Advances in the site-selective C-5, C-3 and C-2 functionalization of chromones via sp$^2$ C–H activation

Anjitha Theres Benny and Ethiraj Kannatt Radhakrishnan

In this work, site-selective C–H activation at C-5, C-3 and C-2 positions of chromones for the introduction of structural diversity to the chromone scaffold was studied. The keto group of the chromone moiety acts as the directing group for the selective functionalization of chromones at the C-5 position. Furthermore, the C–H functionalization at the electron-rich C-3 position of the chromone can be achieved using electrophilic coupling partners. The C–H functionalization at the C-2 position can be possible using nucleophilic coupling partners. The direct functionalization methods provide a better pathway for the generation of C-5, C-3 and C-2-substituted chromones with good atom economy than that of classical pre-functionalized reaction protocols.

1. Introduction

Chromone 1 scaffolds are ubiquitously present in numerous biologically active compounds and, hence, received considerable attention from synthetic chemists in discovering new drugs.$^1$ The importance of chromones in drug discovery and synthetic transformations is evident from a number of publications in the field. Chromones are well documented in the literature owing to their interesting biological applications such as anticancer,$^2$–$^5$ antimicrobial,$^6$–$^{10}$ anti-inflammatory,$^{11}$–$^{14}$ antioxidant,$^{15}$–$^{18}$ anti-Alzheimer,$^{19}$–$^{22}$ and antidiabetic$^{21}$–$^{26}$ activities. Apart from their biological applications, they show promising fluorescent activity, and hence can be used as sensors in detecting various metal ions and in bioimaging.$^{27}$–$^{34}$ The promising biological and photophysical activities of chromones make them a valid scaffold of interest among chemists.

The area of interest in this review covers the site-selective functionalization of chromones in C-5, C-3 and C-2 positions. Few of the C-2, C-3 and C-5-functionalized chromones with biological applications including anti-asthmatic$^2$,$^{35}$ antimalarial$^3$,$^{36}$ anticancer$^4$,$^{37}$ antiviral$^5$,$^{38}$ antimicrobial$^6$–$^9$,$^{39}$–$^{40}$ anti-inflammatory$^{10}$,$^{41}$ antiplatelet$^{11}$,$^{42}$ antioxidant$^{12}$,$^{43}$ and anti-Alzheimer$^{13}$ activities are given in Fig. 1.

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Direct C–H bond activation/functionalization has practical and fundamental challenges but it is a highly attractive strategy in the formation of a new functionality or a new C–C bond. The substrate scope by C–H activation is unlimited, including hydrocarbons, complex organic compounds of small molecular weight, and synthetic and biological polymers.55–57

C–H functionalization of chromones is a useful method for the construction of carbon–carbon and carbon–heteroatom bonds. The cross dehydrogenative coupling (CDC) for the formation of Csp²–Csp, Csp–Csp² and Csp³–Csp bonds enables the synthesis of significant building blocks from a simple chromone moiety.56 The C-5 functionalization of chromones by transition metal-catalyzed reactions is reported through chelation-assisted metal-catalyzed reaction transformation. The method proceeds by the effect of the weakly coordinating ability of the keto group on the chromone to the metal catalyst forming a metallacycle.55,58 Similarly, the C-3 position of the chromone is functionalized by a transition metal-catalyzed reaction through a nucleophile attack of the C-5 site of chromones with various olefins (Scheme 1) was carried out in an atom-economic, highly efficient, and concerted reaction pathway.

Mechanistically, the transition metal-catalyzed directing group (keto group)-assisted dehydrogenative coupling starts with the C–H bond activation (Scheme 2). Initially, the chloride ligand of the complex [(Rh(III)Cp*Cl)₂] will be removed as AgCl by the action of AgSbF₆ to generate an active rhodium complex. The active rhodium species thus generated may contain ligands such as acetate or solvent in addition to pentamethylcyclopentadienyl (Cp*) ligands. Then, the active Rh species coordinates with the keto group of chromone and a concerted metalation/deprotonation (CMD) process activates the ortho C–H. The CMD process yields a five-membered rhodacycle 14 intermediate. The intermediate 14 transforms to a seven-membered rhodacycle 16 via an alkene insertion to the Rh–C bond of 14. The β-H elimination from the seven-membered intermediate afforded desired products 17(a–f) and the reduced inactive [RhCp*²] complex. The catalytic cycle was completed by the oxidation of inactive [RhCp*] to [Rh(III)Cp*] by copper acetate.55

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2. C-5 activation of chromones

2.1. Transition metal-catalyzed C-5 activation of chromone

R. Samanta et al. in 2012 introduced Rh(III)-catalyzed C–C bond formation without a pre-functionalized reaction protocol for the synthesis of hybrid molecules of synthetic and medicinal importance. The direct oxidative cross-coupling reaction at the C-5 site of chromones with various olefins (Scheme 1) was carried out in an atom-economic, highly efficient, and concerted reaction pathway.

