

Synthesis of *ortho*-Acylphenols through the Palladium-Catalyzed Ketone-Directed Hydroxylation of Arenes**

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In memory of Michael W. Day

ortho-Acylphenols are an important structural motif found in a diversity of bioactive molecules ranging from natural products to drugs (Figure 1).^[1] Moreover, they also serve as

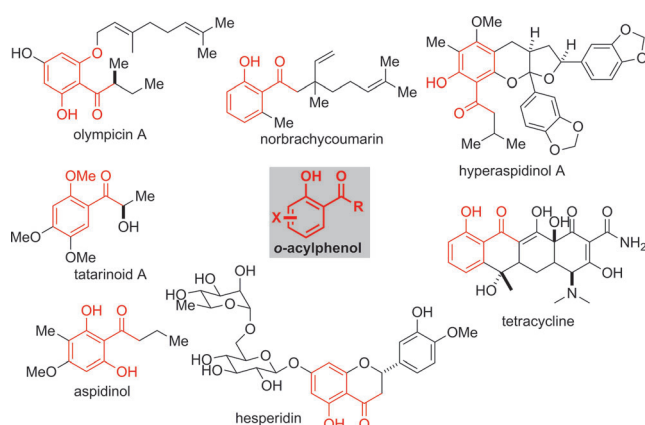
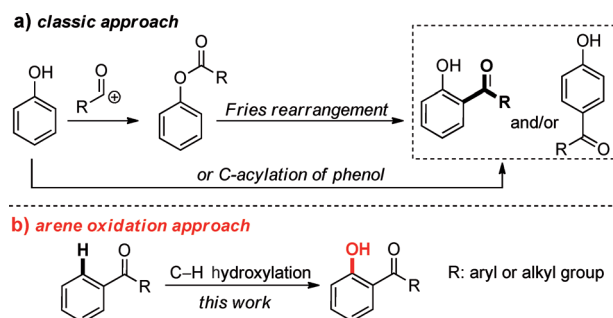


Figure 1. Natural products and drugs containing *o*-acylphenols.

versatile building blocks for the synthesis of various pharmaceuticals, such as warfarin,^[2] as well as agrichemicals, flavors, and fragrances.^[3] Classic approaches to the synthesis of *o*-acylphenols generally involve a two-step process: acylation of phenols followed by Fries rearrangement of the resulting phenyl esters (Scheme 1a).^[4] On the other hand, direct C-acylation of phenols has also been known under more forcing conditions.^[5] Although effective, these approaches are often complicated by the formation of undesired *p*-substituted products when bulky acyl groups need to be introduced, as well as the limited variety of ketones that can be generated.^[6,7]



Scheme 1. Synthesis of *o*-acylphenols.

An alternative approach would be to use ketones as the starting material and introduce the hydroxy group (phenol) by direct arene oxidation (Scheme 1b). This is attractive because aryl ketones are a readily available feedstock and can be prepared in one step through electrophilic aromatic acylation,^[8] benzylic oxidation,^[9] Wacker oxidation of the corresponding styrenes,^[10] or other means.^[11] The direct hydroxylation of arylketones either through a radical approach^[12] or through heterogeneous catalysts^[13] is known; however, these reactions lack site-selectivity and provide a mixture of all *o*-, *m*-, *p*-isomers. A highly *ortho*-selective hydroxylation of arylketones remains to be developed.

