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Divergent synthesis of flavones and flavanones from 2'-hydroxydihydrochalcones via palladium(11)catalyzed oxidative cyclization†

Seung Hwan Son,‡^a Yang Yil Cho,‡^a Hyung-Seok Yoo,^a Soo Jin Lee,^a Young Min Kim,^a

Divergent and versatile synthetic routes to flavones and flavanones via efficient Pd(II) catalysis are disclosed. Received 3rd March 2021 These Pd(III) catalyses expediently provide a variety of flavones and flavanones from 2'hydroxydihydrochalcones as common intermediates, depending on oxidants and additives, via DOI: 10.1039/d1ra01672e discriminate oxidative cyclization sequences involving dehydrogenation, respectively, in a highly atomeconomic manner

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Flavones and flavanones are widely occurring natural flavonoids produced by various medicinal plants and their synthetic derivatives, featuring a main 15-carbon skeleton possessing 2 phenyl rings and 1 oxacycle. In recent decades, they have been considered privileged structures exhibiting various biological activities,2 such as anti-inflammatory,3 anti-cancer,4 neuroprotective,5 and estrogen-related functions6 (Fig. 1). As routes towards such privileged flavonoids, many strategies, including the Allan-Robinson reaction for flavones and intramolecular conjugate addition of 2'-hydroxychalcones for flavanones, have been reported.7 Among these, intramolecular cyclization of 2'hydroxychalcone intermediates was conventionally used for the synthesis of flavanones due to the readily available properties of the substrates through the condensation of 2'-hydroxy-acetophenones and corresponding aldehydes.8 In addition, such flavanones can be readily converted into flavones via further oxidation processes (Scheme 1).9 Thus, these types of flavonoid synthesis involving intramolecular conjugate addition have been generally used for flavonoid synthesis, 10 but they generally have drawbacks of requiring harsh conditions such as acidic or basic reflux conditions for cyclization, indicating that chemically labile compounds may in some cases not be tolerable in reaction conditions.11 Additional oxidation steps in which flavanones are transformed into flavones often require the use of strong oxidants such as I2 that enable side reactions to occur.12 In this regard, novel synthetic routes featuring transformative and compatible reaction conditions toward

flavonoids have been pursued in recent decades. In particular, given the increasing importance of constructing a privileged chemical library for probing biological systems, efficient and divergent synthesis of flavonoids is necessary.¹³

Taken together, it is important to develop a novel and versatile synthetic transformation of common substrates into flavones and flavanones under mild conditions and such a transformation is also expected to enable straightforward reactions for the construction of a privileged flavonoid library. As mentioned above, 2'-hydroxychalcones have been conventionally used as main precursors in flavonoid synthesis, but they are reactive and unmanageable in some cases due to their reactive α,β-unsaturated carbonyl and enol ether moieties. In addition, their use is generally involved with several disadvantages, as mentioned above.14 Thus, we think that the use of common intermediates instead of 2'-hydroxychalcones that is traditionally used but somewhat problematic is required. Recently, direct β-functionalization of simple ketones via several catalysts using transition metals and light was reported,15 indicating that simple ketones can be useful and potential surrogates of α,β-unsaturated ketones for further

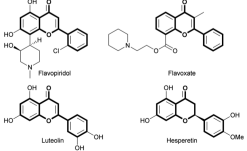


Fig. 1 Examples of bioactive flavonoids.

^aCollege of Pharmacy, Kyung Hee University, Seoul 02447, Republic of Korea. E-mail: kimnj@khu.ac.kr ^bDepartment of Life and Nanopharmaceutical Sciences, Graduate School, Kyung Hee

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[‡] These authors contributed equally to this work.

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Scheme 1 Synthetic strategy for flavonoids pursued in this study.

oxidative catalysis. In addition, we reported that chromanones, simple ketones, can be converted into flavones and flavanones via palladium(π)-catalyzed dehydrogenation-mediated coupling sequences with arylboronic esters and acids, respectively, and also recently reported that N-substituted azaflavanone can be synthesized from N-substituted aminodihydrochalcones via Pd(π)-catalyzed oxidative cyclization. In Spired by these previous works, we sought to find novel and divergent synthetic routes to a variety of privileged flavonoids, particularly flavones and flavanones using common starting materials.