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applicable for the selective perfluoroalkenylation of chromones at its C-5 position. The reaction proceeds under a mild condition with low catalyst loading and very good functional group tolerance and good to excellent yield. The synthetic scope of the method was enhanced by the excellent reactivity of the coupling partners and the regioselectivity of the reaction.

The reaction proceeds by initial electrophilic deprotonation of chromone 1 forming an Ar–Rh(III) intermediate 18. The coordination of the intermediate with CF₃-substituted acrylate 19 then activates the olefin for further oxidative addition and β-hydrogen elimination to afford the expected product 22.⁵⁵

Minsk Min et al. 2014 put forward stereodivergent hydroarylation of chromones with alkynes (Scheme 5). The reaction proceeds in the presence of catalytic amounts of [Ru(p-cymene)Cl₂]₂ and AgSbF₆ was a diastereodivergence in the reaction achieved by the quantitative loading of AgSbF₆. The controlled quantitative addition of AgSbF₆ to the reaction led to the switching of E/Z isomers in the product instead of the sole E-isomer.

Ru(n) for the reaction was generated from [Ru(p-cymene)Cl₂]₂ upon treatment with AgSbF₆ (Scheme 6). A coordinative insertion of alkyne to intermediate 23 yields seven-membered ruthenacycle intermediate 25. Subsequent protonolysis of Ru–C bonds of 25 affords E-alkene 27 regenerating the Ru(n) catalyst, and the rotation of the bond generates thermodynamically stable Z-isomer 28.⁶⁶

Kiho Kim et al. 2015 studied the direct C–H oxygenation of chromones using a Ru(n)-catalyzed synthetic protocol for the installation of a hydroxyl group on the C-5 position of chromones (Scheme 7). The oxygenation of chromones was achieved in good yield in the presence of monomeric ruthenium complex [Ru(p-cymene)(CF₃CO₂)₂(H₂O)].
Chelate-directed C–H bond activation of chromone generates five-membered ruthenacycle 32, which on oxidation with iodine affords a Ru(IV) 33 species. A subsequent reductive elimination of 33 yields 5-(trifluoroacetyloxy)-avone 34 which on aqueous workup yields the corresponding C-5-functionalized chromones (Scheme 8).

Rhodium(II)-catalyzed C–H functionalization of bi- or tricyclic scaffolds such as chromones and xanthones was carried out by Sang Hoon Han et al. in 2016 (Scheme 9). This biologically active compound upon site-selective functionalization with maleimide yields a succinimide substitute at their respective C-5 position with good functional group compatibility and excellent site selectivity. The succinimide generated in the C-5 position of the chromones was obtained by olefin insertion into the metallacycle reaction intermediate followed by a protonation step.68

The C–C bond formation with metal-catalyzed C–H activation of aryl ketones was studied by R. Shang et al. 2016 (Scheme 10). The main attraction of the reaction is the usage of inexpensive methylaluminum as the methylating agent and the economic and environmental merits of iron catalysts. The reaction proceeds by the combined activity of 4-(bis[2-(diphenylphosphanyl)phenyl]phosphanyl)-N,N-dimethylaniline (Me2N-TP) and methylaluminium for the transformation of ortho C–H to C–CH3 under an iron-catalyzed reaction condition. The reaction proceeds well for various functional groups.69