Transition metal (TM)-catalyzed C–H oxidation presents an increasingly important approach for preparing heteroatom-substituted aromatic compounds.^[14] Recently, such transformations driven by weak coordination have become a powerful means for further functionalization of simple organic feedstocks.^[15] The first carbonyl-directed (carbamate) Pd^{II}-catalyzed sp³ C–H oxidation was developed by Yu et al.;^[16] subsequently, a remarkable achievement using carboxylate-directed (–CO₂Na) C–H carboxylation was reported by the same group, which significantly expanded the scope of using weakly coordinating σ donors as directing groups (DGs) in C–H oxidations.^[17] In terms of *ortho*-oxidation of arenes, Yu and co-workers further demonstrated an *ortho*-hydroxylation with carboxylic acids as DGs using air or O₂ as the source for the hydroxy group.^[18] On the other hand, an *ortho*-acetoxylation using oximes as DGs was first reported by Sanford et al., wherein oximes can be considered as a ketone surrogate.^[19] Very recently, Ru-catalyzed *ortho*-hydroxylations using esters and amides as DGs were reported by the groups of Rao^[20] and Ackermann,^[21] respectively. These seminal works could potentially serve as effective ways to access *o*-acylphenols, although additional steps are needed to reveal the phenol or to obtain the ketone moiety. Thus, it

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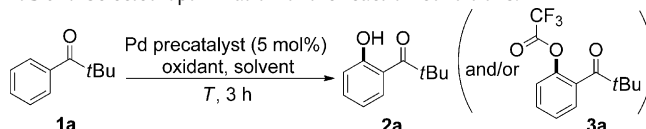
would be a more step-economical approach to synthesize *o*-acylphenols if ketones could be directly employed as the DG for the catalytic arene hydroxylation reaction.^[22]

Although rare compared to other directed reactions, ketone-directed catalytic C–H functionalization has been documented for almost two decades since the pioneering work on Ru-catalyzed C–H/olefin coupling reactions by Murai et al.^[23] However, most of these transformations are restricted to C–C bond formations (by the groups of Murai,^[24] Chatani,^[25] Miura,^[26] Brookhart,^[27] Cheng,^[28] Glorius,^[29] Jegannathan,^[30] Shi,^[31] and others). Only recently, the groups of Liu^[32] and Glorius^[33] reported the first ketone-directed C–N and C–Br/I bond-forming reactions using Pd and Rh catalysts, respectively. To the best of our knowledge, catalytic ketone-directed arene oxidation to form C–O bonds has not been previously described. The likely challenges are: 1) As a hard Lewis base and even weaker σ donor (compared to oximes, amides, carboxylic acids, and esters), ketones generally do not coordinate strongly with TMs,^[34] which leads to lower reactivity. 2) Undesired oxidations (particularly with oxygen-based oxidants) can occur either at the enolizable α C–H bond or at the α C–C bond of the ketone through a Baeyer–Villiger-type oxidation.^[35]

Stimulated by the previous success of C–H activation driven by weak coordination^[15,17] and the need for efficient synthesis of *o*-acylphenols from arylketones, we pursued this challenge using a Pd-catalyzed formal arene hydroxylation reaction. It was envisioned that the above-mentioned challenges could potentially be overcome by: 1) increasing the Lewis acidity of the metal and 2) using milder oxidants and reaction conditions. Herein, we report the use of ketone carbonyls as DGs for a Pd-catalyzed *ortho*-selective formal hydroxylation of arenes to provide various 2-acylphenols; preliminary results of the first ketone-directed oxidative carbonylation to give an unusual ketal–lactone are also described.

We initiated our studies by examining the *ortho*-oxidation of 2,2-dimethylpropiophenone, a problematic substrate for the Fries-rearrangement.^[36] Different Pd^{II} precatalysts, oxidants, and solvents were tested (Table 1). Not surprisingly, the standard Pd(OAc)₂ and PhI(OAc)₂ (DIB) combination, which is commonly used in nitrogen-based DG-directed C–H oxidations,^[19,14c] did not give any of the desired product at 80°C in 1,2-dichloroethane (DCE) or AcOH (Table 1, entries 1 and 2). A search for more electrophilic catalysts and oxidants was pursued. Eventually, Pd(TFA)₂ (5 mol %) with PhI(TFA)₂ (BTI; 2 equiv) was found to give quantitative conversion with 83 % yield of the desired *ortho*-oxidation product (entry 3). Trifluoroacetate **3a** is the initial product generated, which is confirmed by both GC-MS and NMR analysis. We were delighted to discover that compound **3a** is labile, and readily converts into the corresponding phenol (**2a**) upon purification by column chromatography with silica gel. The use of other solvents, such as dioxane, THF, and MeCN, in place of DCE proved much less effective (entries 4–6). Interestingly, when BTI is used as an oxidant, comparable yields can be obtained when Pd(TFA)₂ is replaced by either Pd(OAc)₂ or Pd(OPiv)₂, whereas this was not the case for PdCl₂^[37] (entries 7–9). In contrast, DIB cannot be used in

