_2$pin$_2$), and using an expensive metal-based photocatalyst [fac-Ir(ppy)$_3$] under inert atmosphere. We have recently demonstrated that aerial oxygen could be captured by alkyl radicals to install a keto-functionality onto alkenes in an organophotocatalytic way [23]. We aimed to extend this aerobic oxygenation approach to imidazo heterocycles II to install the hydroxymalonate unit onto I through sequential photoredox C–H functionalization.

Till date, there is no report of the direct incorporation of a quaternary hydroxyalkyl, specifically a hydroxymalonyl group at the C-3 position of IPs using air as the sole oxygen source. Keeping in mind the progress in photochemical relay catalysis [24] and the attention paid to photocatalytic carbon-bond functionalization in the past several years [25], here we developed an organophotoredox-catalyzed C–H functionalization of imidazo[1,2-a]pyridines and related heterocycles with active bromomethylenes under mild conditions (Scheme 1B). Importantly, using simple Zn(OAc)$_2$ as the additive, the first photocatalytic direct acetoxymalonylation of imidazo heterocycles was developed under aerobic conditions. Here, the additive Zn(OAc)$_2$ plays a crucial triple role as activator of IPs, halide scavenger, and acetylating agent.

**Results and Discussion**

**Optimization**

In the quest for the optimal reaction conditions, we started our investigations with 2-phenylimidazo[1,2-a]pyridine (1a) and diethyl bromomalonate (2a) as model substrates. Initially, the reaction was carried out between 1a and 2a in dry CH$_3$CN as
solvent under N₂ atmosphere using 4CzIPN as the photocatalyst. Irradiating the reaction mixture for 10 h under blue LEDs (450 nm) led to the isolation of products 5 (54%) and 6 (28%) (Table 1, entry 1). However, the same reaction, under aerobic conditions, delivered compounds 3a (47%) and 6 (22%) (Table 1, entry 2). Keeping in mind the ability of Zn(OAc)₂ as a bromide ion scavenger [26], we used Zn(OAc)₂ (2 equiv) as an additive to prevent the formation of the bromo product 6. While the additive successfully prevented the formation of compound 6, we were delighted to isolate the unexpected acetylated product 4a with a promising yield of 38% (Table 1, entry 3), reflecting the ability of Zn(OAc)₂ to act as an acetylating agent.

While screening other organophotocatalysts, we detected no desired product 4a (Table 1, entries 4–6) [27], except for photocatalyst 10-phenylphenothiazine (PTH) under violet LEDs which uplifted the yield up to 52% (Table 1, entry 7). Now with the optimal catalyst in hand, we screened some common solvents, out of which 1,2-DCE positively impacted the yield (Table 1, entries 8–11). However, the best result was obtained when 3.0 equiv of Zn(OAc)₂ was used as an additive (Table 1, entry 12). To check the viability of other acetylating agents, Zn(OAc)₂ was replaced with AcOH, generating the desired product in a comparatively lower yield (Table 1, entry 13). Finally, control experiments without a catalyst (Table 1, entry 14), light (entry 15) or acetylation agent (entry 16) failed to provide the desired product 4a, displaying the necessity of each component for developing the reaction.

### Substrate scope

With suitable reaction conditions (Table 1, entry 12), we systematically investigated the scope of this acetoxymalonylation strategy with substrate 2a (Scheme 2). Several imidazo[1,2-a]pyridines with diverse aryl substituents in the C-2 position were acetoxymalonylated, providing the desired products 4a–k regioselectively in good to excellent yields. Reflection of electronic properties was shown by the substituents attached to the aryl ring – as electron-releasing groups (Me, OMe) showed little more reactivity than electron-withdrawing groups (CN) at the same position (4b, 4f, and 4g). Halogen-substituted IPs also followed the general reactivity trend of the respective halogens (4c–e). Excellent reactivity was found for o-F and m-Br-substituted IPs (4h and 4i). Similarly, IPs associated with biphenyl

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**Table 1: Reaction optimization.**

<table>
<thead>
<tr>
<th>Entry</th>
<th>Catalyst</th>
<th>Solvent</th>
<th>Additive</th>
<th>Yield (%)&lt;sup&gt;b&lt;/sup&gt; 3a:4a:5:6</th>
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<td>1&lt;sup&gt;c&lt;/sup&gt;</td>
<td>4-CzIPN</td>
<td>CH₃CN</td>
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<tr>
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<tr>
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<td>Zn(OAc)₂</td>
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<tr>
<td>7&lt;sup&gt;d&lt;/sup&gt;</td>
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<td>8</td>
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<tr>
<td>14</td>
<td>–</td>
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<td>–</td>
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<tr>
<td>15&lt;sup&gt;f&lt;/sup&gt;</td>
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<td>–</td>
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<tr>
<td>16</td>
<td>–</td>
<td>1,2-DCE</td>
<td>–</td>
<td>57:0:0:24</td>
</tr>
</tbody>
</table>

<sup>a</sup>Reaction conditions: 1a (0.2 mmol), 2a (0.4 mmol), catalyst (5 mol %), additive (0.4 mmol) in dry solvent (2 mL) under aerobic conditions, irradiation with 12 W blue LEDs for 10 h. <sup>b</sup>Isolated yield. <sup>c</sup>Under N₂ atmosphere. <sup>d</sup>Irradiation with violet LEDs (λ<sub>max</sub> = 390 nm). <sup>e</sup>3.0 equiv of zinc acetate used. <sup>f</sup>In the dark, without light source.
and naphthyl groups in the C-2 position were also suitable substrates giving the corresponding products 4j and 4k in 77% and 82% yield, respectively. However, the yield of the products varied when different groups with diverse electronic properties were present in the pyridine ring of the IP moieties (4l–q). With substrates having a methyl substitution at C-7 and C-8 of the pyridine ring, the yields and regioselectivity were still excellent (4l and 4m), but reduced significantly upon introducing a halogen group onto the pyridine ring. Except for the 6-bromo-substituted compound (4o), all other substrates having a
halogen substituent in the pyridine ring showed reduced yields (4n, and 4p,q). The number of substituents also seemed to negatively affect the yield, as observed for products 4p and 4q, featuring two substituents each on the pyridine ring. Moreover, IPs with a non-aromatic C-2 substituent like an ester group were also included (4r). We also explored bromo analogues of other active methylenes such as ethyl cyanoacetate, ethyl acetoacetate, dimethyl, and diisopropyl malonates, as extension of diethyl malonate (4s,t and 4x,y). Lastly, we explored a few heterocycles that resemble imidazo[1,2-a]pyridine to vindicate the generality of this method. Gratifyingly, 6-phenylimidazo[2,1-b]thiazole, 2-phenylbenzo[d]imidazo[2,1-b]thiazole, and 2-phenylimidazo[1,2-a]pyrimidine participated well under the standard reaction conditions, generating the corresponding acetoxymalonolylated products 4u-w in good to excellent yields.

Several control experiments were performed to gain insights into the mechanistic pathway of this reaction. Firstly, a radical scavenging experiment using the radical scavenger TEMPO was performed (Scheme 3A). Upon analyzing the reaction mixture of 1a and 2a under standard conditions in the presence of TEMPO, we found only a trace of the desired product 4a. At the same time, a TEMPO-DEM adduct 7 and TEMPO-OAc adduct 8 were identified by the HRMS analysis of the crude reaction mixture, indicating the involvement of a malonyl radical and an acetyl radical in the course of the reaction (see Supporting Information File 1 for details). Additionally, when an aliphatic alkene, 5-hexen-1-ol was introduced into the reaction mixture under standard conditions without Zn(OAc)2, an ATRA product 9 was isolated, further confirming the involvement of a malonyl radical generated by the cleavage of the C–Br bond of 2a [28]. Next, an attempt was made to identify the key interme-

Scheme 3: Mechanistic investigations.
diate of the reaction (Scheme 3B). When compound 5 was subjected to the acetylation reaction individually with Zn(OAc)$_2$ and AcOH under optimized reaction conditions, the acetylated product 4a was produced with excellent conversion (>90%). These results suggest the involvement of compound 5 as an intermediate, and Zn(OAc)$_2$ or AcOH may be effective acetylating agents via generation of acetyl radicals. Control experiments under degassed conditions with or without water only delivered a trace amount (<5%) of the desired products, indicating that aerial oxygen plays a crucial role in the second catalytic cycle for the conversion of 5 to 3a or 4a (Scheme 3C). To determine the role of zinc acetate, a standard reaction of 1a and 2a in the absence of Zn(OAc)$_2$ was conducted (Scheme 3D). The results showed the formation of hydroxymalonated product 3a (57%) and bromo derivative 6 (24%). Notably, the hydroxymalonated product 3a under the reaction conditions was not converted to the acetylated derivative 4a, confirming 3a is not the intermediate for the final product 4a. So, Zn(OAc)$_2$ is crucial in shutting down the formation of 6 by scavenging free bromide in the reaction as ZnBr$_2$ salt (confirmed by HRMS). In addition, an excellent yield of the final product 4a [4a: 94% vs (6, 24% + 3a, 57%)] with additive indicates that zinc acetate plays a crucial role in activating IP towards the photoredox coupling reaction. Shifting of protons in the $^1$H NMR spectrum of 2-phenylimidazo[1,2-a]pyridine (1a) in the presence of Zn(OAc)$_2$ in CDCl$_3$ indicates a weak interaction of Zn(OAc)$_2$ with 1a (see Supporting Information File 1 for details) [20,21].

Finally, the reaction of 5 with benzoic acid and zinc acetate (in a 1:1 ratio) under standard reaction protocol resulted in the competitive formation of products 4a and 10 (Scheme 3E), indicating the susceptibility of other acids towards this method. These results, along with the Stern–Volmer fluorescence quenching study (Scheme 3F), expressed that the photoredox reaction started with the reductive generation of a malonyl radical from bromomalonate by interaction with the photocatalyst.

Analyzing all the observations from the above mechanistic studies, we propose a plausible mechanism involving sequential activation and functionalization of sp$^2$ and sp$^3$ C–H bonds via relay catalysis (Scheme 4). The relay can be divided into two cycles; the first cycle (cycle-1) deals with the C(sp$^2$)–H functionalization at the C-3 position of the imidazo heterocycles, while the second cycle (cycle-2) is all about the C(sp$^3$)–H functionalization at the newly incorporated active methylene center.

**Cycle-1** is initiated with the reduction of bromomalonate 2a by the photoexcited catalyst PC$^+$ to malonyl radical I. This is followed by the Minisci-type addition of radical I to the imidazo-pyridine, preactivated by Lewis acidic Zn(OAc)$_2$ [29]. PC$^+$ then oxidizes the resulting radical II to carbocation III which rearomatizes by losing a proton to generate the intermediate IV and closing the first catalytic cycle. Meanwhile, the bromide ions in the medium undergo anion exchange with the Zn(OAc)$_2$ to release free acetate ions, along with the conversion into ZnBr$_2$ (confirmed by HRMS). These in situ-generated free acetate ions function as a base, deprotonating carbocation III to produce the intermediate IV and AcOH.

The first step of cycle-2 involves the oxidation of the excited photocatalyst by aerial oxygen to generate superoxide anion and

---

**Scheme 4: Plausible reaction mechanism.**
PC^+. The superoxide anion (O_2^{-}) then captures the proton from the active methylene center of intermediate IV to generate the malonoyl radical V, which undergoes single electron oxidation by PC^-- generating the malonoyl radical VI [30,31]. Meanwhile, the hydroperoxy radical (\text{OAc})^-- forms, reacts with AcOH produced in cycle-1 to give the acetoxy radical (\text{OAc})^-- and H_2O_2. Then, the radical recombination between AcO^-- and radical VI furnishes the desired product 4. In the absence of the acetoxy radical (\text{OAc})^--, the hydroperoxy radical (\text{OAc})^-- may combine with radical VI to produce VII, which then easily converts into hydroxymalonated product 3 [31].

Conclusion
Thus, we have reported the successful C-3 acetoxymalonylation of imidazo[1,2-a]pyridines and related heterocycles by an organophotocatalytic relay C–H functionalization strategy with Zn(OAc)_2 in the triple role of an activator, bromide scavenger, and acetylating agent. The developed method is heavy-metal free, as shown by the use of inexpensive PTH, as well as a base-free approach, and involves aerial oxygen to generate exciting derivatives, which may prove to be valuable in the field of radical chemistry research in future.

Supporting Information

Supporting Information File 1
Experimental section and characterization of synthesized compounds.
[https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-19-48-S1.pdf]

Funding
Financial support from SERB (CRG/2021/004140), India, is gratefully acknowledged. D.S and S.P thank IIT(ISM) and CSIR, New Delhi, for doctoral fellowships, respectively.

ORCID® iDs
SoumitraMaity - https://orcid.org/0000-0003-0944-8162

References
With these catalytic systems (entries 4–6) no desired products (3a, 4a, 5, 6) were identified: instead a new product N-(pyridin-2-yl)benzamide was isolated in 8–14% yield. Jain et al. recently reported that aerobic oxygen under photoredox conditions oxidatively cleave the imidazoyl ring of 1a to benzamide derivatives.
See for details for photoredox ATRA reaction between diethyl bromomalonate 2a and 5-hexen-1-ol.


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Sulfate radical anion-induced benzylic oxidation of N-(arylsulfonyl)benzylamines to N-arylsulfonylimines

Joydev K. Laha*, Pankaj Gupta and Amitava Hazra

Abstract
A mild, operationally convenient, and practical method for the synthesis of synthetically useful N-arylsulfonylimines from N-(arylsulfonyl)benzylamines using K$_2$S$_2$O$_8$ in the presence of pyridine as a base is reported herein. In addition, a “one-pot” tandem synthesis of pharmaceutically relevant N-heterocycles by the reaction of N-arylsulfonylimines, generated in situ with ortho-substituted anilines is also reported. The key features of the protocol include the use of a green oxidant, a short reaction time (30 min), chromatography-free isolation, scalability, and economical, delivering N-arylsulfonylimines in excellent yields of up to 96%. While the oxidation of N-aryl(benzyl)amines to N-arylimines using K$_2$S$_2$O$_8$ is reported to be problematic, the oxidation of N-(arylsulfonyl)benzylamines to N-arylsulfonylimines using K$_2$S$_2$O$_8$ has been achieved for the first time. The dual role of the sulfate radical anion (SO$_4^{2-}$), including hydrogen atom abstraction (HAT) and single electron transfer (SET), is proposed to be involved in the plausible reaction mechanism.

Introduction
Among various imine compounds [1], N-arylsulfonylimines are perhaps the most prominent due to their unique stability, defined reactivity, and versatility in organic synthesis [2]. Leveraging their electron-deficient nature, N-arylsulfonylimines are widely used in organic transformations including nucleophilic addition, cycloaddition, imino-aldol reaction, ene reactions, aza-Friedel–Crafts reactions, and C–H functionalizations ([3] and references therein), leading to the synthesis of diverse nitrogen heterocycles of pharmaceutical relevance [4]. The traditional synthetic method for the preparation of N-arylsulfonylimines, similar to the preparation of N-arylimines, is based on the condensation of aromatic aldehydes and sulfonamides (Scheme 1a) [3,5-8]. Because of the poor nucleophilicity of sulfonamides, the condensation reactions generally require
harsh reaction conditions involving the use of strong acids, elevated temperature, and metal catalysts. Other methods include a non-dehydrative reaction of aldehydes with isocyanate analogs ([3] and references therein) (Scheme 1b) and an oxidative reaction of primary benzylic alcohols with sulfonamides or chloramine-T ([3] and references therein), and although they are elegant, they use substrates that are not readily accessible or toxic in nature. To overcome these limitations, oxidation of N-(arylsulfonyl)benzylamines to N-arylsulfonylimines, as opposed to the traditional methods, under mild and neutral reaction conditions has been reported, although limited to a few methods. However, these methods of oxidation involving the use of CrO₃ [9], Phl(OAc)₃/I₂ [10], TEMPO [11], NHPI [12], and metal catalysts [13], suffer from serious limitations including the use of metal catalysts, high temperature, risk of explosive hazards, production of large waste, and often low yield (Scheme 1c). Thus, an environmentally benign method that could deliver N-arylsulfonylimines under mild reaction conditions is highly desirable.