1,2-Diacylbenzene as a valuable precursor for the synthesis of medicinally important compounds such as isobenzofurans, phthalazines, and isoindoles can be generated by decarboxylative acylation of aromatic ketones via a palladium-catalyzed reaction (Scheme 11). The decarboxylative acylation reaction yields 1,2-diacylbenzenes in excellent yields with ortho selectivity. This reaction protocol put forward by Wing-Yiu Yu et al. 2017 demonstrates a successful attempt for the direct C–H acylation reaction of aromatic ketones with ω-oxocarboxylic acids, which proceeds without pre-derivatization to imines.67
The plausible reaction mechanism for the acylation of chromones under a palladium-catalyzed reaction condition is depicted in (Scheme 12). First, Pd(OAc)$_2$ gets transformed to Pd(TFA)$_2$ in situ by the action of trifluoroacetic acid (TFA). The acylation gets initiated by the directing group (keto) via ortho-selective electrophilic palladation on the olefin by Pd(TFA)$_2$. Palladacycle 43 obtained will then undergo oxidative coupling with acyl radical 44a generated in situ by decarboxylation of a-oxocarboxylic acid. Reductive elimination from Pd(III) or Pd(IV) intermediate would then furnish the final product 1,2-diacylbenzene 46 and an active Pd(II) catalyst.

Ir(III)-catalyzed selective C–H bond alkynylation is a useful synthetic transformation of ketones and esters for the construction of molecular complexity and developing drug molecules. The alkynyl functionality introduced with the facilitation of the carbonyl group on ketones and esters was catalyzed by Ir(III) and Ag(I) (Scheme 13). The feasibility of the reaction with both electron-donating and electron-withdrawing substituents was investigated. The yield of the reaction implies the feasibility of electron-rich species over electron-deficient substituents. The substrate scope was also extended with halogen-substituted aromatic rings, which may increase the scope of reaction transformation further. The terminal alkynes were afforded using the desilylation reaction under mild reaction conditions.

Li, X et al. 2017 explained the mechanism for the alkynylation reaction of chromones, as depicted in Scheme 14. The reactive Ir(III) is initially generated from [Cp*IrCl$_2$]$_2$ and AgNTf$_2$ via ligand exchange. Later, the carbonyl-anchored Ir(III) facilitates the bond cleavage of C-5–H with the aid of acetate ligands. The reaction yields cyclometalated Ir(III) complex 49 the key intermediate of the reaction. The Ir(III) complex obtained undergoes oxidation with Ag(i) and alkynyl bromide in a concerted step or a multistep pathway yielding the product along with the regenerated Ir(III) catalyst.

Direct C-5 arylation of chromones catalyzed by [IrCP*Cl$_2$]$_2$ for the synthesis of C-5-functionalized chromones was introduced by K. Shin et al. 2018 (Scheme 15). The reaction afforded C-5 aryl chromones in good yields with functional group tolerance at ambient temperature. The reaction proceeds via C–H cleavage at the C-5 position and subsequent transmetallation.
The transmetallated complex forms a high-valent aryl species by one-electron oxidation. The formation of this high-valent species reduces the barrier of reductive elimination by approximately 19 kcal mol\(^{-1}\). Finally, the oxidatively induced reductive elimination yields the C-5 arylated chromones.

Tan, G et al. 2018 discovered a [IrCp*Cl\(_2\)]-catalyzed C–H arylation of diverse arenes with heteroarenes in a modest yield (Scheme 16). The C–H heteroarylation of aromatic ketones and enamides was carried out by the Cp*Ir catalyst via oxidatively induced reductive elimination. The reaction protocol works well for the synthesis of aryl–heteroaryl and vinyl–heteroaryl scaffolds.

S. Debbarma further studied the wide applicability of chromones. S. Debbarma et al. in 2019 synthesised a set of C-5 substituted chromones by metal-catalyzed C-5 activation of chromones (Scheme 17). The ketone-directed C–H functionalization of C-5 carbon leads to the diversification of chromones by transition metal-catalyzed allylation, amidation, alkenylation, and 1,4-addition reactions.

A C-5 functionalization method for the synthesis of β-aryl- and β-heteroarylethenesulfonyl fluoride was carried out by G. Ncube et al. 2019. The reaction utilizes a catalytic system of tris(acetonitrile)pentamethylcyclopentadieny]rhodium(III) hexafluoroantimonate, [Cp*Rh(MeCN)\(_3\)], and (SbF\(_6\))\(_2\) by oxidative C–H alkenylation of arenes with ethenesulfonyl fluoride (Scheme 18). The directing group strategy allows the synthesis of β-aryl- and β-heteroarylethenesulfonyl fluoride without the pre-installation of halogens, boronic acids, or diazonium functional groups on arenes with modest to good yields for a broad variety of directing groups. The reaction strategy provides an excellent and harmonizing approach for the attainment of novel building blocks in sulfur(IV) exchange click chemistry.