Herein, we report palladium(II)-catalyzed oxidative cyclization of 2'-hydroxydihydroxychalcones that are novel and tolerable isosteres of 2'-hydroxychalcones as an efficient and divergent route for the synthesis of flavones and flavanones. To find an optimal reaction condition where flavone and flavanone were synthesized via Pd(II)-catalyzed oxidative cyclization from the common intermediate in independent manners, simple 2'hydroxydihydrochalcone was selected as the model compound to screen the reaction conditions (Table 1). Based on the speculation that the dehydrogenation process from 2'-hydroxydihydrochalcone to 2'-hydroxychalcone might proceed further reactions into flavone and flavanone, we initially tested the reaction conditions which are known to enable Pd(II)-catalyzed dehydrogenation to occur with Pd(TFA)₂ and DMSO under an O₂ atmosphere, which is eco-friendly oxidant.17 The use of this reaction condition resulted in the formation of flavanone 4a as the major product (31% yield, entry 1) and flavone 3a with 12% yield, along with a small amount of 2'-hydroxychalcone 2a (14%). This result implied that 2'-hydroxydihydrochalcone could be dehydrogenated and slightly transformed into desired flavonoids under the reaction conditions featuring Pd(II) catalysis. Based on the results, we tried to optimize the reaction conditions by using additives such as heterocyclic amines or inorganic bases as Pd(II) ligands. In the presence of most of the heterocyclic amines and K₂CO₃, the overall yields of the reactions for the synthesis of flavonoids were significantly increased (entries 2–5). Interestingly, the catalytic system featuring 2,2'bipyridine (bpy) as a ligand provided flavone 3a in 55% yield as a major product, along with flavanone 4a in 10% yield, indicating that bidentate amine was better than the other

Table 1 Optimization of the reaction conditions^a

Entry	Additive	Oxidant	Yield ^b [%]		
			2a	3a	4a
1	_	O_2	14	12	31
2	K_2CO_3	O_2	8	28	20
3	DMAP	O_2	6	29	36
4	Pyridine	O_2	5	41	37
5	Pyrimidine	O_2	7	48	32
6	bpy	O_2	5	55	10
7	5-NO ₂ phen	O_2	3	81	2
8	_	Benzoquinone	7	1	12
9	_	$K_2S_2O_8$	11	4	19
10	_	AgOAc	12	13	29
11	_	$Cu(OAc)_2$	33	6	44
12	Phenanthroline	$Cu(OAc)_2$	23	27	34
13	Pyridine	$Cu(OAc)_2$	14	13	33
14	DMAP	$Cu(OAc)_2$	25	22	35
15	K_2CO_3	$Cu(OAc)_2$	7	1	16
16	AcOH	$Cu(OAc)_2$	20	11	42
17	HCO_2H	$Cu(OAc)_2$	9	6	43
18	p-TsOH	$Cu(OAc)_2$	15	21	27
19 ^c	_	$Cu(OAc)_2$	30	3	55
$20^{c,d}$	_	Cu(OAc) ₂	8	10	79
$21^{c,e}$	_	$Cu(OAc)_2$	_	16	_

 $[^]a$ Reactions were carried out in the presence of 0.3 mmol of **1a**, 10 mol% Pd(TFA)₂, 20 mol% additive, molecular oxygen or 1.0 equiv. oxidant and DMSO 1 mL at 100 °C for 15–48 h. b Isolated yield. c DMSO 3 mL. d Addition of 2 N HCl 20 mL and ethyl acetate 10 mL for 24 h after **1a** was consumed. e Cu(OAc)₂ 2 equiv.

monodentate amines, including pyridine as a Pd(II) ligand, under incorporating conditions (entry 6). Notably, the use of 5-nitro-1,10-phenanthroline as a ligand enables the reaction to exclusively yield flavone 3a with the highest yield of 81% and flavanone 4a in 2% yield (entry 7).

Next, we focused on optimizing the conditions where flavanone, rather than flavone, is converted from 2'-hydroxydihydrochalcone that is a common starting material. In the view of the oxidation process, we supposed that the use of a stoichiometric amount of oxidant rather than molecular oxygen can result in a stepwise conversion of 2'-hydroxydihydrochalcone into 2'-hydroxychalcone, leading to the flavanone through the concurrent intramolecular 1,4-addition, rather than to the flavone for which an additional oxidation process may be needed. With this insight, we tested stoichiometric amounts of various oxidants instead of molecular oxygen for obtaining flavanone exclusively based on the entry 1 conditions (entries 8-11). As anticipated, in most of the conditions, flavanone was synthesized as a major product, but the yields were generally poor. When Cu(OAc)₂ was used as the oxidant, the yield of flavanone was increased to 44%, along with the formation of flavone at a yield of 6% (entry 11). In addition, we screened

various bases and acids as additives or potential ligands for improving the reaction under the conditions where Cu(OAc)₂ was used as the oxidant.