Previously, we reported a tandem oxidative intramolecular cyclization of N-aryl(benzyl)amines, having an internal nucleophile substituted at the ortho-position in the aniline ring, to nitrogen heterocycles using potassium persulfate (K₂S₂O₈) as the exclusive reagent [14]. The mechanistic study revealed that an initial oxidation to an iminium ion could be the key intermediate in the intramolecular cyclization step. In sharp contrast, when N-aryl(benzyl)amines that do not have an ortho-substituted nucleophile in aniline ring were used as the substrates in this reaction, N-arylimines were not isolated. Rather, an amide, in some cases, was isolated via oxidation of the benzylic methylene to a carbonyl group [14]. In the quest of a new method for
the synthesis of N-arylsulfonylimines, we questioned ourselves whether N-(arylsulfonyl)benzylamines would behave similarly as N-aryl(benzyl)amines under K₂S₂O₈-mediated oxidative conditions and could provide a platform for the synthesis of N-arylsulfonylimines.

To this endeavor, we have developed a method for the synthesis of N-arylsulfonylimines from N-(arylsulfonyl)benzylamines using K₂S₂O₈ in the presence of pyridine as a base. The key findings include a) requirement of a mild base for the formation N-arylsulfonylimines, and b) stability of N-arylsulfonylimines, unlike N-arylimines, under the oxidative conditions. Further, to demonstrate the scope and applicability of this approach, a gram-scale synthesis and a “one-pot” tandem synthesis of pharmaceutically relevant N-heterocycles by the reaction of in-situ-generated N-arylsulfonylimines with various ortho-substituted anilines were also developed. The mechanism of the oxidation is believed to occur via hydrogen atom abstraction (HAT) followed by single electron transfer (SET) enabled by the sulfate radical anion (SO₄²⁻).

### Results and Discussion

Initially, we investigated the reaction of N-benzene-sulfonyl(benzyl)amine (1a) as a model substrate with K₂S₂O₈ in MeCN at 80 °C for 12 h, conditions that were used earlier in our previous study [14]. Unfortunately, no product formation was observed under these conditions, while substrate 1a remained unreacted (Table 1, entry 1). When the solvent was changed to H₂O, a trace quantity of product formation was observed (Table 1, entry 2). To our surprise, when 2 equiv of pyridine were used as an additive along with the oxidant K₂S₂O₈ in MeCN, the desired product N-benzenesulfonylimine 2a was obtained in 90% yield (Table 1, entry 3). Subsequently, we carried out further optimization studies by changing the additive, solvent, temperature, and reaction time to obtain the best possible yield of the product 2a (Table 1). Interestingly, when duration of the reaction was reduced to 1 h, product 2a was obtained in 96% yield with complete conversion of substrate 1a (Table 1, entry 4). Further shortening the reaction time to 30 min resulted in the formation of 2a also in 96% yield (Table 1, entry 5). Lowering the temperature to 60 °C had a deleterious effect (Table 1, entry 6). Likewise, reducing the stoichiometry of pyridine to 1 equiv proved detrimental (Table 1, entry 7). Replacing pyridine with other organic and inorganic bases such as Et₃N, DBU, DABCO or K₂CO₃ also gave product 2a, however, in varying yields (Table 1, entries 8–11). While replacing the solvent MeCN with DCE delivered 2a in 89% yield, and a dramatic reduction in the yield of 2a was observed when H₂O was used as the solvent (Table 1, entries 12 and 13). Therefore, the conditions listed in entry 5 of Table 1 were chosen as the best conditions for further evaluating the substrate scope. Unlike the oxidation of N-aryl(benzyl)amines to N-arylimines

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**Table 1: Optimization of reaction conditions.**

<table>
<thead>
<tr>
<th>Entry</th>
<th>Additive (equiv)</th>
<th>Solvent</th>
<th>Temp. (°C)</th>
<th>Time (h)</th>
<th>Yield (%)&lt;sup&gt;b&lt;/sup&gt;</th>
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<td>90</td>
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<td>1</td>
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<td>80</td>
<td>1</td>
<td>20</td>
</tr>
</tbody>
</table>

<sup>a</sup> Reaction conditions: 1a (0.25 mmol), K₂S₂O₈ (0.5 mmol), additive (0.5 mmol) in solvent (1 mL) at 80 °C for the specified period of time. n.d. = not detected. <sup>b</sup> Isolated yield.
using K$_2$S$_2$O$_8$ in the presence or absence of a base is unsuccessful, the oxidation of N-arylsulfonylbenzylamines 1a to imines 2a was achieved under the optimized conditions. Distinctly, the use of a base is the key to success in this oxidation. Perhaps more importantly, the stability of N-benzenesulfonylimine 2a, unlike N-arylimines, under the oxidative conditions is noteworthy.

With the optimized reaction conditions in hand (Table 1, entry 5), we further investigated the substrate scope for the above transformation (Scheme 2). A limited variety of N-(arylsulfonyl)benzylamines 1a–m carrying substitutions on the aromatic rings was examined. Firstly, N-(arylsulfonyl)benzylamines having substitution(s) on one or both rings delivered the N-arylsulfonylimines 2a–h in 80–96% yield. The presence of a disubstitution in 1i gave product 2i in 78% yield. Replacing phenyl with naphthyl in N-(arylsulfonyl)benzylamines 1j and 1k resulted in the formation of N-arylsulfonylimines 2j and 2k also in very good yield (82–84%). Interestingly, when the arylsulfonyl group was replaced by methylsulfonyl, as in substrate 1l, the desired N-sulfonylimine 2l was obtained in 90% yield under the optimized reaction conditions. However, an attempted synthesis of N-arylsulfonylketimines was unsuccessful. Thus, N-(arylsulfonyl)benzylamine 1m having a phenyl substituent at the benzylic position gave benzophenone in 80% yield with a trace of N-benzenesulfonylketimine 2m under the optimized reaction conditions. Likewise, N-(arylsulfonyl)benzylamine 1n having a methyl group present at the benzylic position gave product 2n only in a trace quantity. To demonstrate further the scalability of the developed protocol, we carried out a gram-scale synthesis of 2a from 1a under the optimized reaction conditions. A complete conversion of substrate 1a was observed within 2 h under the optimized reaction conditions giving the product with an isolated yield of 92%.

Furthermore, to demonstrate the synthetic utility of the developed protocol, a tandem “one-pot” synthesis of N-heterocycles was successfully executed (Scheme 3). Thus, exposition of substrates 1 under the optimized reaction conditions followed by the addition of ortho-substituted anilines 3 and

**Scheme 2:** Substrate scope for the synthesis of N-arylsulfonylimines. Reaction conditions: 1a (0.25 mmol), K$_2$S$_2$O$_8$ (0.5 mmol), pyridine (0.5 mmol) in MeCN (1 mL) at 80 °C for 0.5 h. Yields refer to isolated compounds. aGram-scale synthesis (1a, 5 mmol).
Scheme 3: Tandem “one-pot” synthesis of N-heterocycles. Reaction conditions: 1a (0.25 mmol), K$_2$S$_2$O$_8$ (0.5 mmol), and pyridine (0.5 mmol) in MeCN (1 mL) at 80 °C for 0.5 h followed by the addition of 1 equiv of K$_2$S$_2$O$_8$ and the corresponding ortho-substituted anilines 3 (1.2 equiv) and stirring at 80 °C for 2 h. Yields correspond to isolated products.

Scheme 4: Control experiment with TEMPO. K$_2$S$_2$O$_8$ (1 equiv) and heating the reaction mixture at 80 °C for 2 h furnished the desired N-heterocycles 4. Thus, treatment of substrate 1a under the standard conditions, followed by reaction of the intermediate N-benzenesulfonylimine 2a with 2-aminobenzamide in one-pot gave 2-phenylquinazolin-4(3H)-one (4a) in 86% yield. Similarly, the reaction of the intermediate product 2c and 2-aminobenzamide gave 2-(p-tolyl)quinazolin-4(3H)-one (4b) in 85% yield.

Furthermore, when various other ortho-substituted aniline derivatives such as 2-aminobenzylamine, 2-aminothiophenol, and o-phenylenediamine are reacted with imine 2a in a similar manner, the corresponding N-heterocycles 4c–f were obtained in good to moderate yield. However, the reaction with 2-aminophenol did not give the corresponding cyclized product 4g. This could be possibly due to the poor nucleophilicity of the ortho-OH group in 2-aminophenol thereby restricting the intramolecular nucleophilic addition and as a result the corresponding cyclized product is not formed. The synthesis of these nitrogen heterocycles signifies the innate ability of in-situ-generated N-arylsulfonylimines in a variety of reactions with various ortho-substituted anilines without the need for pre-isolation or purification.

Next, in order to determine whether the reaction proceeds via a radical pathway, we performed a control experiment. When substrate 1a was treated with the radical scavenger TEMPO under the optimized reaction conditions, the formation of product 2a was completely suppressed (Scheme 4). This confirms that the reaction proceeds via a radical pathway.

Based on the literature [15,16], our previous experience [14,17,18], and current understanding, a plausible mechanism for the benzylic oxidation is depicted in Scheme 5. Initially, a sulfate radical anion (SO$_4^{-}$) is generated by homolytic cleavage of the peroxy linkage under heating conditions [17]. The hydrogen atom is abstracted from the benzylic position of 1 by SO$_4^{-}$, generating benzylic radical 1aa [14-16]. A single electron transfer (SET) could subsequently occur from 1aa to form the reactive species 1ab. Finally, the base abstracts the activated...
NH proton to produce imine 2. The dual role of SO$_4^{2-}$ involving HAT and SET is proposed in this plausible mechanism, which requires further investigation.

Similarly, a plausible mechanism for the one-pot synthesis of $N$-heterocycles is shown in Scheme 6. Initially, the $N$-arylsulfonylimine 2, generated in situ from the corresponding $N$-(arylsulfonyl)benzylamine 1, undergoes transimination with the ortho-substituted aniline 3 to form imine 3ab via 3aa. Subsequent intramolecular nucleophilic addition in imine 3ab produces intermediate 3ac, which upon oxidation delivers the desired $N$-heterocycle 4.

**Conclusion**

In conclusion, we have developed a complementary approach to the currently available methods for the oxidation of $N$-(arylsulfonyl)benzylamines to $N$-arylsulfonylimines using K$_2$S$_2$O$_8$ and pyridine as a base. While $N$-arylimines are difficult to prepare by the oxidation of $N$-aryl(benzyl)amines using K$_2$S$_2$O$_8$, $N$-arylsulfonylimines are successfully prepared and are quite stable under the oxidative conditions. In addition, we demonstrated a “one-pot” tandem synthesis of pharmaceutically relevant $N$-heterocycles through the reaction of in situ-generated $N$-arylsulfonylimines with ortho-substituted anilines. The key features including the use of a green oxidant, a short reaction time, chromatography-free isolation, and scalability mark a distinction from the contemporary methods. Although we propose a dual role for SO$_4^{2-}$ involving both hydrogen atom abstraction (HAT) and single electron transfer (SET), further investigation of the mechanism would enrich our understanding of persulfate-mediated oxidative reactions.

**Supporting Information**

**Supporting Information File 1**

General procedures, product characterization, and copies of $^1$H NMR and $^{13}$C NMR spectra of all compounds. [https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-19-57-S1.pdf]

**Acknowledgements**

The authors acknowledge NIPER S.A.S. Nagar for providing excellent research facilities.

**Funding**

We greatly appreciate the generous financial support from the Science & Engineering Research Board (SERB) of DST, New Delhi (award number CRG/2020/000462). PG thanks the DST INSPIRE (IF180484) for a research fellowship.
ORCID® iDs
Joydev K. Laha - https://orcid.org/0000-0003-0481-5891
Pankaj Gupta - https://orcid.org/0000-0002-0004-077X
Amitava Hazra - https://orcid.org/0000-0003-1129-7025

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The definitive version of this article is the electronic one which can be found at: https://doi.org/10.3762/bjoc.19.57
Pyridine C(sp²)–H bond functionalization under transition-metal and rare earth metal catalysis

Haritha Sindhe¹, Malladi Mounika Reddy², Karthikeyan Rajkumar², Akshay Kamble¹, Amardeep Singh², Anand Kumar² and Satyasheel Sharma *²

Review

Address:
¹Department of Medicinal Chemistry, National Institute of Pharmaceutical Education and Research - Ahmedabad, Gandhinagar, Gujarat, 382355, India and ²Department of Natural Products, National Institute of Pharmaceutical Education and Research - Ahmedabad, Gandhinagar, Gujarat, 382355, India

Email:
Satyasheel Sharma* - sharma.satyasheel@gmail.com

* Corresponding author

Keywords:
C–H functionalization; heterocycles; pyridine; rare earth metal; transition-metal-catalyzed

Abstract

Pyridine is a crucial heterocyclic scaffold that is widely found in organic chemistry, medicines, natural products, and functional materials. In spite of the discovery of several methods for the synthesis of functionalized pyridines or their integration into an organic molecule, new methodologies for the direct functionalization of pyridine scaffolds have been developed during the past two decades. In addition, transition-metal-catalyzed C–H functionalization and rare earth metal-catalyzed reactions have flourished over the past two decades in the development of functionalized organic molecules of concern. In this review, we discuss recent achievements in the transition-metal and rare earth metal-catalyzed C–H bond functionalization of pyridine and look into the mechanisms involved.

Introduction

Pyridine, one of the most important azaheterocyclic scaffolds, is found in a diverse range of bioactive natural products, pharmaceuticals, and functional materials [1-10]. Due to its different characteristics such as basicity, stability, water solubility, small molecular size, and ability to form hydrogen bonds, pyridine continues to be a suitable moiety in organic synthesis. In addition, it has been observed that pyridine rings serve as bioisostere for aromatic rings, amines, amides, and N-containing heterocycles. Due to the aforementioned qualities, numerous U.S. FDA-approved medications have pyridine scaffolds in their molecules (Figure 1).

In this context, the synthesis of the pyridine motif is always there in trend. Many pyridine syntheses have relied on the con-
densation of carbonyl compounds and amines for a very long time [11]. The classical methods for the synthesis of functionalized pyridine include the Hantzsch pyridine synthesis and the Bohlmann–Rahtz synthesis (Scheme 1a and b). Furthermore, alternative methodologies are being developed for the synthesis of functionalized pyridines or its integration into an organic molecule [12-20]. Although classical organic synthesis is incredibly effective, it frequently requires the prefunctionalization of substrates and involves stoichiometric waste.

The challenges associated with the functionalization of pyridine are based on the low reactivity of the pyridine ring system for undergoing substitution reactions. This is attributed to the electron-deficient nature of the ring system due to the presence of the sp²-hybridized nitrogen atom. In addition, the lone pair electrons of the nitrogen atom interact with Lewis acids instead of the π-electrons of the ring system thus resulting to its reduced reactivity for electrophilic aromatic substitution reactions, such as a Friedel–Crafts reaction [21-23]. Hence, it is challenging to functionalize a C–H bond in pyridine with traditional chemical transformations. On the other hand, intriguing developments have been made for the functionalization of inert C–H bonds in organic synthesis during the past two decades. In this regard, the transition-metal-catalyzed C–H functionalization has made its way towards the synthesis and functionalization of various complex organic molecules [24-31]. In addition, rare earth metal-catalyzed C–H functionalization reactions have been known for a few decades, however, they received growing interest only recently [32-34]. Thus, diversely functionalized pyridines have been synthesized via C–H activation under transition-metal and rare earth metal catalysis, including C–H alkylation, alkenylation, arylation, heteroarylation, borylation, etc.
Recently, metal-free approaches have also been developed for the C–H functionalization of N-heterocycles [35-39]. However, due to the vastness of reports on C–H functionalizations of N-heterocycles, in this review we have summarized recent progress (from year 2010 to 2023) in the C–H functionalization of the pyridine ring only. Herein, we discuss transition-metal as well as rare earth metal-catalyzed directed and undirected, proximal as well as distal pyridine C(sp²)–H bond functionalizations in detail under different types of reactions. Further, this review excludes the use of pyridine as a directing group for C–H functionalizations and the C–H functionalization of fused pyridines.