Vinylarenes are structurally important scaffolds due to their applicability as a starting material for various fine chemicals, polymers, and drugs. The synthesis of vinylarenes with ketone-directed C–H vinylation using vinyl acetate was reported by M. R. Sk et al. 2020 (Scheme 19). The vinylation on a set of compounds including diaryls, acetophenones, chalcones, and chromones was carried out by first-row transition metals via ketone-directed C–H bond activation. This is the recently reported reaction for the cobalt-catalyzed carbonyl group-directed C–H vinylation with vinyl acetate.

The active catalyst for the vinylation reaction was obtained from [Cp*CoL\(_2\)] complexes in the presence of silver hexafluoroantimonate and acetate by removal of AgI (Scheme 20).
The active catalyst and aromatic ketone moiety form a cyclo-
metzallated species $69$. Subsequently, a 1, 2 migratory insertion
of Co–C bond into the vinyl acetate provides the intermediate
$72$. The β-hydride elimination/reinsertion of the intermediate
$72$ and the β-acetate elimination of the formed final interme-
diate $73$ gives the desired vinylated product $74$.$^{76}$

Y. Zhou et al. 2020 reported the Ru(II) catalyst for the
coupling of chromones with maleimides yielding a C-5-
activated chromone by utilizing keto as a weak directing

The mechanism of the reaction was studied by deuteration
techniques. Initially, the ruthenium complex $[\text{Ru}(p$-cymene)
$\text{Cl}_2]_2$ reacts with AgNTf$_2$ and benzoic acid or AgOAc accordingly
for the generation of an active monomeric form or $A'$ respec-
tively (Scheme 22). The oxygen of the keto group coordinates
with the active ruthenium species, which provides a five-
membered metallacycle $76$. Subsequent insertion of mal-
eimide to the carbon–ruthenium bond of $76$ gives a bicyclic

![Scheme 19](image)

**Scheme 19** Cobalt(II)-catalyzed C–H vinylation of chromones.

![Scheme 20](image)

**Scheme 20** Proposed mechanism for cobalt(III)-catalyzed C–H
vinylation of chromones.

![Scheme 21](image)

**Scheme 21** Ruthenium(II)-catalyzed C–H activation of chromones
with maleimides to synthesize succinimide/maleimide-containing
chromones.

![Scheme 22](image)

**Scheme 22** Proposed mechanism for ruthenium(II)-catalyzed C–H
activation of chromones with maleimides to synthesize succinimide/
maleimide-containing chromones.
intermediate 78. The addition of benzoic acid or silver acetate to the intermediate 78 yields the 1,4-addition product or Heck-type product respectively. The catalyst is regenerated as a Ru(II) species by the deprotonation of β-hydrogen of 78 and subsequent oxidation by silver acetate.77

2.2. Metal-free C-5 activation of chromones

The C-5 functionalization with the nitro group was carried out by Recanatini, M et al. in 2000 with the traditional nitration method using HNO3 in H2SO4 at 0 °C to room temperature to yield 8-methyl-5-nitro-4H-benzopyran-4-one (Scheme 23). The nitration compound was utilized for further reaction transformation.78

Tsukasa Ishihara et al. 2004 utilized base-catalyzed C-5 activation of xanthenone by allylbromide (Scheme 24). The C-5 alkylated compounds were synthesized using DMF with allyl bromide at room temperature in excellent yields. The alkylated compounds were further utilized for other transformations and biological studies.79

3. C-3 activation of chromones

3.1. Transition metal-catalyzed C-3 activation of chromones

C-3-functionalized chromones were synthesized by Jeganmohan, M et al. 2010 using a tmp4Zr·4MgCl2·6LiCl base in THF (Scheme 25). The base tmp4Zr·4MgCl2·6LiCl initiates the zirconation of heterocycles of pharmaceutical importance like chromones. Chromones upon reaction with the zirconium base at −35 °C yield zirconated chromones in excellent yields. The zirconated chromones upon copper-catalyzed allylation yields a Sn2 substituted C-3 allylated product.80

A direct C-3 alkenylation of chromones through a palladium(n) catalyzed methodology was achieved by D. Kim et al. in 2011 (Scheme 26). In this reaction, the reactivity of the coupling reaction between chromones and alkenes was attained by the combined activity of pivalic acid and Cu(OAc)2/Ag2CO3 systems. The cross-coupling method leads to the formation of 3-vinylchromones, which are good structural scaffolds for the generation of biologically and synthetically active compounds. The method represents a C-3 derivatization of chromones with considerable advances over the current two-step methods with a good functional group tolerance and remarkable yield.