Heterocyclic ligands, including monodentates and bidentates, slightly decrease the yields of the reaction for providing flavanones (entries 12–14). This trend was also observed for the addition of potassium carbonate or acids (entries 15–18). Thus, unlike for flavone synthesis, additives did not significantly improve the reaction efficiency. With the entry 11 condition (no additive), the concentration was diluted to 0.1 M for facilitating intramolecular cyclization, and the reaction afforded flavanone 4a in 55% yield (entry 19). Next, with the conditions of entry 19, an additional 2 N HCl was added to cyclize the remaining

Pd(TFA)₂ 5-NO₂ phen DMSO (0.3 M) В O₂, 100 °C R = F. 3b. 70% R = Cl, **3c**, 84% R = Br, **3d**, 92% R = OTf 3e 72% = OPiv, **3f**, 55% 3a 81% (82%) 3j, 83% 3i 59% 31 79% 3m 71% 3o. 47% (72%)^a 3a. 87% 3s 75% R = F, 3v, 78% R = I, 3y, 65%

Scheme 2 Reaction scope of the flavone synthesis. Reaction conditions: 1 (1.0 equiv.), $Pd(TFA)_2$ (10 mol%), $5-NO_2-1,10$ -phenanthroline (20 mol%) and DMSO (0.3 M) at 100 °C under O_2 for 48 h. ^a Yield determined by ¹H NMR analysis. ^b 1.2 g scale reaction.

3ac, 15%

chalcone to flavanone under acidic conditions when 2'-hydroxydihydrochalcone **1a** was completely consumed. Under these conditions, flavanone **4a** was successfully obtained with the highest yield of 79% (entry 20). The use of 2 equivalents of Cu(OAc)₂ generated flavone **3a** as a detectable major product in lower yield (16%) than expected, indicating that excess Cu(OAc)₂ is less effective than molecular oxygen for flavonoid synthesis.

With the optimized conditions in hand (Table 1, entry 7), we applied our methodology for the synthesis of flavones with various substituents from the corresponding 2'-hydroxydihydrochalcone (Scheme 2). A variety of aryl A ring derivatives were successfully synthesized under these conditions. The reaction was well-tolerated and proceeded with the 2'-hydroxydihydrochalcone derivatives with electron-withdrawing groups such as halogens, triflate, pivalate and nitro groups (3b-f, 3h, and 3v-y), regardless of the A or B ring. In the case of using the substrates possessing electron-donating substituents such as methyl (3g), hydroxy (3i, 3o, and 3p), methoxy (3j, 3l and 3n), benzyloxy (3k), dimethoxy (3 m, 3q and 3r) and trimethoxy

Scheme 3 Reaction scope of the flavanone synthesis. Reaction conditions: 1 (1.0 equiv.), $Pd(TFA)_2 (10 \text{ mol}\%)$, $Cu(OAc)_2 (1.0 \text{ equiv.})$ and DMSO (0.1 M) at $100 \,^{\circ}\text{C}$ under Ar for 15 h; then addition of 2 N HCl 20 mL and ethyl acetate 10 mL for 24 h at $100 \,^{\circ}\text{C}$. ^a No addition of 2 N HCl and ethyl acetate. ^b 1.0 g scale reaction.

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groups (**3s** and **3t**), the reactions progressed well in moderate to good yields. In particular, the highest yield of 95% was obtained for 3'-hydroxyflavone **3p**. 2-(Naphthalen-2-yl)-4*H*-chromen-4-one **3u** was formed in 72% yield. However, 3-methylchromenone **3z** was obtained from a corresponding 2'-hydroxydihydrochalcone in a relatively lower yield (30%). The substrates with heterocycles such as *N*-methyl indol-3-yl, thiophen-2-yl, and furan-2-yl, as B ring, were also converted to corresponding flavones **3aa**, **3ab**, and **3ac** in 32%, 54%, and 15% yields.

Next, under the optimized conditions (Table 1, entry 20), we performed the reaction for the synthesis of flavanones using various derivatives of 2'-hydroxydihydrochalcone 1 that is the common substrate for the synthesis of flavones (Scheme 3). The desired flavanone products possessing electron-deficient substituents such as halogens and ester groups (4b-c, and 4o) as well as electron-rich substituents such as hydroxy and alkoxy moieties regardless of the A or B ring (4d-m) were readily obtained in moderate to good yields. In particular, flavanones with naturally abundant oxygen-containing moieties (4e and 4g-m) were generated effectively in the reaction. Flavanone 4n with a naphthyl moiety was also formed well in 72% yield. A substrate containing a thiophene-2-yl group as B ring was also converted to flavanone 4r in 29% yield. 3-Methylchromanone 4q was obtained from a corresponding 2'-hydroxydihydrochalcone in a lower yield (18%) compared to the other flavanones, which is similar to the case of 3-methylchromenone 3z. In the case of reactions for preparing 3z and 4q, lots of 1-(2-hydroxyphenyl) butan-1-one, a starting material, remained not to be reacted. A few flavones such as 3ac and flavanones such as 4p were not obtained in good yield, along with several side products. Collectively, these results indicated that 2'-hydroxydihydrochalcones that are chemically compatible substrates can be obviously transformed into flavones and flavanones in divergent and efficient manners, respectively.