**C–H Alkylation of pyridine**

The C–H bond is the backbone of an organic molecule and the conversion of a C–H bond to a C–X bond (X = carbon or heteroatom) forms the basis in organic synthesis. The functionalization of C–H bonds is challenging due to a large kinetic barrier for C–H bond cleavage and also achieving selectivity is difficult due to its ubiquitous nature [40]. The metal-catalyzed C–H bond functionalization is a good strategy for synthesizing highly functionalized organic frameworks. In this context, the C–H alkylation is one of the most important C–C bond formation reactions [41-45]. On the other hand, a metal-catalyzed functionalization of arene/heteroarene C–H bonds to the corresponding C–C bonds is an area of great interest and has been well studied [46,47]. Pyridine, being an important heterocyclic scaffold, various studies have been conducted for the C(sp²)–H alkylation of the pyridine ring. In this part, we describe pyridine C–H alkylation reactions sub-sectioning based on the position of the alkylation reported.

**ortho-C–H Alkylation**

Inspired by the pioneering work of Jordan and co-workers [48] on the ortho-selective C–H alkylation of 2-picoline with propene using a cationic zirconium complex under a H₂ atmosphere in 1989 and the work done by Bergman and Ellmann [49] in 2010 for the ortho-C–H alkylation of pyridines under Rh(I) catalysis at high temperature, in 2011 Hou and Guan reported an atom economical method for the selective ortho-alkylation of pyridines by C–H...
addition to olefins under cationic half-sandwich rare-earth catalysis [50]. They carried out the reaction in the presence of dialkyl complexes of scandium (Sc) or yttrium (Y) such as \((\text{C}_5\text{Me}_5)\text{Ln}((\text{C}_2\text{H}_5)_2\text{NMe}_2)_2\) \((\text{Ln} = \text{Sc}, \text{Y})\) in combination with \(\text{B}((\text{C}_6\text{F}_5)_3)\) as an activator. The method demonstrated a wide substrate scope of both pyridines and olefins including \(\alpha\)-olefins, styrenes, and conjugated dienes. The yttrium complex was found to be superior as compared to the scandium complex for the alkylation reaction of bulkier 2-tert-butylpyridine with ethylene. In addition, the yttrium catalyst was also found to have a higher catalytic activity for the ortho-alkylation of pyridines with styrenes to give the linear alkylation products [5b,c, Scheme 2]. Further, the authors proposed that the C–H bond activation could be the rate limiting step based on kinetic isotope experiments (KIE). The proposed mechanism involves the coordination of pyridine to the metal center of the cationic catalyst and \(\text{B}((\text{C}_6\text{F}_5)_3)\) promotes the ortho-C–H activation (deprotonation) of pyridine to afford pyridyl species 6. Next, the 2,1-migratory insertion of alkene 2 into the metal–pyridyl bond in 6 gives the intermediate 7, which on subsequent deprotonation gives the branched alkylation product 4. Whereas, in case of styrene 3 a 1,2-insertion takes place possibly due to the formation of the stable benzallylic species 8, which on deprotonation gives the linear alkylation product 5.

The C–H activation/C–C cross-coupling reaction with 1° alkyl electrophiles has been known in the past, however, the C–H alkylation with nonactivated secondary (2°) alkyl electrophiles and tertiary alkyl electrophiles was little known. In this context, in 2013, Fu and co-workers came across an unexpected finding with Pd-catalyzed C–H activation/C–C cross-coupling of pyridine N-oxides with nonactivated secondary (2°) alkyl bromides [51]. The cross-coupling is difficult to achieve as the Pd-catalyzed S_n2 process is sensitive towards the steric bulk of the secondary or tertiary alkyl electrophiles. The optimized conditions for the ortho-alkylation of pyridine N-oxides 9 with nonactivat-

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**Scheme 2:** Rare earth (Ln)-catalyzed pyridine C–H alkylation.
ed secondary (2°) alkyl bromides 10 required 5 mol % of the Pd(OAc)₂-dppf catalyst, Cs₂CO₃ (2.0 equiv) as base in toluene at 100 °C as shown in Scheme 3. Under these conditions, the reaction provided diverse 2-alkylpyridine derivatives 11 in moderate to good yields starting from both cyclic and acyclic alkyl bromides. The findings of the reaction’s stereoreactivity and observations made during some cyclization or ring-opening reactions indicated that the C–H alkylation may proceed through a radical-type mechanism. Next, in 2013, Wang and co-workers [52] reported a protocol using CuI (10 mol %) as inexpensive catalyst and LiO-t-Bu (3.5 equiv) as the base for the C–H alkylation of N-iminopyridinium ylides 12 with N-tosylhydrazones 13 showing good substrate scope for both coupling partners (Scheme 4). A substituent on the aromatic ring of the tosylhydrazones did not significantly affect the C–H alkylation reaction and the reaction also proceeded well with hydrazones 13 obtained from aliphatic aldehydes or ketones. Based on mechanistic experiments and DFT calculations, the reaction presumably proceeds via a Cu–carbene migratory insertion (Scheme 4b). In the presence of CuI and the base the initial direct C–H activation of the ylide 12 gives the copper pyridinium ylide 15. The latter reacts with the diazo compound formed through reaction of hydrazone 13 with the base to give the copper–carbene species 16. Then, the intermediate 16

![Scheme 3: Pd-catalyzed C–H alkylation of pyridine N-oxide.](image)

![Scheme 4: Cul-catalyzed C–H alkylation of N-iminopyridinium ylides with tosylhydrazones (A) and a plausible reaction mechanism (B).](image)
undergoes a Cu–carbene migratory insertion giving intermediate 16', which upon protonation gives the desired alkylated product 14.

Later, in the year 2018, Yao and co-workers [53] developed the first example of a group 4 metal-based catalyst protocol for the C–H alkylation of pyridine 1 with alkenes 18 and 20 as coupling partners. They demonstrated that the reaction in the presence of cationic zirconium complexes derived from zirconium dibenzyl complexes bearing tridentate [ONO]-type amine-bridged bis(phenolato) ligands and [Ph₃C][B(C₆F₅)₄] (Scheme 5), gave rise to ortho-selective C–H alkylated pyridines 19 and 21. It was observed that the cationic Zr complexes provided good transformations, probably due to good
accessibility of the coordination site and an increased Lewis acidity of the metal center. The authors also demonstrated that this catalytic system also catalyzes the alklylation of benzylic C–H bonds (C(sp^3)–H) of various dialkylpyridines with alkenes. It is to be noted that the ligands’ backbones were found to be crucial for the regioselectivity of the addition to benzylic C(sp^3)–H bonds, as N-arylamine-bridged bis(phenolato) Zr complexes provided branched products whereas N-alkylamine-bridged bis(phenolato) Zr complexes provided the linear addition products. The proposed mechanism (Scheme 5b) involves the initial formation of Zr complex 22 through the reaction of neutral Zr complex 17 with [Ph_3C][B(C_6F_5)_4], which on coordination with the pyridine resulted in the formation of the 3-membered zirconacyclic intermediate 23. The migratory insertion of the alkene into the metal–C bond of 23 gives the intermediate 24a on reaction with styrene 18 and intermediate 24b in the presence of alkene 20. The intermediates 24a and 24b then undergo further hydrolysis to give the desired linear products 19 and branched products 21, respectively.

In the same year, Tsurugi and Mashima reported the use of rare earth metal complexes for the insertion of nonpolar unsaturated substrates (C=N) into the ortho-C–H bond of pyridine derivatives [54]. They carried out the C–H aminoalkylation of pyridines 1 using yttrium complex 26 with nonactivated imines 25 (Scheme 6). The authors also demonstrated the enantioselective aminoalkylation, using chiral diamines as ligands. The introduction of chiral diamines in the metal complex produced the aminoalkylated products enantioselectively with good ratio of enantiomeric excess. The plausible mechanism involves the formation of (dibenzylamido)yttrium complex 28 by the reaction of yttrium complex 26 with HNBr_2. Then σ-bond metathesis between the Y–N bond of 28 and the ortho-C–H bond of pyridine gives η_2-pyridyl species 29 which on imine insertion produces species 30. Subsequent protonation then provides the aminoalkylated product 27 (Scheme 6b).

The selective C–H monoalkylation of pyridines with alkenes is a challenging task. Most ortho-C–H alkylation reactions have been achieved starting from C2-substituted pyridines. There are a few studies reported for the selective C–H monoalkylation of unsubstituted pyridines, which, however, displayed limited substrate scope [55,56]. In this regard, in 2021, Nakao and co-workers [57] reported a selective C2-monoalkylation of 2,6-unsubstituted pyridines with alkenes 31 using a heterobimetallic Rh–Al catalyst. The reaction provided the linear product 32 with aliphatic alkenes 31, whereas vinylarenes produced the branched product 33 and also alkenylated products 34. The reaction gave excellent yields of the ortho-alkylated products with good functional group tolerance (Scheme 7).

The C–H functionalization of pyridines through action of different catalyst systems including transition metals and rare earth metals has been described and some other organometallic
systems also were shown to have catalytic reactivity. Adopting this catalytic reactivity of organometallics and also the special bidentate nature of phosphinoamide ligands, in 2021, Chen and group [58] described the catalytic ortho-C(sp²)–H functionalization of pyridines with polar imines 35 and nonpolar alkenes 37 by using mono(phosphinoamido)-ligated rare earth complexes (NP2-Gd and NP1-Sc) as shown in Scheme 8. Complex NP2-Gd was found to be effective in the functionalization of pyridines with imines providing various ortho-aminoalkylated products 36 whereas ortho-alkylated pyridine derivatives 38 were obtained when using NP1-Sc as the catalyst (Scheme 8). Attributing to the strong coordination of unsubstituted pyridine with Rh(I) catalysts, C–H alkylations of pyridine lacking ortho-blocking groups is a challenge. In this context, a regioselective alkylation of ortho-unsubstituted or substituted unactivated pyridines with acrylates and acrylamides under Rh(I) catalysis has been demonstrated by Ellman and co-workers [59]. The...
authors observed that in the presence of [Rh(cod)Cl]₂ as catalyst, dppe as ligand, and potassium pivalate (KOPiv) as base, linear C–H-alkylated products 40 were obtained from both acrylates and acrylamides in moderate to high yields (Scheme 9, reaction conditions a). However, when K₃PO₄ was employed as the base under otherwise identical conditions, the authors observed a switch in regioselectivity and branched products 41 were obtained with acrylamides as coupling partners (Scheme 9, reaction conditions b). Thus, the authors demonstrated a switch in regioselectivity (linear/branched) which was controlled exclusively by the base used. During further investigations the authors found that the use of ligand dArFpe at reduced reaction temperature resulted in a significant increase in the yield of the branched alkylated product 41 (Scheme 9, reaction conditions c) compared to using the ligand dppe (Scheme 9, reaction conditions b). Moreover, when ethyl methacrylate was used as the coupling partner under the reaction conditions c, branched alkylated products 41’ were obtained selectively in high yields (Scheme 9). A high functional group tolerance was observed in both linear and branched alkylated products.

It is known that the strong coordination of the nitrogen atom in pyridine rings with metals inhibits the metal–chiral ligand coordination, thus making the C–H alkylation of pyridine substrates challenging. In addition, transition-metal-catalyzed enantioselective C–H alkylation reactions of pyridine still remain a great challenge. In this regard, in 2022, Ye and co-workers [60] reported for the first time an enantioselective C-2 alkylation of pyridine using a chiral phosphine oxide-ligated Ni–Al bimetallic catalyst system and the protocol was found effective for a wide range of pyridines including unsubstituted pyridines,

**Scheme 9:** Rhodium-catalyzed pyridine C–H alkylation with acrylates and acrylamides.
C2, C3 and C4-substituted pyridines and complex pyridines containing bioactive molecules (Scheme 10). To attain enantioselectivity a chiral phosphine oxide (43)-ligated Ni–Al bimetallic catalyst was used that was critical in improving the reactivity and controlling the selectivity of the reaction. Further, based on deuterium labelling experiments, KIE studies, and DFT calculation, a plausible mechanism (Scheme 10b) has been proposed. Initially, a reversible ligand-to-ligand H-transfer process occurs for C–H activation between the intermediates 46 and 47. Next, isomerization of the η1-allyl complex 47 forms the η3-allylic nickel complex 48, which on reductive elimination delivers the desired product 44 via the intermediate 49 (Scheme 10b). It was proposed that the enantioselectivity was mainly due to the C–C reductive elimination of the R-pathway, which is lower in energy than the S-pathway.

Remote C–H alkylation
Several remarkable studies have been reported for proximal C–H functionalizations in pyridine substrates under different catalytic systems. However, the intermolecular undirected distal C–H functionalization in pyridine remained unstudied. In these circumstances, the distal C–H alkylation by addition of the pyri-
dine C–H bond to an aldehyde 50 under iridium catalysis was achieved by Shi [61] in 2010 through an unusual meta-selectivity for the first time (Scheme 11a). To achieve meta-selectivity, the group has screened various transition metals and revealed that a silyl-iridium complex promoted the addition of meta-pyridyl C–H bonds to aldehydes 50 which resulted in C3-alkylated pyridines 51. Based on the reactions performed for the catalytic activity of the silyl-iridium complex, the authors proposed a catalytic mechanism (Scheme 11b). The mechanism involves the initial formation of the active silyl-iridium catalyst A which through oxidative addition of 1 gives the silyl-iridium complex 52. The insertion of aldehyde 50 into the Ir–Si bond of 52 provides the pyridyl alkyl iridium species 53 that finally by C–C formation via reductive elimination furnishes the desired products 51 along with the formation of an iridium hydride species (Scheme 11b).