The possible mechanism for the transformation of the chromones to 3-vinylchromones initiates with a nucleophile attack of chromones on palladium and the subsequent generation of palladium(n) intermediate 88 by the deprotonation by a pivalate ligand (Scheme 27). Next, the C-3 palladated species was inserted into the alkene substrate followed by reductive elimination, which generates the desired C-3 functionalized chromones. The catalyst Pd(n) was then regenerated from Pd(0) using Cu(OAc)2/Ag2CO3 to continue the catalytic cycle further.81

Oxidative arylation of chromones catalyzed by a Pd(n) catalyst was conducted by K. H. Kim et al. in 2012 (Scheme 28). The C3-functionalized product was then selected for the arylation reaction was C=2, and C=3 was unsubstituted, and hence, the reaction yields a trace amount of C3-functionalized product.82

F. Chen et al. 2012 investigated the regioselective intermolecular arylation of chromones with polyfluoroarenes using a palladium-catalyzed reaction protocol (Scheme 29). The reaction afforded fluoroarylated enones by a two-fold C–H bond functionalization method. The scope of the reaction extends to C3 fluoroarylation of quinolones, chromones and pyrimidones in reasonable yields. The proposed reaction mechanism for the cross-coupling reaction proceeds by the formation of a palladated intermediate of both enone and fluoroarene. The intermediates thus formed couple with fluoroarene and enone and the succeeding reductive elimination yields desired products along with the Pd(n) catalyst.83

The C-3 functionalization of flavones with tert-butylperoxbenzoate (TBPB)/potassium persulphate (K2S2O8) oxidant combinations catalyzed by an iron(n) catalyst was carried out by B. A. Mir et al. in 2016 (Scheme 30). A Csp2–Csp3 coupling product was afforded for the reaction of flavones in cycloalkanes with exclusive C-3-functionalized cycloalkylated products. The reaction afforded a C-3 amidation product when the
solvent was altered from cycloalkane to N,N-dimethylformamide. In both reactions, the solvent itself acts as a coupling partner for the progress of the reaction. The radical forming capability of cycloalkane and N,N-dimethylformamide ensures the formation of a product with the active participation of solvents in reactions. Again, when the solvent was changed to chlorobenzene, the C-3 methylated product was obtained by utilizing tert-butylperoxybenzoate (TBPB) as the source of the methyl group.

The C-3 functionalization reaction is set off by thermal homolytic bond cleavage of peroxide oxidant tert-butyl peroxybenzoate (TBPB) forming peroxybenzoate and tert-butoxy radicals. Both of these radicals generated are capable of scavenging hydrogen from cyclohexane 96 and N,N-dimethylformamide 98. A stable benzyl radical will generate upon the addition of any of these radicals on flavone 1. The radical species 110 generates a cationic intermediate 111 through the reduction of catalyst Fe^{3+} to Fe^{2+}. Finally, the C-3 proton of the flavone gets abstracted by the base affording the C-3-functionalized product. In the absence of radical-generating solvents, TBPB acts as the source of the CH₃ group. The method allows the synthesis of a C-3-functionalized product through cycloalkylation, amidation and methylation by varying the solvents** (Scheme 31).

Youngtaek Moon et al. 2012 put forward an efficient and straightforward methodology for the direct cross-coupling of

Scheme 27 Proposed mechanism for palladium(II)-catalyzed intermolecular C-3 alkenylation of chromones.

Scheme 28 Palladium-catalyzed C-3 arylation of chromones.

Scheme 29 Palladium-catalyzed dehydrogenative coupling of chromones with polyfluoroarenes.

Scheme 30 Iron(III)-catalyzed functionalization of the C-3 position of chromones.

Scheme 31 Proposed mechanism for iron(III)-catalyzed functionalization of the C-3 position of chromones.
the C-3 position of chromones with quinones using palladium acetate as the catalyst. The reaction afforded good to excellent yields of quinones with tolerance for a wide range of functional groups on both the chromosome and quinone moieties. Later, H. Choi et al. 2016 developed Pd(n)-catalyzed direct arylation of chromones (Scheme 32) by site-selective C-3 functionalization. The C–H activation progresses through a single-step metatillation deprotonation method, which selectively activates the C-3 position of chromones.85

The synthesis of a C-3 acylated chromosome via acylation of 2,3-unsubstituted chromones through decarboxylative C–C cross-coupling with α-keto acid was described by Satenik Mkrtchyan et al. 2018 (Scheme 33). The title transformation was effectively catalyzed by using a set of silver and palladium salts.