To further investigate the divergent utility of our synthetic methodology, we applied it to the synthesis of natural flavonoids (Scheme 4). Under the optimal conditions, geraldone dimethyl ether 5a (ref. 18) and butin trimethyl ether 5b, 19 the

Optimized condition
for flavone

OMe

OMe

OMe

OMe

OMe

Optimized condition
for flavanone

Optimized condition
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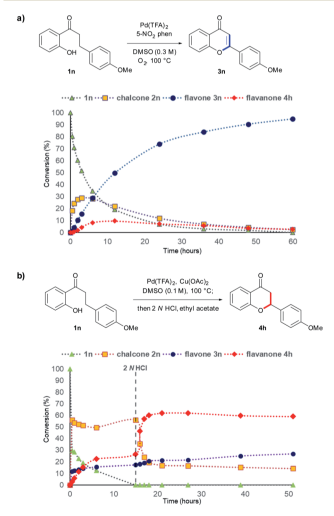
Optimized condition
For flavone

Optimized condition
OMe

Scheme 4 Application for the synthesis of natural products.

biologically active natural flavonoids, were successfully transformed from common substrate **1aa** in moderate yields (88% and 48%, respectively, Scheme 4). Tithonine **5c**, an anti-inflammatory flavone known as a selective COX (cyclooxygenase)-1 inhibitor, ²⁰ was also synthesized well under our reaction conditions.

Next, to investigate the mechanism of our synthetic methodology, we carried out kinetic analysis of the reaction using high-pressure liquid chromatography (HPLC) to examine the time-dependent conversion 2'-hydroxy-4methoxydihydrochalcone 1n to the corresponding flavone 3n or flavanone 4h under optimized reaction conditions. For flavone synthesis, it was observed that the amount of 2'-hydroxy-4- methoxydihydrochalcone 1n gradually decreased over time, and 2'-hydroxy-4-methoxychalcone 2n was increasingly formed in a dramatic manner with the concurrent formation of flavone 3n within 5 h (Scheme 5a). Then, 2n was gradually diminished, and 3n was significantly formed, indicating that 2n may be consumed for the synthesis of the desired flavone 3n. In addition, in the conditions of flavanone synthesis (Scheme 5b), it



Scheme 5 Kinetic experiments for flavone and flavanone synthesis from 2'-hydroxy-4-methoxydihydrochalcone **1n**. (a) Flavone synthesis condition. (b) Flavanone synthesis condition.

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Fig. 2 Plausible mechanism.

was observed that the reaction proceeded rapidly at first (Fig. S1†), resulting in the drastic formation of 2'-hydroxy-4-methoxychalcone **2n** with the concurrent formation of flavonoids. After 15 h, the reaction was observed to reach the plateau state. Upon adding 2 N aqueous HCl, flavanone **4h** was immediately formed with a significant disappearance of **2n**, indicating that **2n** may also be consumed for the synthesis of the desired flavanone **4h**.

Based on the kinetic results, a plausible mechanism for this divergent synthesis is proposed, as shown in Fig. 2. First, $Pd(\pi)$ -catalyzed dehydrogenation may result in the oxidative conversion of 2'-hydroxydihydrochalcone **1a** into 2'-hydroxychalcone **2a.**¹⁶ Then, common intermediate **2a** can be transformed into the desired flavanone **4a** through Michael addition and flavone **3a** through oxidative Heck coupling, respectively, depending on the optimized reaction conditions that are used in the synthesis. During oxidation processes such as dehydrogenation and oxidative cyclization, Pd(0) species regenerated in the reaction would be reoxidized into $Pd(\pi)$ *via* the [O] process, where O_2 or $Cu(OAc)_2$ acts as the main oxidant.²¹

Conclusions

We reported novel and versatile approaches for the synthesis of flavones and flavanones from common intermediates via Pd(II)-catalyzed oxidative cyclization. These methodologies greatly facilitate the construction of chemical libraries of flavones and flavanones in moderate to high yields using easily accessible reagents and offer good compatibilities with various functional groups under mild conditions in atom-economic manner. Further studies on the biological evaluation of a wide variety of flavonoids synthesized by our methodology are ongoing.

Conflicts of interest

There are no conflicts to declare.

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