A direct selective C4-alkylation of pyridine has been reported by the groups of Hiyama [62] (Scheme 12a) and Zhang [63] (Scheme 12c) in 2010 and 2020, respectively. The Hiyama group developed a C-4-selective alkylation of pyridines using a Ni/Lewis acid cooperative catalytic system in combination with a bulky N-heterocyclic carbene ligand and (2,6-t-Bu₂-4-Me-C₆H₃)₂AlMe (MAD) as the Lewis acid which allowed the direct C-4 alkylation of pyridines 1 (Scheme 12a). With the optimized reaction conditions in hand the group also screened the alkene and pyridine substrate scope which resulted C4-alkylated products 55 in moderate to high yields. A possible mechanistic cycle (Scheme 12b) was also proposed, comprising an initial formation of η²-arenenickel species 56A, which undergoes oxidative addition to the C(4)–H bond of pyridine to form intermediate 56B. Next, coordination and migratory insertion of the alkene provides the intermediate 57 which on subsequent re-

\[
\begin{align*}
&\text{A)} \\
&\text{1 (1 equiv)} + \text{ArCHO (50 (3 equiv))} \\
&\text{Ir} \cdot \text{(CO)}_{\text{12}} \text{ (2 mol %)} \text{1,10-phenanthroline (4 mol %)} \\
&\text{HSiEt}_3 \text{ (3 equiv)} \text{PhH, 135 °C, 12 h} \\
&\text{51} \\
&\text{51a, 73%} \quad \text{51b, 53%} \quad \text{51c, 55%} \quad \text{51d, 76%}
\end{align*}
\]

\[
\text{B) proposed mechanism}
\]

\[
\begin{align*}
&\text{Ir} \cdot \text{(CO)}_{\text{12}} / \text{Phen} \\
&\text{HSiEt}_3 \\
&\text{H}_2 \\
&\text{L}_n \text{Ir-SiEt}_3 \\
&\text{A} \quad \text{oxidative addition} \\
&\text{HSiEt}_3 \\
&\text{L}_n \text{Ir-H} \quad \text{reductive elimination} \\
&\text{51} \\
&\text{52} \\
&\text{53} \\
&L = \text{Phen, CO, and/or SiEt}_3
\end{align*}
\]

\text{Scheme 11: Iridium-catalyzed pyridine C–H alkylation.}
ductive elimination furnishes the C4-alkylated products 55. Based on the deuterium exchange experiment, the author suggested that the steps involved in the catalytic cycle from 56A to 57 are reversible in nature, which may activate the C2 or C3 position as well. However, the reductive elimination at the C4-position was suggested to be irreversible in nature and does not take place at the C2 and C3 position. On the other hand, the Zhang group reported the C4 alkylation of pyridines using alkenes 58 catalyzed by an organoborohydride (NaBEt₃H) and aided by organoboranes (Scheme 12c). The proposed mechanism (Scheme 12d) involves the formation of the organoborate intermediate 60 from alken 58 in the presence of the NaBEt₃H catalyst and the organoborane. Next, the organoborane-activated pyridine species 61 undergoes an addition reaction regio-
selectively at the C4 position of the organoborate intermediate 60 delivering the σH-adduct intermediates 62 and 63. Subsequently, hydride elimination with the help of the organoborane gave the desired alkylated product 59 and regenerates the hydride catalyst.

Further enantioselective pyridine C–H alkylation reactions are very scarcely reported which specifically include the intramolecular C–H alkylation of pyridine with alkenes at the C3 or C4 positions. Hence, very recently in 2022, Shi and co-workers [64] adopted an intermolecular process and reported the enantioselective para-alkylation of pyridines with styrenes 64 using a Ni–Al bimetallic system and NHC ligand 65 through intermolecular hydroarylation with high levels of enantio- and regioselectivity in the alkylated products 66 (Scheme 13). Also, the authors performed DFT studies revealing the reaction mechanism and supported that the interaction of the NHC aryl part with trans-styrene was highly important for the reaction to proceed and for the enantiocontrolled formation of the products.

Alkenylation

The C–H alkenylation is another important C–C bond-forming reaction. Olefinated organic molecules like vinylarenes play a significant role as key intermediates in organic synthesis and are also present in various natural products as well as drug molecules [65–68]. Though there are traditional methods available for C–H olefinations they suffer from some disadvantages such as for example requiring prefunctionalized substrates as in case of the Heck cross-coupling [69,70]. However, researchers have developed various methods for the transition-metal-catalyzed C(sp²)–H olefination using various types of alkenes as coupling partners [71–73]. This part of the review covers reports for the alkenylation of pyridine with terminal alkenes, acrylates, allenes, and alkynes as coupling partners achieving the functionalized C(sp²)–H-olefinated pyridine frameworks via metal catalysis.

**ortho–C–H Alkenylation**

In 2012, Huang and co-workers [74] disclosed a ligand-free oxidative cross-coupling reaction of pyridine with acrylates, acrylamides, and styrenes (Scheme 14). Their preliminary investigation provided both C2 and C3-olefinated products, with the C2-selective product 69 as the major product (Scheme 14a). With the optimized conditions of Pd(OAc)₂ (10 mol %), AgOAc (3 equiv), PivOH (2.5 equiv) in DMF, the method showed wide substrate scope and good yields. Based on the experimental findings the authors proposed a catalytic cycle (Scheme 14b) which commences with the coordination of Pd(II) with the pyridine nitrogen to provide intermediate 70. A strong trans-effect results in the C–H cleavage for the formation of Pd(II) species 71. Subsequently, insertion of alkene 68 provides the cyclic Pd(II) intermediate 72 which undergoes β-hydride elimination to produce the desired product 69.

In the same year, Ramana and Goriya [75] proposed an unexpected C-6 (C-2)-propenylation reaction of pyridine in the presence of allyl bromide (73) and a Ru catalyst using 2-arylpyridines (Scheme 15). Earlier reports described the

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**Scheme 13**: Enantioselective pyridine C–H alkylation.
propenylation took place on the ortho-position of the phenyl ring [76, 77], whereas this group achieved the propenylation of the pyridine moiety. The authors screened different allyl halides and Ru complexes as catalysts. With the optimized conditions in hand, diverse 2-arylated pyridines were screened resulting in the corresponding products 74 in good yields.
Allene, a cumulated diene and an important building block in organic synthesis has versatile biological properties and is also an important subunit in various natural products and pharmaceutical compounds [78]. Allenes have been applied as useful substrates for the alkenylation of organic molecules [79]. There are various reports for the C–H alkenylation of aromatic C–H bonds using allenes [80]. To this end, Hou and group in 2015 [81] demonstrated the C–H allenylation of pyridines with excellent substrate scope using a scandium catalyst (Scheme 16). A vast number of pyridines and allenes were studied as substrates to provide the C2-alkenylated pyridines in good to high yields. Based on the mechanistic experiments a possible catalytic cycle has been proposed (Scheme 16b). The half-sandwich scandium complex 76 along with the

Scheme 16: C–H addition of allenes to pyridines catalyzed by half-sandwich Sc metal complex.
tetrakis(pentafluorophenyl)borate and pyridine forms a cationic Sc-pyridyl complex 78, which after addition of allene 75, forms a transient pentacyclic intermediate 80 via intermediate 79. Next, another molecule of pyridine adds to intermediate 80 to furnish the transient complex 81 which undergoes σ-bond metathesis to give the product 77 and regenerating 78 (Scheme 16b).

While speaking regarding the alkenylation, the geometrical isomerism, i.e., the stereoselectivity between the cis- and trans-alkenylation, has not been considered so far. Except lately, in 2020, Chen and group [82] reported a Pd/Cu-catalyzed regio- and stereoselective synthesis of C2-alkenylation pyridines starting from internal alkynes 84 and pyridinium salts in a stereodivergent manner (Scheme 17a). The interesting part of this work was the switching of the alkene configuration of the products by modifying the substituents on the nitrogen of the pyridinium salts. Further, the method showed a wide substrate scope for both the Z- and E-alkenylated products in which Z-selectivity was achieved when N-methylpyridinium salts were used and E-selectivity when N-benzylpyridinium salts were used.

In the proposed mechanism (Scheme 17b) the E- and Z-isomers can be assessed through point at which dealkylation occurs, i.e., if it occurs as last step the Z-isomer 85 is obtained and if it

\[ \text{Scheme 17: Pd-catalyzed stereodivergent synthesis of alkenylated pyridines.} \]
Remote alkenylation

In 2011, a study for weakening the strong coordination of the pyridyl N-atom with Pd in the presence of a bidentate ligand was reported by Yu and co-workers [83]. They showcased the C3-selective olefination of pyridines using 1,10-phenanthroline, a bis-dentate ligand that weakens the coordination of the Pd catalyst with the pyridyl N-atom through the trans-effect (Scheme 18). The trans-effect is the switching of the metal coordination between the π-ring system and the hetero (nitrogen) atom of pyridine [84,85]. In comparison to coordination with nitrogen, which is strong in nature, the coordination with the ring is weaker and cleavable. The usage of a bidentate ligand will enhance the trans-effect and shift the coordination towards the ring (Scheme 18b).

There are numerous studies reported for directing group or chelate-assisted metal-catalyzed C–H functionalization reactions. However, non-chelate-assisted or undirected C–H functionalizations under ligand-controlled conditions are underexplored. Hence, differentiating from this co-coordinative model, in 2013, Zeng and co-workers [86] reported the MPAA (mono-N-protected amino acids) ligand-promoted non-chelate-assisted C–H activation via Pd-catalyzed dehydrogenative Heck reactions on pyridines with simple alkenes 96, leading to the C3-alkenylated products 97 (Scheme 19). The reaction was based on the previous reports of using of the MPAA ligands in the Pd-catalyzed oxidative cross-coupling reactions discovered by Yu et al. [87]. When 2-methoxypyridine was screened, the reaction resulted in a mixture of C3- and C5-selective C–H-functionalized products 97f and 97f' in a regioisomeric ratio of nearly 1:1. Further, during the substrate scope study, when 1,1'-disubstituted butyl methacrylate was used as coupling partner a mixture of 97g and the isomeric product 97g' was observed in 42% yield.

Further, Shi and co-workers reported the rhodium-catalyzed directed C–H olefination of pyridines using different directing groups in 2013 [88] (Scheme 20a) and 2014 [89] (Scheme 20c), respectively. In the former study, under optimized conditions of...
Scheme 19: Mono-N-protected amino acids in Pd-catalyzed C3-alkenylation of pyridines.

Scheme 20: Amide-directed and rhodium-catalyzed C3-alkenylation of pyridines.
[RhCp*Cl₂]₂ (5 mol %), AgSbF₆ (20 mol %) in DCE at 120 °C, Cu(OAc)₂ was found crucial for the transformation in comparison to other additives and showed good substrate scope while unactivated alkenes like styrene resulted in no reaction. Also, the authors successfully applied the developed protocol to a multigram-scale synthesis of compound 101, a tricyclic imidazophenaphthyridine derivative having antibacterial properties, with low catalyst loading (0.1 mol %) (Scheme 20b). Later, in 2014, the same authors, using an amide as directing group (DG), developed a protocol for the regioselective C3-alkenylation of pyridines through syn-addition of alkynes, displaying broad substrate scope and high yields (Scheme 20c). Based on literature reports and experimental studies, a possible mechanism (Scheme 20d) was proposed in which coordination of the DG 102 to the rhodium cationic species followed by ortho-metalation and migratory insertion of 103 into the Rh–C bond of 105 provides a seven-membered rhodacyclic intermediate 106. The protonation at the Rh–C bond of intermediate 106 in the presence of RCOOH furnishes hydroarylation product 104.

Nitrogen heterocyclic carbenes (NHCs) are of central importance in organometallic chemistry and in organic synthesis. Also, metal–NHC complexes have wide application in catalysis and various organic transformations and a range of metal–NHCs served as catalysts. In 2010, using NHC ligands, Yap and co-workers [90] developed a method for the direct para and meta-C–H alkenylation of pyridines with 4-octyne (107) using a nickel Lewis acid catalyst with amino pendant linked NHC complex (Scheme 21). In addition, the authors were able to isolate the bimetallic intermediate structure η²,η¹-pyridine–Ni(0)–Al(III) complex 112, as a support for their mechanism for the para-C–H functionalization. They further investigated the scope and limitations of the dual catalyst Ni–AlMe₃ and also the sensitivity of the reaction towards the steric environment on the pyridine ring. The complex 112 undergoes oxidative addition followed by an alkyne insertion reaction to give intermediate 113, which after reductive elimination provides the alkenylated product 109 (Scheme 21b).

Arylation
C-2 Arylation
Owing to the remarkable role of aromatic C–H arylation reactions in organic synthesis abundant methods have been reported for aromatic C–H arylations using different arylating coupling partners, such as for instance, aryl halides. In 2014,
using organoboronic coupling partners, Wu and co-workers [91] reported a protocol for the Cu-catalyzed C–H arylation of pyridine N-oxides 9 with arylboronic esters 114 and prepared C2-arylated pyridines 115 in moderate to good yields (Scheme 22). By using an inexpensive Cu catalyst, the method allows for the simple and practical synthesis of 2-arylpyridines. The reaction starts with the formation of arylated pyridine N-oxide 116 by reaction of pyridine N-oxide 9 with the arylboronic ester 114 in the presence of Cu catalyst and base which is followed by deoxygenation to furnish the desired product 115 (Scheme 22b).

In 2015, a palladium-catalyzed cross dehydrogenative coupling of pyridine N-oxides with toluene for the regioselective arylation and benzylation of pyridine N-oxide was reported by Khan and co-workers [92] (Scheme 23). The authors have shown toluene 117 when used as benzyl and aryl source remained intact under the reaction conditions without any further oxidation. Different oxidants resulted in different products such as the monoarylated product 118 formed in the presence of TBHP as oxidant and the benzylated product 119 was obtained when potassium persulfate was used. Interestingly, aza-fluorene N-oxide 119b was formed during benzylation of 2-ethylpyridine N-oxide. A possible mechanism has also been reported (Scheme 23b). Electrophilic palladation at the C2-position of pyridine N-oxide 9 provides intermediate 120. The radical intermediate 121 is generated in situ by H-atom abstraction from toluene 117 by sulfate radical anion. Coordination of intermediate 120 and 121 leads to complex 122 which undergoes reductive elimination to provide product 119.

In 2016, Wei and co-workers [93] reported the arylation of pyridine N-oxides 9 employing potassium (hetero)aryltrifluoroborates 126 as coupling partner using palladium acetate and TBAI (Scheme 24). Electron-withdrawing and donating groups on the pyridine N-oxide 9 resulted in the corresponding C2-arylated products 127 in good to excellent yields with high site selectivity. A catalytic mechanism was proposed in which the electrophilic C–H palladation of pyridine N-oxide 9 occurs preferentially at the C-2 position leading to heterocoupling intermediate 128. Subsequent transmetalation provides the arylpalladium intermediate 129 which after reductive elimination furnishes the desired product 127.
Scheme 23: Pd-catalyzed C–H arylation/benzylation with toluene.
In 2017, Chen and group [94] developed a protocol for the C2,C6-arylation of pyridine under Pd catalysis (Scheme 25). In their study, N-alkylpyridinium salts were used as a directing group, facilitating the C–H arylation of pyridine. Dimethyl sulfate was used as a good N-methylating agent, which acts as transient activator. The group performed HRMS and KIE studies and proposed a catalytic cycle (Scheme 25b). The oxidative addition of ArBr 130 to the in situ-formed Pd(0) species gives species 132 followed by transmetalation with CuI pyridyl species 133 generated from the reaction of Cu2O with the methylated pyridine to afford intermediate 134 that on reductive elimination results in salt 135. Subsequent demethylation of 135 gives monoarylated product 136 or the intermediate 135 reenters the catalytic cycle to produce the diarylated N-methylpyridinium species, which again undergoes demethylation to produce product 131.

C-3 Arylation
In 2011 and 2013, the groups Yu [95] and Tan [96], reported a ligand-assisted distal arylation selectively taking place at the meta-position in pyridine. Both groups used Pd(OAc)2 as catalyst with 1,10-phenanthroline as ligand. The group of Yu used aryl halides 137 as coupling partner, whereas the group of Tan utilized aryl tosylates 142 as coupling partner (Scheme 26). The Yu group also applied the developed protocol for the synthesis of the drug molecule preclamol (139, Scheme 26b). The presumed catalytic cycle (Scheme 26c) involved the coordination of Pd(II) to the pyridine nitrogen to give N-bound pyridine substrate A followed by the formation of Pd(II) intermediate (B) involving the π-system of pyridine, which initiates the activation of the C(3)–H of pyridine to form aryl–Pd(II) species 140 via intermediate C. Subsequently, oxidative addition takes place in the presence of the aryl halide to give the Pd(IV) complex 141 followed by reductive elimination furnishing 3-arylpyridines 138.

Almost at the same time, Yu and co-workers reported the selective Pd(0)/PR3-catalyzed C3 or C4-arylation of nicotinic and isonicotinic acids using amide as a directing group (Scheme 27) [97]. This method provides a way for arylated nicotinic acid derivatives which serve as building blocks for biologically important molecules. This was the first report for a directing group-

Scheme 24: Pd-catalyzed pyridine C–H arylation with potassium aryl- and heteroaryltrifluoroborates.
assisted C3/C4-arylation of pyridines. The authors screened various \( N \)-arylamide directing groups 144 out of which \( N \)-phenylamide was found to be the better directing group. Then, the authors screened various nicotinic and isonicotinic acids which afforded the desired products 145 and 146 in good yields generating a library of isonicotinic and nicotinic acid derivatives.