The mechanism for the C-3 acylation of chromones involves an initial O-acylated radical-chromone adduct formation (Scheme 34). Subsequently, the adduct formed get oxidised into an O-acyl pyrylium moiety, which further undergoes Fries-type rearrangement followed by deprotonation yielding the preferred product in moderate quantity.86

3.2. Metal-free C-3 activation of chromones

Lv, Z et al. 2010 studied the metal-free C-3–H activation of chromones (Scheme 35). The chromones were functionalized at their C-3 position by base-catalyzed iodination under reflux conditions. The iodochromones were further utilized for the generation of a terminal alkyl chain on the C-3 position.87

Lydia Klier et al. in 2012 and 2017 explained the Lewis acid accelerated zincation of chromones and thereby the transformation of the compound to other useful products (Scheme 36). TMPZnCl2·LiCl (TMP = 2,2,6,6-tetramethylpiperidyl) leads to the zinication of chromones and quinolones by coordination of TMPZnCl2·LiCl to the basic carbonyl oxygen. The zinication yields Zn-coordinated chromones through a complex induced proximity effect (CIPE). The zinication can be done at either the C-3 or C-2 position of chromones depending on the 2,2,6,6-tetramethylpiperidine (TMP) base. A similar zinication product of chromone can be obtained on the C-2 position of the chromone if the TMP base selected is TMP2Zn2·MgCl2·2LiCl. The resulting zinc organometallic intermediate undergoes various reactions including Pd/Cu-catalyzed cross couplings or acylation/allylation reactions.88,89

Y. Ding et al. 2016 put forward ammonium iodide-induced sulfenylation of flavones, arylimidazo[1,2-a]pyridines and indoles for the generation of thioethers (Scheme 37). The reaction proceeds with sodium benzenesulfonates as the sulphur source and odourless ammonium iodide as the reaction inducer. Sodium benzenesulfonate was selected as the sulfur source due to its stability and efficiency in synthesising regioselective products in good yields. The regioselectiveness of the reaction on flavones was studied by substitution with methyl groups at the C-3 and C-2 positions. Once the C-3 position was blocked by the methyl group, no thioether was
formed, verifying the regioselectivity of the sulfonylation reaction.\textsuperscript{90}

W. Zhao \textit{et al.} 2015 introduced arylthiols as a new sulfur source and the same colourless odourless ammonium iodide as the reaction inducer (Scheme 38). The sulfonylation method introduced here was an efficient and regioselective methodology for obtaining organosulfur compounds of flavones in good to excellent yields in a metal-free synthetic protocol.

At 135 °C, the reaction inducer ammonium iodide was split into NH\textsubscript{3} and HI (Scheme 39). The obtained HI gets oxidized into iodine in the air atmosphere. The generated iodine will then react with thiophenol to form an electrophilic species ArS–I. ArS–I upon reaction with flavone 1c at the electron-rich C-3 position gives the corresponding intermediate 128. The intermediate 128 on deprotonation yields the desired thioether 130.\textsuperscript{91}

Later in 2015, W. Zhao \textit{et al.} introduced another sulfonylation method with dimethylsulfoxide or benzenesulfonyl hydrazide as the sulfonylating reagents and ammonium iodide as the reaction inducer (Scheme 40). The reaction proceeds at elevated temperatures, leading to the splitting of ammonium iodide into HI and NH\textsubscript{3}. HI reduces the reaction inducers, DMSO and benzenesulfonyl hydrazide. The electrophilic intermediate obtained from DMSO easily reacts with nucleophilic flavone and subsequently attacks I and NH\textsubscript{3} to give the final thioether product from DMSO. Similarly, the benzenesulfonyl hydrazide counterpart undergoes dehydration followed by reductive dehydration supported by HI to give thiadiazonium. The thiadiazonium regioselectively attacks the nucleophilic flavone, yielding the corresponding thioether. The regioselectivity of the reaction was studied by blocking the C-3 position of flavone, yielding no desired product, which implicates the regioselectivity of the reaction.\textsuperscript{92}

T. Guo \textit{et al.} 2020 synthesised a C-3-functionalized chromone by the C-3 vulcanisation of imidazoheterocycle (Scheme 41). The dual C-H sulfonylation was achieved by odourless elemental sulphur under an additive and catalyst-free reaction condition. The C-3-functionalized asymmetric sulphides were obtained in good yields with C-H activation of both chromones and imidazoheterocycles using DMSO as both solvent and internal oxidant. The method acts as an efficient methodology for the synthesis of different organosulfur compounds using commercially available S\textsubscript{8}.\textsuperscript{93}

4. C-2 activation of chromones

4.1. Transition metal-catalyzed C-2 activation of chromones

K. H. Kim \textit{et al.} in 2012 carried out the synthesis of C-2 arylated chromones by palladium-catalyzed oxidative arylation in
benzene (Scheme 42). The reaction proceeds with regioselective C-2 arylation of chromones via a double C–H activation process. The reaction yields C-2 arylated chromones in moderate yields through regioselective palladation on the C-2 position of chromones via a CMD process and a subsequent C–H bond activation. Similarly, M. Min et al. 2012 put forward the regiocontrolled C–H activation at the C-2 position of chromones. The method enables the palladium-catalyzed oxidative cross coupling reaction of nonactivated arenes with the C-2 position of chromones. The reaction offers a direct route to the C-2 arylation under mild conditions with a wide substrate scope.

The mechanism proceeds via regioselective palladation at the C-2 position of chromones via a CMD process with 138 for the generation of intermediate 139. A subsequent C–H bond activation of benzene could occur via CMD mechanism or an electrophilic palladation (SEAr) process to generate the intermediate 141. The intermediate 141 on the removal of palladium (0) yields the final C-2 arylated product. Furthermore, Pd(0) was oxidized to Pd(II) by the action of AgOAc for the continuation of the catalytic cycle (Scheme 43).

D. Kim et al. in 2012 reported the C-2 functionalization of chromones via 1,4-conjugate addition of aryl boronic acid with substituted chromones. The reaction yielded a mixture of flavones and flavanones in the presence of the catalytic amount of Pd(OAc)₂ and Lewis acid, whereas the addition of the catalytic amount of oxidants DDQ and KNO₂ to the reactions solely yielded flavones (Scheme 44). The scope of the reaction was fairly well giving good yields for a wide range of flavanones and flavones.

The plausible mechanism for the direct arylation of chromones at the C-2 position by boronic acid is depicted in Scheme 45. Initially, Fe(OTf₃) coordinates with the carbonyl oxygen of the chromone moiety, thereby activating the electrophilicity of enolone of chromones. The phenylpalladium formed in situ via transmetalation of the aryl group of supplied boronic acid adds to the C-2 position of chromones forming alkylpalladium adduct 146 and enolate 147. Further protonolysis of the formed enolate yields the conjugate addition product flavanone. Similarly, β-elimination of the palladium adduct 146 generates a Heck-type product flavone as a minor product. The usage of the oxidant system DDQ/KNO₂ provides flavone exclusively via oxidation of the resulting flavanone. Finally, the reoxidation of Pd(0) to Pd(II) completes the catalytic cycle.

B. Niu et al. 2015 introduced copper-catalyzed dehydrogenative coupling of coumarins and flavones with various ethers yielding new substituted ethers (Scheme 46). The reaction proceeds in the presence of a catalytic amount of copper oxide in combination with tert-butylhydrogen peroxide (TBHP) and 1,4-diazabicyclo[2.2.2]octane (DABCO) in good yields with excellent functional group tolerance.

The cross-dehydrogenative coupling of chromones with ethers initiates the formation of two radicals under thermal conditions.
conditions (Scheme 47). The radicals abstract hydrogen from the α position of supplied ether. The relatively electron-rich ether radical 150 thus forms the electron-deficient C-2 position of flavones. The radical species 151 thus generated upon oxidation with a copper catalyst yields C-2-functionalized products.