Another inexpensive and non-toxic iron-catalyzed C–H arylation of pyridines has been reported by DeBeof and co-workers [98]. Using the imine in 147 as directing group, afforded the arylated pyridine products 150 in good to high yields (Scheme 28). In this reaction, Grignard reagent 148 was used as arylation source in excess amount as the reagent underwent homocoupling leading to the formation of biaryl systems under the reaction conditions. 1,2-Dichloro-2-methylpropane (149) was found to be an effective oxidant under the reaction conditions. Also, the additive KF was employed in order to minimize the oxidative iron-catalyzed homocoupling of 148. An imine directing group at the \( para \)-position in pyridine 147 lead to activated \( ortho \)-position products 150 within 15 minutes. The imine group of the products can further be hydrolyzed to get the corresponding ketones.

In 2018, Albéniz and group [99] reported the direct C3-arylation of pyridines with the help of bipy-6-OH as coordinating ligand under palladium catalysis (Scheme 29). In most of the cases the arylated pyridines 152 were obtained as mixtures of \( ortho \)-/\( meta \)-/\( para \)-substitution, however, the authors found that the yield of the \( meta \) (C-3)-arylated pyridines were drastically higher, thereby showcasing the regioselectivity of the reaction. The chelating anionic ligand acted as base in the catalytic cycle, allowing for the oxidative addition of the arene to the Pd complex. The proposed mechanism (Scheme 29b) involves the ox-
Scheme 26: Ligand-promoted C3-arylation of pyridine.
dative addition of the aryl halide to the Pd(0) complex in the presence of base ligand to afford 153. Subsequently, the substitution of the halide by pyridine 1 provides the intermediate 154 which undergoes C–H activation followed by reductive elimination to furnish the C3-arylated product 152.

**Scheme 27:** Pd-catalyzed arylation of nicotinic and isonicotinic acids.

**Scheme 28:** Iron-catalyzed and imine-directed C–H arylation of pyridines.

**Heteroarylation**

**C-2 Heteroarylation**

Heteroaryl groups are a common core in natural products and pharmaceuticals. In addition, the heterodiaryl systems widely occur in biologically important organic molecules, dyes,
Thus, the functionalization of the pyridine core with a heterocycle is a desirable transformation in organic synthesis. Manickam and co-workers [100] carried out a palladium-catalyzed decarboxylative ortho-(hetero)arylation of pyridine N-oxides 9 with heteroarylcarboxylic acids 156 (Scheme 30).

The reaction showed good compatibility with various functional groups. The proposed mechanism (Scheme 30b) involves the silver-catalyzed decarboxylation of heteroaryl acid 156 followed by transmetalation providing palladium intermediate 160. Further, C–H activation of pyridine N-oxide 9 provides intermediate 161 which upon reductive elimination furnishes the desired product 157 and regeneration of Pd(0) (Scheme 30b).

Later in 2014, Kuang and co-workers [101] developed a highly efficient and regioselective oxidative cross-coupling of pyridine N-oxides 9 with five-membered heterocycles 162 and 163 through a two-fold C–H activation under palladium catalysis. Silver carbonate and 2,6-lutidine were found to be an effective base and ligand, respectively, for providing the desired products 164 and 165 in good yields (Scheme 31).

In 2015, an economic route for copper-catalyzed biaryl coupling of azine(pyridine)-N-oxides 9 with oxazoles 166 was reported by Miura and group [102]. Although their work majorly covered quinoline N-oxide substrates, they also investigated three pyridine substrates in the reaction leading to the corresponding products in moderate yields (Scheme 32). The N-oxide plays a role as an activator and is subsequently eliminated via deoxygenative elimination furnishing the C-2-functionalized pyridines 167. The reaction mechanism (Scheme 32b) involves the initial C–H-cupration of 166 producing an oxazolyl–copper intermediate 168. Nucleophilic addition fol-
Scheme 30: Pd-catalyzed pyridine N-oxide C–H arylation with heteroarylcarboxylic acids.

lowed by C–H activation of 9 provides the hydroxy copper species 169, which on deoxygenative elimination furnishes the desired product 167.

C-3 Heteroarylation
In 2013, Su and co-workers [103] developed a catalytic methodology for the distal heteroarylation of pyridines 170 via Rh(III)-catalyzed dehydrogenative cross-coupling showcasing a good substrate scope (Scheme 33). Initially, their investigation involved evaluating the reaction between N-phenylisonicotinamide 170 and 2-methylthiophene 171 which resulted in the desired product 172. The plausible mechanism (Scheme 33b) starts with the initial coordination of the pyridine directing group 170 with rhodium providing a five-membered rhodacyclic intermediate I which further forms the aryl–rhodium(III) complex II by reaction with 171. Subsequently, this intermediate undergoes reductive elimination from the rhodium(III) center to furnish the desired ortho-C–H-arylated product 172 releasing a Rh(I) species. The Rh(III) species is regenerated in the presence of the copper salt.
Scheme 31: Pd-catalyzed C–H cross-coupling of pyridine N-oxides with five-membered heterocycles.

Scheme 32: Cu-catalyzed dehydrative biaryl coupling of azine(pyridine) N-oxides and oxazoles.
In another case of C3-(hetero)arylation, Yu and group [104] using palladium for C–H activation of pyridine with phenanthroline as a ligand developed a method in 2016 (Scheme 34). The authors achieved both arylation and heteroarylation at the C-3-position in pyridine and showcased the formation of bipyridines 174. The mechanism is depicted in Scheme 34b, where the complex A undergoes C3–H activation to provide intermediate 176 which similarly undergoes one more step of C–H activation to provide the bi(hetero)arene–Pd(II) species 177 which undergoes reductive elimination furnishing the desired products 174/175.

**C–H Annulation of pyridine to fused heterocycles**

Annulation reactions in organic synthesis have achieved great attention toward the construction of various carbocycles and heterocycles. These annulations can be either intermolecular or intramolecular and various substrates have been studied resulting in diverse products. Pyridine has been also reported for the construction of pyridine-fused heterocycles via C(sp^2)–H functionalization and further annulation. In this aspect, considering the use of pyridines for the formation of quinolines and isoquinolines, an oxidant-dependent rhodium-catalyzed C–H annulation of pyridines with alkynes was reported by Li and co-workers [105] in 2011 for the direct synthesis of quinolines 180 and isoquinolines 181 involving a two-fold C–H activation of pyridine at the C2 and C3 position (Scheme 35a). Further, during optimization when silver additives like Ag2CO3, Ag2O, and AgOAc were used the reaction resulted in the formation of isoquinoline derivative 181. In addition, the reaction showed high regioselectivity in the presence of unsymmetrical alkynes 179. Different directing groups 178 were employed resulting in
diversified products 180. The proposed mechanism (Scheme 35b) involves coordination of rhodium with iso-
icotinamide 178 and subsequent ortho-C–H activation generating the five-membered rhodacycle 183. Next, first alkyne 179 insertion results in the five-membered rhodacycle 184 which is followed by a second regioselective insertion of alkyne 179 into the Rh–C bond of 184 providing the seven-membered cyclic intermediate 185. Further reductive elimination furnishes the quinoline product 180 and a Rh(I) species, with the latter being oxidized by Cu(II) to complete the catalytic cycle.

Next, considering the role of N-heterocyclic carbene (NHC) ligands acting as directing group as well as functionalizing unit in arene C–H functionalization reactions with alkynes, Choudhury and group [106] in 2015 developed a protocol for the intermolecular C–H annulation of NHC-substituted pyridines with a variety of internal alkynes 187 under rhodium catalysis for the synthesis of annulated and highly decorated pyridines 188 (Scheme 36). The authors used the N-heterocyclic carbene ligand as directing group to prepare imidazo[1,2-α][1,6]naphthyridine motifs 188 as desired products. Based on the experimental results and annulation chemistry a catalytic mechanism has been proposed (Scheme 36b) that involves the C3 hydrogen of pyridine undergoing a cyclorhodation with the catalyst in the presence of NaOAc, directed by in-built NHC ligand coordination to provide intermediate 189. The further coordination of
Scheme 35: Rhodium-catalyzed oxidative C–H annulation of pyridines to quinolines.

189 with the alkyne 187 results in intermediate 190 and subsequent insertion provides rhodacycle intermediate 191 which undergoes reductive elimination to furnish the product 188 via dissociation of intermediate 192 along with oxidative regeneration of 189 (Scheme 36b).

In 2019, using NHC ligands, a protocol for the regio- and enantioselective C–H cyclization of pyridines was reported by Shi and co-workers [107] toward the direct asymmetric pyridine C–H alkylation (Scheme 37). The authors found that alkene-tethered C2 pyridine 193, C3 pyridine 195 and C4 pyridine 197 can undergo endo-cyclization reactions in the presence of Ni(cod)$_2$, a chiral NHC ligand, and MAD as Lewis acid to afford optically active 5,6,7,8-tetrahydroquinolines 194 and 5,6,7,8-tetrahydroisoquinolines 196 and 198. The endo-selective annulation approach was compatible with various tethered
alkenes, such as 1,1-disubstituted alkenes, styrene, diene, trisubstituted alkene and enamines. To get insights into the mechanism the authors conducted additional experiments including deuterium labelling reactions and proposed the mechanism depicted in Scheme 37b. Initially, the sterically bulky additive MAD coordinates to the pyridine nitrogen, which pushes the tethered alkene close to the nickel center subsequently providing the intermediate 201. Then, the C–D bond on cleavage via oxidative addition of Ni(0) forms the Ni–D species 202 which after anti-Markovnikov hydronickelation of the alkene provides
The seven-membered cyclic intermediate 203. Subsequent reductive elimination furnishes the endo-annulated product 194 (Scheme 37b).

Out of various pyridine-fused heterocyclic hybrids, azaindolines are important scaffolds in natural products and pharmaceuticals serving different biological activities. Hence, looking at the importance of azaindolines in drug discovery a protocol of rare earth metal-catalyzed intramolecular insertion of the pyridine C–H bond into unactivated vinyl C–H bonds has been developed by Chen and co-workers [108] (Scheme 38). Using this protocol azaindolines were accessed in moderate to excellent yields and also naphthyridine derivatives (205k and 205l) were synthesized. In the proposed mechanism, the initial deprotonation of HNBn2 by Ln[N(TMS)2]3 provided the lanthanide amide. Activation of the vinyl-substituted pyridin-3-amine by the lanthanide amide gives a lanthanide–pyridine complex 206. Then, coordination and sequential insertion of C=C into

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**Scheme 37: Ni/NHC-catalyzed regio- and enantioselective C–H cyclization of pyridines.**
the Ln–pyridine bond of 206 provided intermediate 207, which undergoes intermolecular protonation with 204 to afford the desired product 205 and regenerating the lanthanide species (Scheme 38b).

**C(sp\(^2\))–H Functionalization of pyridine rings in bipyridine systems**

The functionalization and synthesis of bipyridine derivatives is of great interest and importance in synthetic chemistry. These compounds are well-studied for their roles as chelating ligands in transition-metal-catalyzed reactions, coordination chemistry including materials science [109,110]. The challenge associated with the C–H functionalization of bidentate molecules is the finding strategy in the subduing the high activation barrier of rollover cyclometallation pathway. In this section we discuss the C(sp\(^2\))–H functionalization of the pyridine ring in bipyridine ring systems. In early 2009 Miura and co-workers [111] reported the rhodium-catalyzed regioselective reaction of aryl-N-heterocycles and aromatic imines with terminal silylacetylenes 209 to synthesize C–H-alkenylated products 210. The terminal silylacetylenes were employed as effective substrates for catalytic cross-dimerization reactions. The reaction was performed in the presence of [RhCl(cod)]\(_2\) (3 mol %), taking PPh\(_3\) or (4-ClC\(_6\)H\(_4\))\(_3\)P as the ligand at 160 °C, for about 48 h (Scheme 39). This work provides an effective way for preparing C–H-alkenylated bipyridines 210.

In 2012, a hydroarylation of alkenes 211 and alkynes 212 with 2,2′-bipyridines 208 and 2,2′-biquinolines was reported by Chang and co-workers [112] in the presence of Rh(acac)\(_3\) as catalyst, IMes·HCl (3 mol %) as ligand and t-BuONa (30 mol %) in toluene for 2 h (Scheme 40). The authors demonstrated theoretically and mechanistically the important role of the NHC ligand in the resultant catalyst Rh(NHC) for the hydroarylation of alkenes and alkynes with chelating 2,2-bipyridine and 2,2-biquinoline molecules. The experimental studies revealed that the trans-effect of the NHC ligand in the complex assisting in the reduced energy barrier of a rollover cyclometallation pathway and results in selective and efficient hydroarylation of the alkenes and alkynes. This was the first report for the role of a “rollover” cyclometallation pathway catalytically leading to double C–H bond functionalization of chelating mol-
Scheme 39: Rh-catalyzed alkenylation of bipyridine with terminal silylacetylenes.

Scheme 40: Rollover cyclometallation in Rh-catalyzed pyridine C–H functionalization.
Scheme 41: Rollover pathway in Rh-catalyzed C–H functionalization of N,N,N-tridentate chelating compounds. Next, a protocol for the selective and catalytic C–H functionalization of N,N,N-tridentate chelating compounds using a rollover cyclometallation strategy was reported by the same group in 2016 [113]. The reaction involves the Rh-catalyzed alkylation of 2,2',6',2”-terpyridine 221 with 3,3-dimethyl-1-butene coupled in the presence of a catalytic amount of t-BuONa providing the mono- and dialkylated products in low combined yields. The alkylation of terpyridines with aliphatic olefins 222 afforded only anti-Markonikov linear products 223 (Scheme 41). The authors also expanded their study to tridentate heteroarenes. Delightfully, they observed the dialkylation...
products 223 in good yields. The plausible reaction mechanism (Scheme 41b) was explained by the formation of a cationic Rh–terpyridine complex 224 generated from terpyridine 221 and a Rh(NHC) species formed from the Rh(I) precursor and the NHC in the presence of an external base and successive decomplexation of 224 provides complex 225. The latter undergoes an initial key rollover cyclometallation followed by oxidative addition leading to the metal–hydride intermediate 226 which on olefin insertion and subsequent reductive elimination resulted in the monoalkylated rhoda complex 227. Complex 227 then undergoes recomplexation to form 228 and enters the subsequent catalytic cycle furnishing the bisalkylated product 223.

In 2018, Cheng and co-workers [114] reported a straightforward approach to 3'-aryl-2,2'-bipyridine-6-carboxamide derivatives 231 with exclusive selectivity starting from 2,2'-bipyridine-6-carboxamides 229 under Pd catalysis (Scheme 42). The arylation reaction of N-butyl-2,2'-bipyridine-6-carboxamide with iodobenzene 230 in the presence of Pd(OAc)$_2$ as catalyst, Cs$_2$CO$_3$ as a base in DMSO at 160 °C furnished the desired products 231 (Scheme 42). It was found that non-polar solvents resulted in good yields of the products 231. It is reported that 2,2'-bipyridine-6-carboxamides 229 can bind to the transition metal, such as Pd(II), to form stable $N,N,N$-chelates I (Scheme 42b). The amide moiety of the $N,N,N$-chelates I exerts a strong trans-effect which weakens the Pd(II)–pyridyl bond trans to the amide anion, thus, allowing the decomplexation to afford complex II which is key intermediate for furnishing the desired C–H functionalization product (Scheme 42b).