B. M. Trost et al. 2019 reported the enantio- and diasteroselective 1,4-addition of butenolides to chromones (Scheme 48). The zinc-prophenol catalyst was compatible for both α,β- and β,γ-butenolide nucleophiles without any preactivation. The reaction proceeds with moderate to good yields for electrophiles including chromones, thiochromones and quinolones.97

4.2. Metal-free C-2 activation of chromones

R. Narayan et al. in 2014 reported hypervalent iodine-mediated selective oxidative functionalization of chromones by aliphatic alkanes proceeds well in both cyclic and acyclic alkanes (Scheme 49). They suggested that iodine used in the reaction generates alkyl radicals in situ, which will attack the most electrophilic C-2 position of chromones. The presence of the catalytic amount of PhI(O2CCF3)2 and NaN3 under room temperature conditions yields C-2 alkylated chromones.98

The reaction mechanism was proposed as a radical pathway and supported by the suppression of the reaction in the presence of radical scavengers (Scheme 50). Initially, trifluoroacetate of PhI(O2CCF3)2 was exchanged by azides of sodium azide by a ligand exchange method to form 156, and thermolysis of 156 generates 160 and an azide radical. The azide radical abstracts H from the alkane to afford alkyl radical 157, which upon reaction with the electrophilic C-2 position of chromones and further oxidation gives C-2 alkylated chromones.

R. Samanta et al. in 2015 introduced transition metal-free oxidative, regioselective cross-coupling of chromones with non-functionalized azoles for the generation of C-2-functionalized chromones with a new C–N bond (Scheme 51). The reaction is having prior importance due to the formation of...
the C-N bond, which is ubiquitous in natural products and bioactive molecules.99

Initially, 162 reacts with molecular iodine forming an electrophilic iodine source 163 (Scheme 52). The reaction of 164 with 1c gives an oxonium ion. The oxonium ion 164 thus formed exists in equilibrium with the corresponding iodonium ion 165. Subsequently, 164 or 165 gets trapped by iodide to form product 166. The compound 166, which is not stable under basic conditions, thus undergoes a base-mediated dehydroiodination to generate 3-iodochromone 167.

Yi-Han Chang et al. 2021 synthesised 2-(1H-1,2,4-triazol-1-yl)-4H-chromen-4-one by dehydroiodination (Scheme 53). The method used 1,2,4-triazole 162 as a catalyst for the generation of 3-iodochromones by reaction of molecular iodine with chromones at the C-3 position. Dehydroiodination on 3-iodochromone yields 2-(1H-1,2,4-triazol-1-yl)-4H-chromen-4-one 170a. Yi-Han Chang et al. used the method for the generation of biologically relevant compunds with phenols and 3-fluorothiol substitution at the C-2 position. The anti-inflammatory effect of phenols and 3-fluorothiol-substituted chromones against superoxide anion generation was studied. The tetrazole group acts as the leaving group in the reaction of chromones with phenols and thiophenols.100

5 Conclusion

To summarize, the review covered the site-selective functionalisation of non-prefunctionalized chromones at the C-5, C-3 and C-2 positions. All the functionalization reactions either catalyzed by a transition metal or occurring under metal-free conditions were mentioned in the review. A detailed review of all the reaction transformations by hydrogen at the active positions of chromone into different functional groups was presented. The review classifies the reactions into two categories to enhance the understanding. They are transition metal-catalysed reactions in which the reaction proceeds by the formation of a metallacycle intermediate and the other one as a metal-free reaction transformation. The second topic is a wider one where the reaction can be nitration, a base-catalysed allylation, a nucleophilic or an electrophilic reaction, and a catalyst- and ligand-free reaction. The C-2 position of chromones is more electron deficient and it may go for reactions with nucleophilic coupling partners or alkyl radicals. Similarly, the C-3 position is electron rich and it undergoes reactions with electrophilic coupling partners. There are also reports for catalyst and additive-free sulfonylation at the C-3 position. Therefore, to include all the functionalization reactions in the review, we summarise all these reactions under metal-free functionalization.

The non-pre-functionalized reactions are having greater importance compared to conventional prefunctionalized reactions. The reaction occurs in fewer steps without the liberation of halogen wastes when compared to conventional reaction methods. Despite these immense advancements in the site-selective sp² C-H activation of chromones, the methodologies suffer from serious drawbacks. It is difficult to attain site-selective substitution on the C-3 site of the chromone. Most of the reactions depend on the pre-functionalization of the C-2 sites before the functionalization at the C-3 position. On studying the yield of the functionalization reactions, we can see that the yield of these reactions is comparatively low. The lower yield of these site-selective products needed to be addressed in future. Similarly, all the C-H functionalization reactions on the arene ring of chromone take place at the C-5 position alone. This is because of the inherent direction of the carbonyl group on the chromone moiety. Methodologies focusing on these
drawbacks will facilitate the future advancement of these reactions.

Conflicts of interest

There are no conflicts to declare.

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