In 2019, Cheng and co-workers reported an approach for the C3-selective acylmethylation of [2,2’-bipyridine]-6-carboxamides 232 with sulfoxonium ylides 233 in the presence of a Rh(III) catalyst (Scheme 43) [115]. Sterically hindered amide directing groups were also well tolerated under the optimal conditions. A H/D exchange reaction exclusively at the C3-position suggested C–H-bond cleavage is reversible. The catalytic cycle involves the coordination of the carboxamide 232 with the Rh(III) species affording Rh(III) complex 235, which on rollover cyclometalation gives the complex 236. The addition of sulfoxonium ylide 233 to the intermediate complex 236 generates the Rh–carbene complex 237 with the release of DMSO and further migratory insertion of complex 237 and subsequent

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Scheme 42: Pd-catalyzed rollover pathway in bipyridine-6-carboxamides C–H arylation.
protonolysis furnishes the acylmethylated product 234 (Scheme 43b).

Recently, in 2020, Zhu and co-workers [116], developed a novel annulation reaction of bipyridine systems 211 with alkynes 239 via a Rh(III)-catalyzed dual C–H functionalization. The authors have initiated their studies with 6-bromo-2,2'-bipyridine as their model substrate and with diphenylacetylene as coupling partner. The optimized conditions included [RhCp*Cl₂]₂ (5 mol %), AgOAc (2.5 equiv), NaOAc (5 equiv) in DCE, at 110 °C for 24 h to obtain the annulated product 240 (Scheme 44). The proposed mechanism (Scheme 44b) involves the formation of Rh(III) complex 241 by coordination of the bipyridine with rhodium and complex 241 via a rollover cyclometallation process gives the intermediate 242. It was suggested that the substitution at the 6 position of the bipyridine ring system facilitates the rollover cyclometallation process by weakening the Rh–N bond. Next, intermediate 242 coordinates with alkyne 239 to give the seven-membered rhodacycle 243. The excess Ag⁺ help in the dissociation of the N–Rh bond in
complex 243 and give the five-membered rhodacyclic interme-
diate 244 which again coordinates with the alkyne 239
furnishing another seven-membered rhodacyclic intermediate
245 or 246. Finally, reductive elimination delivers the desired
product 240.

In the subsequent year, the same group reported a method for
the rhodium-catalyzed acylmethylation of bipyridines [117].
The group has demonstrated a switchable reaction, wherein
changing the additive can deliver the acylmethylated product
248 or the annulation product pyrido[2,3-a]indolizine 249
(Scheme 45). Under action of the Rh(III) catalyst, zinc acetate
and PivOH as additives, the acylmethylation of bipyridines
takes place at the C-2 position to furnish acylmethylated prod-
ucts 248 and the reaction was found suitable for various sub-
strates. On the other hand, the usage of silver acetate as an addi-
tive provided the annulated (intramolecular cyclization of
bipyridine) product 249.

Miscellaneous reactions
C–H Borylation
Due to the broad utilities of arylobonic esters in organic syn-
thesis, various protocols have been reported till date for their in-
corporation into an organic molecule. In 2017, Nakao and group
reported a method for the iridium-catalyzed para-C–H boryla-
tion of pyridines using bis(pinacolato)diboron (250) for the syn-
thesis of borylated pyridines 251, which are important interme-
diates for various derivatization reactions (Scheme 46) [118]. In
common, site-selective borylations have been in less focus, due
to the lack of suitable strategies, however, this group achieved
the para-selective borylation of pyridines using a cooperative catalyst strategy. The authors used [Ir(cod)(OMe)]$_2$ as a metal catalyst, along with a sterically bulky Lewis acid such as methylaluminum bis(2,6-di-tert-butyl-4-methylphenoxide) as a cooperative catalyst.

Later, in 2019, the same group reported a protocol for the selective C5(C3)-borylation of pyridines under iridium–Lewis acid bifunctional catalysis (Scheme 47) [119]. With the optimized conditions in hands, the authors screened for the substrate scope of substituted pyridines. Also, they employed the reported protocol for the late-stage functionalization of brompheniramine (252d), an antihistaminic drug.

C–H Silaboration

In 2011, a protocol for the synthesis of highly functionalized dihydropyridines via palladium-catalyzed silaboration providing silylated dihydropyridines 255 and 256 (Scheme 48) was developed by Suginome and co-workers [120]. This reaction involved a dearomatizing conversion of pyridines to dihydropyridines under mild conditions with the introduction of a silyl group on a carbon atom of pyridine ring. Various pyridines

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**Scheme 46:** Iridium-catalyzed C4-borylation of pyridines.

**Scheme 45:** Rh-catalyzed C–H acylmethylation and annulation of bipyridine with sulfoxonium ylides.
Scheme 47: C3-Borylation of pyridines.

Scheme 48: Pd-catalyzed regioselective synthesis of silylated dihydropyridines.
were subjected to this elaboration using the Pd/PCy$_3$ catalytic system providing the corresponding products in good yields. The proposed mechanism (Scheme 48b) involves the oxidative addition of silylboronic ester 254 to Pd(0) and coordination of pyridine 1 providing the intermediate 257 which on further regioselective insertion of pyridine into the Pd–B bond resulted in the π-allyl palladium complex 258. Subsequent reductive elimination furnishes the silaboration products 255 and 256 with the regeneration of Pd(0).

Conclusion

Significant progress has been made in the area of ortho- and distal C–H-functionalization of pyridines, as evidenced by the reactions outlined in this review. The previous research and their mechanistic insight provided us with more information to approach the new avenue of catalytic C–H functionalization of the pyridine nucleus. The challenges still remain for the distal C–H functionalization, particularly at the C4 position. Even the directing group on pyridine ring system has been less explored for ortho- or distal C–H functionalization. Although the C–H functionalization with transition-metal catalysis and rare earth metal catalysis has advanced, the functionalization of the pyridine ring system can further be explored by employing new catalytic systems and merging of different strategies. Taking this into account, we hope that the efforts for the development of novel protocols for the preparation and incorporation of functionalized pyridine scaffolds will continue and could be applicable for applications in industry.

ORCID® iDs

Haritha Sindhe - https://orcid.org/0000-0002-4510-2209

Satyasheel Sharma - https://orcid.org/0000-0002-4983-1572

References

Aromatic C–H bond functionalization through organocatalyzed asymmetric intermolecular aza-Friedel–Crafts reaction: a recent update

Anup Biswas

Abstract
The aza-Friedel–Crafts reaction allows an efficient coupling of electron-rich aromatic systems with imines for the facile incorporation of aminoalkyl groups into the aromatic ring. This reaction has a great scope of forming aza-stereocenters which can be tuned by different asymmetric catalysts. This review assembles recent advances in asymmetric aza-Friedel–Crafts reactions mediated by organocatalysts. The mechanistic interpretation with the origin of stereoselectivity is also explained.

Introduction
The ease of a chemical transformation depends on the thermodynamic instability of a chemical bond owing to its fast cleavage under mild reaction conditions. A C–H bond is thermodynamically stable and possesses a high bond dissociation energy opposing the bond to easy chemical transformation. Therefore, harsh reaction conditions and the necessity of an external activator like catalysts are common prerequisites for processes involving C–H bond breaking. Among different types of C–H bonds, an aromatic C–H bond is even more inert rendering this type of bond functionalization more difficult. Herewith the term “bond functionalization” is defined as the cleavage of an existing bond with substitution by another bond.

Aromatic C–H bond functionalizations have gained considerable attention by organic chemists because of the strategic importance of this process as well as the ability to synthesize functionalized aromatic molecules in a straightforward way. Many organic name reactions have been discovered utilizing the C–H bond functionalization concept [1].

Metals were exclusively explored to assist substitutions of aromatic C–H bonds by other bonds and this area of research is more than a century old. However, many disadvantages are associated with metal-mediated organic transformations including harsh reaction conditions (e.g., high temperature) and toxic...
solvents. With the tremendous progress in organic chemistry over the last few decades, metal catalysis has been increasingly and successfully replaced by organocatalysis, i.e., accelerating the rate of chemical transformations by using small organic molecules as catalysts. Although being discovered more than 100 years ago, the concept became increasingly accepted and popular only by the last decade of the last century [2,3].

Nowadays, organocatalysis is especially applied to asymmetric synthesis and a huge number of organocatalysts has been introduced in last three decades for the asymmetric synthesis of acyclic, carbocyclic, heterocyclic, and polycyclic molecular architectures with high molecular complexity. In particular, asymmetric organocatalysis plays a pivotal role in the construction of optically active, bioactive, and natural products. The main advantages of organocatalyzed stereoselective reactions include mild reaction conditions and the use of a sole catalyst without the need of other chiral ligands [4,5]. In these reactions, stereoinduction in the products is achieved by the chiral environment present in the catalyst itself. Depending upon the reactivities, organocatalysts can be categorized into two major divisions: 1) covalent bonding and 2) noncovalent bonding catalysts. A covalent bonding organocatalyst reacts with a substrate to form an activated chiral intermediate which undergoes a stereoselective reaction with another reagent. A noncovalent bonding catalyst usually assembles the reaction partners in a highly ordered three dimensional transition state through noncovalent interactions (like H-bonding, π–π interactions) thus promoting the stereoselective reaction. Examples of covalent bonding organocatalysts are amines [6,7], N-heterocyclic carbenes [8,9], phosphines [10], amidines [11], isothioureas [12,13], whereas thioureas [14,15], ureas [16], phosphoric acids [17,18], and squaramides [19,20] fall into the second category.

The Friedel–Crafts reaction, discovered by Charles Friedel and James Crafts in 1877 allows the aromatic C–H bond functionalization through the formation of a new C–C bond [21]. The reaction requires an electrophilic reagent/intermediate present in the reaction system on which an electrophilic attack by the π-electron cloud of the aromatic ring can occur spontaneously to form a dearamatized species. The latter is rearomatized in a succeeding step with the elimination of a H⁺ ion to form the functionalized aromatic moieties. The aza-Friedel–Crafts reaction is a subclass of the originally reported transformation that incorporates an aza-tertiary stereocenter into the 2′ position of the heteroaromatic product (Scheme 1) [24].

The first organocatalyzed asymmetric aza-Friedel–Crafts protocol was published by Terada and co-workers in 2004. In this methodology, a 1,1′-bi-2-naphthol (BINOL)-derived chiral phosphoric acid P1 was used as the catalytic reagent to couple 2-methoxyfuran (1) and N-Boc-protected aldimines 2 to incorporate an aza-tertiary stereocenter into the 2′ position of the heteroaromatic products 3 (Scheme 1) [24].

This review summarizes the recent advances (2018 till date) on organocatalyzed asymmetric aza-Friedel–Crafts reactions. The examples have been segmented according to the different types of catalysts.

**Review**

**Phosphoric acids**

Chiral phosphoric acids have been envisaged as versatile organocatalysts for various asymmetric chemical transformations. These compounds play a dual role in the catalytic cycle due to their intrinsic Brønsted acidity and the ability to H-bond formation. Organophosphoric acids can perform as both H-bond acceptors and donors. 1,1′-Bi-2-naphthol (BINOL) and 1,1′-spiroindane-7,7′-diol (SPINOL)-derived phosphoric acids with different substituents in the 2,2′-positions of the aromatic framework have been extensively explored as axially chiral catalysts in the field of asymmetric transformations including aza-Friedel–Crafts reactions.

In 2018, Nakamura and co-workers designed an aza-Friedel–Crafts process between indoles 4 and cyclic N-sulfonyl ketimines 5. The authors employed the BINOL-based chiral...
phosphoric acid P2 bearing two imidazoline moieties at the ortho-positions as the catalyst which activates both reactants through H-bonding where the NH group of the nucleophile performs as an H-bond donor towards the imidazoline nitrogen and the electrophile acts as H-bond acceptor from the OH group of the catalyst. These interactions rearrange the three molecules in a chiral pocket as shown by transition state 7, favoring stereoinduction in the products through C3-functionalization of the indole (Scheme 2) [25].

In 2018, Lin and co-workers deployed pyrroles 9 in an aza-Friedel–Crafts reaction with trifluoromethyldihydrobenzoazepinoindoles 8 to achieve the aromatic electrophilic substitution at the C2 position of the pyrrole ring. A further extension of the scope of this process was achieved through the C3–H functionalization of indole derivatives 4. The nucleophile favors the attack at the imine carbon included in the seven-membered ring of compound 8 to generate an aza-quaternary stereocenter containing trifluoromethyl, pyrrole/indole, and benzoazepinoindole moieties. Stereoselectivity in the products 10/11 was achieved by using the chiral spirocyclic phosphoric acid catalyst P3 which, through H-bonding interactions with the nucleophile and the electrophile, forces the nucleophile to approach the C=N plane from the Re face. In general, enantiocontrol with pyrroles was better than with indoles (Scheme 3) [26].
In 2018, Kim and co-workers developed an aza-Friedel–Crafts protocol involving pyrroles 9 as the π-nucleophile in combination with cyclic N-sulfimines 12. The chiral phosphoric acid P4 was used to catalyze the introduction of a pyrrole-substituted aza-quaternary stereocenter in cyclic sulfamidate derivatives. N-Alkyl and N-benzyl-substituted pyrroles responded to the process with appreciable enantioefficiency. However, pyrrole was not proved to be an efficient substrate in terms of stereocore [27] (Scheme 4a). In the very next year, pyrrole was successfully replaced by 2-substituted furans 1 as the aromatic reacting partner with imines 12 to execute the asymmetric aza-Friedel–Crafts process modulated by the chiral phosphoric acid P5 as the catalyst. A major concern of this process was the reduced aromatic character of the furan ring and the C2 methoxy-substituted substrate was exclusively employed to make the aromatic ring sufficiently electron rich. The substrate scope was mainly attributed to alterations of the substituents on the benzene ring of imines 12 (Scheme 4b) [28].
In 2018, Morimoto, Ohshima and co-workers reported an aza-Friedel–Crafts process for the functionalization of the C3–H bond in indoles in the presence of BINOL-derived chiral phosphoric acid P6 as the catalytic agent. They utilized tri-fluoromethyl ester-substituted N-unprotected imine 15 as the potential electrophile to install an aza-quaternary stereocenter in the C3 position. The products 16 were achieved with excellent enantioselectivities which were attributed to an attractive interaction between the indole ring and the anthracene substituent of the catalyst’s framework (Scheme 5) [29].

In 2018, Piersanti and co-workers developed a phosphoric acid-catalyzed cascade reaction proceeding through aza-Friedel–Crafts reaction and lactonization steps. Main focus of this article was to demonstrate a racemic process between α-naphthol or phenol derivatives and in situ-generated N-acetyl ketimine from methyl 2-acetamidoacrylate (18) in the course of preparing 3-NHAc-naphthofuran or benzofuran analogues. The achiral phosphoric acid (PhO)2P(O)OH was the catalytic reagent to execute the process delivering the products with low to moderate chemical yields. Attempts to make the process stereoselective, a series of chiral phosphoric acid catalysts were screened in the model reaction between α-naphthol (17) and methyl 2-acetamidoacrylate (18) but promising selectivity was not achieved. The highest enantiomeric excess of 64% was obtained in the presence of P7 as the catalyst (Scheme 6) [30].

In 2018, Reddy and co-workers developed a one pot protocol comprising oxidation and an enantioselective aza-Friedel–Crafts addition. In the first step, the DDQ-promoted oxidation of 3-indolinonecarboxylate 22 generated indolenines that performed as the potential electrophiles towards indoles 4. The chiral catalyst effectively assembled the reacting partners in a chiral transition state through H-bonding interactions to facilitate a highly face-selective nucleophilic attack by π-nucleophile to the cyclic imine (see transition state 22’ in Scheme 7a). The BINOL-derived chiral phosphoric acid P8 was employed as the asymmetric organocatalyst for this transformation to construct the heterodimerized products 23 framed with an aza-quaternary stereocenter. Indole derivatives without any substitution in the

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**Scheme 5:** Aza-Friedel–Crafts reaction involving N-unprotected imino ester as electrophile.

**Scheme 6:** Aza-Friedel–Crafts and lactonization cascade.
heterocyclic ring participated in the reaction through the C3 position smoothly providing the products with appreciable yields and enantiocontrol. Two examples were demonstrated with 3-alkyl-substituted indoles which effectively attacked the electrophile through the C2 position. The reaction was even compatible with pyrroles (Scheme 7a). The utility of this methodology was successfully demonstrated by the synthesis of product 23a, the key intermediate of natural product (+)-trigonolimine (Scheme 7b) [31].

In 2018, Ishihara and co-workers developed a novel C2 and C1-symmetric bisphosphoric acid-catalyzed asymmetric aza-Friedel–Crafts reaction. Both catalysts showed intramolecular H-bonding causing a sharp increase in Brønsted acidity of free OH groups and prevention of catalyst dimerization. The C2-symmetric P9 promoted the reaction between 2-methoxyfuran (1) and aryl-α-ketimino esters 26 to activate the C2′–H bond in 1. The C1-symmetric catalyst P10 was the optimal catalyst for the second reaction furnishing the products with excellent chemical yields and enantioselectivities. To understand the activities of the catalysts, the authors were able to obtain X-ray crystallographic data of the pyridine–catalyst complex which showed two intramolecular H-bonding interactions in the molecular framework of the catalyst where two free OH groups were engaged in interactions with the pyridine. This data clearly indicates the activation of the reaction components through H-bonding engagement with free hydroxy groups of the catalysts also favoring stereoselective addition (see structure 28 in Scheme 8a) [32]. Two years later, the same research group utilized the C1-symmetric catalyst P10 for the functionalization of the C3–H bond of indole 9 through aza-Friedel–Crafts reaction with aryl-α-ketimino esters 26/29. They also utilized unsubstituted and 2,3-disubstituted pyrroles 9 as π-nucleophile towards the same electrophiles to incorporate an amine-substituted quaternary stereocenter at the C2′ position (Scheme 8b) [33].
In 2019, Inokuma, Yamada and co-workers reported the C3–H bond functionalization of indoles 4 through aza-Friedel–Crafts reaction utilizing N-α-nitrophenylsulfonyl (Nps)-iminophosphonates 32 as electrophiles. The chiral phosphoric acid P11 was used as H-bonding catalyst to impart stereoselectivities into the products, i.e., α-3-indolyl-α-aminophosphonic acids 33. The reaction was also well compatible with pyrroles 4 proceeding through C2–H substitution. With C2-substituted pyrrole, the electrophile enters into the C5 position (Scheme 9) [34].

In 2019, Palacios, Vicario and co-workers documented an aza-Friedel–Crafts reaction between indole 4 and α-iminophosphonate 35. The reaction functionalized the C3 position of the heterocyclic ring with an α-aminophosphonate group. Chiral phosphoric acid P12 was the stereoselectivity inducer in the products 36 as explained by π–π stacking and H-bonding interactions between the catalyst and the substrates (see transition state 37 in Scheme 10). The presented substrate scope was not broad and poor to moderate enantioselectivities were obtained. Indoles with a substituent in the carbocyclic ring required shorter reaction times to accomplish in comparison to C2-substituted indoles. The authors also tried the reaction with C3-substituted indoles to functionalize the C2 position. However, a very low enantioselectivity was achieved in the latter case (Scheme 10) [35].

Lin and co-workers designed a planar chiral phosphoric acid containing a [2.2]paracyclophane moiety that efficiently catalyzed the aza-Friedel–Crafts reaction between indole 4 and N-tosyl vinylaldimines 38 to functionalize the C3–H bond of the heterocyclic ring. The authors tried six such catalysts by varying the aromatic substituents, among which P13 was proved to be the best one in terms of both yields and enantioselectivity. The catalyst P13 was an even far superior catalyst than conventional BINOL and SPINOL-derived phosphoric acids. The substrate scope was investigated by varying substitu-
Scheme 9: Aza-Friedel–Crafts reaction using Nps-iminophosphonates as electrophiles.

Scheme 10: Aza-Friedel–Crafts reaction between indole and α-iminophosphonate.

ents in the carbocyclic ring of indole 4. Changing the β-aryl and α-substituents in the styryl-derived aldimines further expanded the substrate scope. Only 1 mol % catalyst loading was sufficiently efficient to deliver the enantioenriched products (Scheme 11a). The compatibility of the reaction was further explored by using N-tosyl arylaldimines 40 as the electrophilic partner to afford (aryl)(indolyl)methanamines 41 with high enantioselectivities. In this case, P14 was identified as the optimal catalyst (Scheme 11b) [36].

In 2019, Kim and co-workers reported a phosphoric acid-catalyzed enantioselective aza-Friedel–Crafts reaction between N-substituted indoles 4 and indol-3-ylsulfamidates 42. The dual reactivity of catalyst P5 initiated with the protonation of amidates 42 to generate intermediate 44 through ring cleavage. Then, the intermediate 44 was paired with the anionic conjugate base of catalyst P5 and acts as electrophile to facilitate the conjugate Friedel–Crafts reaction involving C3 of indole 4 as the nucleophile. This reaction afforded (bis(indolyl)methyl)benzenesulfonamide derivatives 43 but no promising enantioselectivity was achieved for most of the products (Scheme 12) [37].

In 2019, You, Yuan and co-workers reported another enantioefficient aza-Friedel–Crafts reaction between N-unsubstituted pyrroles/indoles 4/9 and isoquinoline-1,3,4(2H)-trione-1-imines 45 installing an aza-quaternary stereocenter in isoquinoline-1,3(2H,4H)-dione frameworks 46/47. The spinol-derived catalyst P15 was applied for the asymmetric induction through H-bonding interaction with the NH group of the heteroarene and amide oxygen of 45 forcing the heteroarene to approach from the Si-face of the imine moiety predominantly (see transition state 48) achieving high enantiocontrol for both heterocycles (Scheme 13) [38].
The carbocyclic ring in indoles is less reactive than the heterocyclic ring and hence the presence of an electron-donating functional group is crucial in the ring to activate it for aromatic electrophilic substitution processes. In 2019, Zhang and co-workers succeeded in the C6-selective aminoalkylation of 2,3-disubstituted indoles 4 without the presence of a directing group in the benzene ring. As the electron-demanding reaction partner, isatin-derived N-Boc-substituted ketimines 49 were employed which effectively functionalized the C6–H bond of substrate 4 to construct 3-oxindole derivatives 50 bearing an indole-substituted aza-quaternary stereocenter at its C3 position. 2,3-Dialkyl-substituted indoles having methyl or cycloalkyl substituents of different ring sizes exclusively reacted as nucleophiles. Chiral phosphoric acid P16 mediated the asymmetric transformation to regulate the stereochemical output of the quaternary stereocenter with good to excellent enantioselectivities. A resonance-assisted accumulation of negative charge on C6 enabled the carbon to add to the electrophile selectively from the Re face of the imine plane because of substrate–catalyst H-bonding interactions (see transition state 51). Beside multiple noncovalent interactions, π–π stacking between the anthracenyl group of the catalyst framework and aromatic rings of both substrates was
also responsible for the stereoselective addition (Scheme 14) [39].

In 2019, Akiyama and co-workers developed a simple enantioselective aza-Friedel–Crafts process using unprotected pyrroles 9 and indoles 4 mediated by BINOL-derived chiral phosphoric acid catalysts P17 and P18. The electrophile was the \( \alpha \)-trifluoromethyl-containing imine 52 which directed the C2 functionalization in the pyrrole moiety with catalyst P17 and a C3 substitution in indole derivatives using catalyst P18 forming the trifluoromethylated aza-quaternary stereocenter. Excellent chemical yields and good to excellent levels of enantioselectivities in the products 53/54 were obtained by the chiral catalysts. The process was robust towards \( \alpha \)-aryl- and \( \alpha \)-trifluoromethylamines and the substrate scope was mainly investigated by the variation of electron-donating groups in the aryl ring of the imines whereas amenability of this methodology was narrow for ring-substituted pyrroles and indoles (Scheme 15a) [40]. In the next year, the same research group reported another aza-Friedel–Crafts reaction between 4,7-dihydroindole (55) and N-unsubstituted trifluoromethylated ketimines 52 proceeding through C2 functionalization and follow up oxidation to provide 2-substituted indoles 56 which are typically difficult to obtain directly from unsubstituted indoles through electrophilic substitution. The process was catalyzed by the chiral phosphoric acid P17 to install a quaternary stereocenter bearing primary amine and trifluoromethyl functionalities associated with appreciable enantiocontrol. The substrate scope was investigated by the

![Scheme 13: Isoquinoline-1,3(2H,4H)-dione scaffolds as electrophiles.](image1)

![Scheme 14: Functionalization of the carbocyclic ring of substituted indoles.](image2)
variation of sterically and electronically divergent aryl substituents in the ketimines but the enantioselectivity was markedly lowered with sterically congested reactants (Scheme 15b) [41]. Very recently, Akiyama and co-workers demonstrated a C2-selective aza-Friedel–Crafts reaction of unmodified pyrroles with (alkynyl)(trifluoromethyl)imines catalyzed by the chiral phosphoric acid P17. This reaction produced an aza-quaternary stereocenter bearing 2-pyrrolyl, trifluoromethyl and alkynyl as other three substituents (Scheme 15c) [42].

In 2020, a completely para-selective aza-Friedel–Crafts protocol with N-monosubstituted aniline derivatives catalyzed by the chiral phosphoric acid P19 was disclosed by Zhu, Zhang and co-workers [43]. The electrophilic aromatic substitution involved isatin-derived ketimines as the electron-demanding partner to achieve this aromatic p-C–H bond functionalization framing an all substituted stereocenter at the C3 position of the oxindole scaffold in the products. A very low reaction temperature (−55/−60 °C) was ideal to obtain the products with satisfactory enantioselectivities. The reaction was compatible with a broad range of substrates using para-substituted phenyl rings as the nitrogen substituents in anilines. Two examples were shown with N-benzyl and N-methyl-substituted anilines which afforded the desired products as well but an elevated temperature was required for these reactions. Further expansion of the substrate scope was achieved by altering functionalities with contrasting electronic and steric nature in the benzene ring of substrate. Generally high enantioselectivities were obtained with N-aryl-substituted anilines which decreased in case of N-alkyl-substituted anilines. This observation led to the development of a plausible transition state of the stereoselective electrophilic addition which included dual H-bonding interactions between both the substrates and the catalyst along with π–π interactions between the catalyst’s aryl group and the aryl substituent at the nitrogen in the aniline (Scheme 16a) [43]. Recently, Fan and co-workers reported a
Scheme 16: Anilines and α-naphthols as potential nucleophiles.

In 2020, Fu and co-workers developed a novel aza-Friedel–Crafts reaction between 3-arylindoles 68 and 2-aryl-3H-indol-3-ones 69 activating the C2–H bond of the heteroaromatic ring. Chiral phosphoric acid P12 catalyzed this transformation generating a complex molecular topology of 2,3-disubstituted indoles bearing both axial and central chirality. The aza-Friedel–Crafts reaction would allow the nucleophile to selectively attack the C=N plane of the electrophile as directed by a triple hydrogen-bonded complex between the catalyst and the substrates (see transition state 75, Scheme 18). This C–C bond formation affords a 3-indolinone moiety bearing an aza-quaternary stereocenter at the C2 position. In addition, the reaction allows to obtain axially chiral products 70/72/74 through restriction of the C–C bond rotation around the heteroaryl and aryl moieties. For this purpose, sterically bulky substituents need to be present in the aryl ring attached to the C3 position of the starting indoles. The axial chirality was attributed to ester and phenolic OH groups at the ortho-positions of the aryl ring and an additional phenolic OH functionality at the meta-posi-

chiral phosphoric acid P20-assisted enantioselective aza-Friedel–Crafts reaction between α-naphthols 17 and isatin-derived ketimines 49 to construct an aza-quaternary stereocenter at the C3 position of oxindole scaffolds 61 bearing a β-naphtholyl substituent (Scheme 16b) [44].

In 2020, Meng, Chan, Zhao and co-workers reported another C3-selective aza-Friedel–Crafts reaction of 4-aminoindole derivatives 63 utilizing N-Boc-α-ketimino esters 62 as potential electrophiles. The chiral phosphoric acid P21 catalyzed this process facilitating the formation of a quaternary stereocenter containing α-amino esters. Switching the solvent from non-polar to polar showed a regioselectivity shift to a C7 alkylation of the indole ring. The solvent-controlled regioselectivity switch of this aza-Friedel–Crafts reaction can be explained by the involvement of the polar solvent (acetonitrile) in the H-bonding with the catalyst thus creating a more hindered environment for a C3 alkylation, rather favoring the reaction through the less congested site (see transition states 66 and 67, Scheme 17) [45].
tion (substrate 68). Some more substrates were prepared by introducing a 2,5-diiodo-3,6-dihydroxyphenyl substitution at the C3 position of the indole ring (substrate 71). The products were formed with high chemical yields and excellent diastereo- and enantioselectivities. A further expansion of the substrate scope was demonstrated by incorporating a β-naphthol ring as the C3 substituent of the indole moiety (substrate 73). In all classes of bi(heteroaryl) substrates, a phenolic OH group at the ortho-position was crucial as it was involved in an intermolecular hydrogen bonding with the carbonyl oxygen of 69 in the ternary complex, thus bringing more rigidity in the three-dimensional transition state (Scheme 18) [46].

In 2021, Chen and co-workers documented a chiral phosphoric acid P17-catalyzed aza-Friedel–Crafts process between racemic 2,3-dihydroisoxazol-3-ol derivatives 76 and pyrroles/indoles 4/9 allowing access to 2,3-dihydroisoxazoles 77/78 bearing an all-substituted stereocenter at the C3 position. A dual catalytic activity of the Brønsted acid catalyst was illustrated by the authors which was initiated with a smooth protonation of the OH group in 76 with a subsequent dehydration to generate isoxazolium cation 80 paired with a phosphate anion. This chiral phosphorus is engaged in H-bonding with the free NH of the heteroarene ring to ease the stereoselective 1,2-addition to in situ generate the cationic heterocyclic scaffold 81. The reaction proceeded faster with pyrroles than with indole (Scheme 19) [47].

In 2021, Zhang and co-workers used 5-aminoisoxazole scaffolds 82 in an enantioefficient aza-Friedel–Crafts reaction with isatin-derived N-Boc ketimines 49. A 2-oxindole-substitutedaza-quaternary stereocenter was installed at the C4 position of the heteroaromatic ring in 83 and the enantioregulation was achieved by BINOL-derived chiral phosphoric acid P22. An amine functionality was crucial in the isoxazole ring to enhance the nucleophilicity of the adjacent carbon atom. In addition, the amine hydrogen forms an H-bond with the catalyst along with another hydrogen bond formed between the imine nitrogen of 49 and the catalyst’s OH group (see transition state 84). These dual H-bonding interactions were assisted by a π–π interaction between the aren rings of both the electrophile and nucleophile that helped in the formation of a stereodefined transition state. The substrate scope was achieved by varying the substituents in the C3 position of the isoxazoles 82 and the carbocyclic ring substituents in ketimines 49. Few more products were added to the library by altering the substituents of the amine in 82 and the ring nitrogen in 49 (Scheme 20a) [48]. The nucleophilicity of C3-substituted 5-aminoisoxazoles 82 was further utilized in another aza-Friedel–Crafts reaction with β,γ-alkynyl-α-ketimino esters 86 to provide N-Boc α-amino esters containing a quaternary stereocenter at the α-carbon. The chiral phosphonic acid P22 was used as catalyst to introduce the aza-ester quaternary stereocenter in the molecular entities 87 with appreciable chemical yields and excellent enantioselectivities. One
example was presented with a 5-aminoisothiazole motif that gave the product with much decreased yield (70%) and enantiomeric excess (36% ee) (Scheme 20b) [49].

In 2022, Sun, Li and co-workers developed an aza-Friedel-Crafts technique involving 3-alkynylated 3-hydroxy-1-oxoisoindolines 88 as electrophiles in combination with unsubstituted indoles 4 in the presence of chiral phosphoric acid ent-P17 as the catalytic agent. Facile dehydration of 88 was facilitated by the Brønsted acid to generate (N-acyl)(propargyl)imine 90 as intermediate which added to the deprotonated phosphoric acid to form phosphate ester 91 as the next intermediate through an equilibrium process. Then, 1,2-addition by the C3 position of the heteroarene ring to the acylimine intermediate afforded the
Scheme 19: Reaction between indoles and racemic 2,3-dihydroisoxazol-3-ol derivatives.

Scheme 20: Exploiting 5-aminoisoxazoles as nucleophiles.
3-indolyl-substituted aza-quaternary stereocenter. Here the stereoselectivity was attributed to an H-bonding interaction between the catalyst and the substrates (Scheme 21) [50].

In 2022, Lin and co-workers reported an unusualaza-Friedel–Crafts reaction using N-aryl-5-aminopyrazoles 92 as potentialπ-nucleophiles in combination with β,γ-alkynyl-α-imino esters 93 acting as the electrophilic reagent. Chiral phosphoric acid P16 was the catalytic agent to access a series of enantioenriched α-amino esters 94 containing 5-aminopyrazolyl and alkynyl substituents at the α-carbon. A library of products was prepared by varying different parts of both nucleophile and electrophile. The enantioselectivity of the reaction was an obvious result of a dual H-bonding interaction between the catalyst and both substrates where the imine nitrogen of 93 acted as H-bond acceptor and the amine functionality in 92 as H-bond donor to the catalyst (see transition state 97, Scheme 22a) [51]. Recently, the same research group documented another aza-Friedel–Crafts reaction between indoles 4 and 95 that frames aza-quaternary stereocenter at the α-carbon.
of unnatural amino acid derivatives 96. Enantiocontrol was rationalized by dual H-bonding interactions between both the reagents and the catalyst. The indole’s NH performed as the H-bond donor whereas the imine nitrogen of 95 was the H-bond acceptor towards the catalyst enabling a face-selective attack by the π-nucleophile to the electrophile C=N plane (see transition state 98). The substrate scope comprised mainly varying aryl or heteroaryl-substituents at the alkyne moiety that imparted high degrees of enantioselectivities to the products (Scheme 22b) [52].

In 2022, Huang and co-workers demonstrated an atroposelective construction of 3,4'-indole-pyrazole frameworks achieved through an asymmetric aza-Friedel–Crafts reaction. As substrate the authors chose the racemate of indole moiety 99 bearing a 5-acetoxy pyrazol substitution at the C3 position which was coupled with the pyrazolone-derived imine 100 to functionalize the C2–H bond of the indole ring. This aromatic electrophilic substitution also gave a quaternaryaza-stereocenter in the pyrazolone moiety. Axial chirality associated with central chirality in the product structures was influenced by chiral phosphoric acid catalyst P23. To freeze the C–C bond rotation, the pyrazole moiety in 99 required sterically demanding substituents. Excellent dia- and enantioselective synthesis of the products were caused by a chiral environment induced in the transition state through a dual H-bonding interaction between both the substrates and catalyst. In addition, π–π stacking between the aromatic moieties in both reagents brought more rigidity in the corresponding transition state (Scheme 23) [53].

In 2023, a chiral phosphoric acid ent-P17-mediated aza-Friedel–Crafts alkylation was reported between 5-aminopyrazole 92 as the π-nucleophile and 3H-indol-3-ones 69 as electrophilic reagents. The presence of an amino group in pyrazole 92 is necessary as it is engaged in the H-bonding interaction with the catalyst P=O moiety whereas the imine nitrogen of 69 accepts an H-bond from the catalyst OH group (see transition state 103). These dual noncovalent interactions were the reason behind a highly face-selective attack by the ortho-carbon of the aromatic amine functionality to the cyclic imine allowing a facile access of indolin-3-ones 102 attached to a 5-aminopyrazoly-substituted aza-quaternary stereocenter via the C2 position. The reaction was very well compatible with various aryl substituents as well as different groups on the benzene ring of indolones 69. Further broadening of the substrate scope was achieved by changing the aryl substituent attached to the pyrazole ring nitrogen. For enantioenrichment of the products, the presence of a methyl group at the C3 position of the pyrazole ring was obligatory. One example was included with a phenyl substituent at the aforesaid position for which a much diminished enantioselectivity (44%) was obtained (Scheme 24) [54].

Pyrophosphoric acids

In 2018, Ishihara and co-workers demonstrated a highly para-selective aza-Friedel–Crafts process using phenols and ortho-monosubstituted phenol analogues 104. As potential electrophiles, N-methoxycarbonyl-substituted aldimines 105 were explored to activate the para-carbon of the phenol derivatives catalyzed by the chiral pyrophosphoric acid Py1. The high regioselectivity was mainly caused by catalyst–substrate interactions via intermolecular H-bonding which could force the π-nucleophile to approach from the less sterically congested para-position. As ortho-substituents in the phenol derivatives, mainly sterically bulky alkyl, silyl, and iodo groups were incorporated to ensure the complete regioselectivity. On the other hand, various aromatic aldehyde-based aldimines were examined as electrophilic partners. Enantioincorporation into the products was explained by a Si-face attack of the nucleophile to the C=N plane. However, this process was not very promising in terms of enantioselectivities (Scheme 25a). The synthetic ap-
Thioureas and squaramides

In 2018, Yang, Deng and co-workers developed an aza-Friedel–Crafts aminoalkylation of 4- and 5-hydroxyindoles [55].

The applicability of this asymmetric process was shown by synthesizing 110, a key intermediate of (R)-bifonazole (Scheme 25b) [55].
As electron-demanding component, N-Boc pyrazolinone ketimines 100 were investigated to install the all-substituted aza-quaternary stereocenter at the C4 position of the pyrazoline scaffold. Stereinduction on this chiral center was regulated by the chiral squaramide catalyst S1 affording the products with excellent enantioselectivities. A stereodefined transition state organized by triple H-bonding interactions between the catalyst and the substrates controls the enantioefficiency of this process (see transition state 114). The substrate scope was broader with 4-hydroxyindoles to functionalize the C5–H bond whereas a bit narrower substrate scope was achieved with 5-hydroxyindoles allowing the 4-indolyl-substituted stereocenter formation. In both cases, few more products were added by altering N1 and C3 substituents of 100 (Scheme 26) [56].

In the same year, a quinine-derived chiral thiourea-mediated aza-Friedel–Crafts reaction between hydroxyquinolines 115 and isatin-derived ketimines 49 was reported by Vila, Pedro and co-workers. Regioisomeric hydroxyquinolines were tested in this reaction to facilitate the electrophilic aromatic substitution on the ortho-carbon atom with respect to the hydroxy group in quinolines 15. The reaction affords oxindole scaffolds 116 with a hydroxyquinoline-substituted aza-quaternary stereocenter in the 3 position. Most of the examples in this report involved 6-hydroxyquinoline as nucleophile whereas two examples each were presented with 5- and 7-hydroxyquinolines, respectively. Both the imine nitrogen and the carbonyl oxygen of the N-substituted Boc group of 49 were H-bonded with NH groups of the thiourea framework whereas the hydroxy functionality of 116 engaged itself in H-bonding with the quaternary nitrogen of the catalyst (see transition state 117). These noncovalent interactions were responsible for the stereochemical output of the reaction furnishing the products with moderate to excellent enantioselectivities. Electronically and sterically divergent functionalities in the benzene ring of 49 expanded the substrate scope whereas variation of 115 was very much limited (Scheme 27) [57].

In 2021, Wang and co-workers developed an azafriedel–Crafts reaction involving β-naphthols 119 as π-nucleophiles and benzothiazolimines 118 as electrophiles. Chiral squaramide S1-assisted this process affording enantioenriched 1-((benzothiazol-2-ylamino)methyl)naphthalen-2-ols 120 with high chemical yields. The activation of the electrophile was achieved through acceptance of H-bonds by the nitrogens in 118 from the NH moieties of the catalyst where a free OH group of 119 donated a H-bond to the tertiary amine moiety of S1. These noncovalent interactions were responsible for the stereochemical output of the reaction. Different aryl substituents on the imine carbon and functionalities in the carbocyclic ring of 118 were tested. One example was shown with an alkyl-substituted imine which provided the product with much
decreased enantioselectivity (45% ee) and four examples were presented by varying the functionalities in the nucleophile (Scheme 28) [58].

In 2021, Wang, Jin and co-workers deployed chiral thiourea T2 as the catalytic agent for executing a highly enantioselective aza-Friedel–Crafts process between β-naphthols 119 and isatin-derived ketimines 49 in the course of accessing enantio-enriched 3-amino-2-oxindoles 122 (Scheme 29) [59].

Other catalysts
In 2019, Vila, Pedro and co-workers reported a functional group-directed activation of the carbocyclic ring of indoles utilizing cyclic imines as electrophiles. The quinine-derived...
compound O1 was the catalytic reagent to functionalize the ortho-C–H bond of 4-, 5-, and 6-hydroxyindoles 111 via an aza-Friedel–Crafts aminoalkylation involving benzoxathiazine 2,2-dioxides 12 as electron-demanding reagents. H-Bonding engagement of both substrates with the catalyst selectively masked the Re face of the imine plane thus forcing the nucleophile to approach from the Si face (see transition state 124, Scheme 30) [60].

In 2019, Zhou and co-workers reported an aza-Friedel–Crafts reaction between α-naphthol derivatives 17 utilizing 7-membered cyclic N-sulfonylimines 125 as electrophiles leading to the facile access of ε-sultams 126 bearing a sulfonylamino-substituted stereocenter. Cinchona alkaloid O2 was the efficient catalyst for this asymmetric C–C bond formation delivering the products with moderate to good enantioselectivities. One example was documented involving β-naphthol as nucleophile and another example included electron-rich phenol (Scheme 31) [61].

Lin, Duan and co-workers demonstrated an enantioselective aza-Friedel–Crafts reaction between indoles 4 and isatin-derived ketimines 49. A chiral phase transfer catalyst O3 derived from urea assisted this organic transformation featuring a C3–H bond functionalization of indoles. Different protecting groups for the imine nitrogen and ring nitrogen of 49 were
screened under optimal reaction conditions where Cbz and benzyl were the best protecting groups in terms of enantioselectivities. A product library was prepared by varying sterically and electronic divergent functionalities in the carbocyclic rings of both reactants. Enantioincorporation into the products was explained by H-bonding engagement between the catalyst NHs groups and an ionic interaction between the anionic indole and quaternary ammonium moiety of the catalyst (Scheme 32) [62].

In 2022, Li, Chen and co-workers employed the BINOL-derived chiral disulfonimide $O_4$ as Brønsted acid catalyst to execute a straightforward aza-Friedel–Crafts reaction between 3-substituted indoles $4$ and $N$-sulfonyl-substituted aldimines $128$. The reaction successfully installed an aza-tertiary stereocenter at the C2 position of the heterocyclic ring. A broad substrate scope was investigated by varying substituents on both substrates. A transition state involving dual noncovalent interactions between the catalyst and substrates directed the face-selective addition of the $\pi$-nucleophile to the electrophilic carbon of the imine (see transition state $130$, Scheme 33) [63].

**Heterogenous catalysts**

In 2020, Pedrosa and co-workers devised a chiral heterogenous thiourea catalyst that was applied in an enantioefficient aza-Friedel–Crafts process. A series of heterogenous catalysts were prepared by condensation between alkaloids and polystyrene-derived isothiocyanates. These polymer-supported materials were utilized as heterogenous catalyst to execute the aza-Friedel–Crafts reaction between 1-naphthols $17$ and isatin-derived ketimines $49$ to produce oxindole motif $61$ bearing a 1-hydroxy-naphth-2-yl-substituted aza-quaternary stereocenter at the C3 position. The best result was obtained with the hydroqui-
nine-based supported catalyst H1 which efficiently promoted five catalytic cycles without loss of its activity. N-Alkyl-substituted ketimines 49 with different functionalities in the benzene ring were well responsive towards the heterogenous reaction to afford the products 61 with moderate to excellent enantiomeric excesses. However, N-unsubstituted 49 (R1 = H) resulted in a much diminished stereoselectivity. As the electrophilic partner, isatin-derived ketimine was also utilized which furnished the product with 68% enantiomeric excess. Replacement of the nucleophile in this methodology for substrate scope expansion was carried out by employing 2-naphthol and 4-hydroxyindole (Scheme 34) [64].

Application in total synthesis

In 2018, a guanidine bisthiourea-catalyzed highly enantioselectiveaza-Friedel–Crafts reaction was applied as a central step in the total synthesis of (+)-gracilamine. The reaction was designed between sesamol (132) and N-Boc-protected ketimine 131 in the presence of T3 as catalyst to introduce the electrophile at the ortho-position with respect to the phenolic OH group. The aza-Friedel–Crafts product was obtained with 94% yield and converted into triflate 133 with 74% yield and 99% ee after recrystallization. Subsequent ozonolysis of the terminal alkene functionality with a follow-up reduction furnished primary alcohol 134 which was transformed into the azide 135. Reduction of the azide 135 was accompanied by debenzylation, was followed by tosylation of the primary amine and exchange of the Boc-protecting group with the Tosc group then gave phenol 136. Compound 136 was then subjected to a highly diastereoselective oxidative phenolic coupling giving fused tetra-cyclic architecture 137. Follow-up acid-mediated intramolecular aza-Michael addition and subsequent alkene reduction provided ketone 138 which was reacted with an α-keto ester in an intramolecular 5-endo-trig-cyclization process to afford 139. Treatment of compound 139 with sodium borohydride afforded secondary alcohol 140 which after conversion of the tosyl group into a methyl group gave the final product 141 (Scheme 35) [65].

In 2019, Piersanti and co-workers reported an organocatalyzed enantioselectiveaza-Friedel–Crafts/lactonization domino reaction sequence as the key step in the course of synthesizing (+)- and (-)-fumimycin. (-)-Fumimycin was first isolated from Aspergillus fumisynnematus and exhibits antibacterial activity against resistant S. aureus strains. It is also an inhibitor of the enzyme peptide deformylases (PDFs). The synthesis comprised the reaction between the highly substituted hydroquinone 142 and dehydroalanine 143 in the presence of chiral phosphoric acid P7 as catalyst to prepare benzofuran-2(3H)-one derivative 144 having an aza-quaternary stereocenter. The achiral Lewis acid tris(pentafluorophenyl)borane was required as additive in the reaction system to enhance the chemical yield and enantioselectivity. After two additional steps, i.e., demethylation of the phenolic ether and ester hydrolysis, (-)-fumimycin (146) was obtained (Scheme 36) [66].

Conclusion

The aza-Friedel–Crafts reaction is a powerful reaction that allows the incorporation of an aminoalkyl functionality in aromatic systems through C–C bond formation. This C–H bond functionalization methodology of aromatic systems also has the possibility of incorporatingaza stereocenters into a product depending upon the choice of a suitable electrophile, i.e., imines. The present review assembled recent (from 2018 till date) examples of asymmetric versions of this important method mediated by different organocatalysts. The mechanistic approaches with explanation about the origin of stereoselectivities has also been elaborated. This reaction has been successfully utilized as the key step in the syntheses of different important natural products which have been included in this article as well. On searching the literature, it has been found that mainly H-bonding chiral organic molecules have been envisaged as the catalytic systems for stereoinduction into products. The asymmetric induction is caused by effective noncovalent interactions between the catalysts and substrates to force a face-selective attack by the nucleophile, i.e., the aromatic π-system to the electrophile.
Scheme 35: Total synthesis of (+)-gracilamine.

Scheme 36: Total synthesis of (−)-lumimycin.
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The definitive version of this article is the electronic one which can be found at: https://doi.org/10.3762/bjoc.19.72