Table 1: Selected optimization of the reaction conditions.^[a]



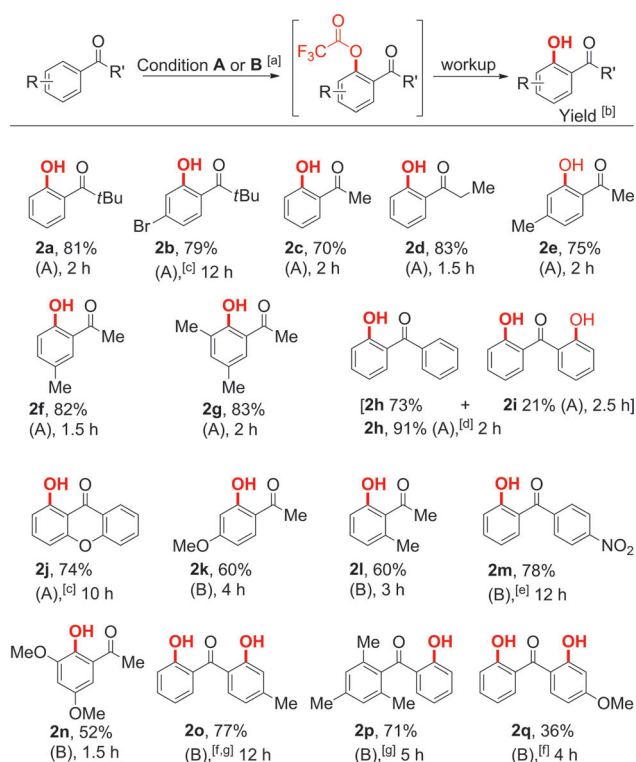
Entry	Catalyst	Oxidant ^[b]	Solvent	T [°C]	Conv. [%] ^[c]	Yield [%] ^[c]
1	Pd(OAc) ₂	DIB	DCE	80	< 5	0
2	Pd(OAc) ₂	DIB	AcOH	80	12	0
3	Pd(TFA)₂	BTI	DCE	80	100	83
4	Pd(TFA) ₂	BTI	dioxane	80	30	7
5	Pd(TFA) ₂	BTI	THF	80	17	8
6	Pd(TFA) ₂	BTI	MeCN	80	22	0
7	Pd(OAc) ₂	BTI	DCE	80	100	79
8	Pd(OPiv) ₂	BTI	DCE	80	100	78
9	PdCl ₂	BTI	DCE	80	15	0
10	Pd(TFA) ₂	DIB	DCE	80	8	0
11 ^[d]	Pd(TFA)₂	BTI	DCE	80	100	84
12	Pd(TFA) ₂	KPS	DCE	80	12	4
13 ^[e]	Pd(TFA) ₂	KPS	DCE	80	36	17
		(TFA)				
14	Pd(OAc)₂	KPS	TFA	50	100	88
15	Pd(OAc) ₂	KPS	TFA/DCE (1:1)	50	100	77
16	–	BTI	DCE	80	< 5	0
17	–	KPS	TFA	50	< 5	0

[a] The reactions were run on a 0.2 mmol scale with 1 mL solvent over 3 h. [b] In all cases, 2 equiv of oxidant was used. [c] Determined by ¹H NMR spectroscopy using mesitylene as the internal standard. [d] 2 mol % catalyst loading, reaction time 9 h. [e] Reaction time 9 h. BTI = [bis(trifluoroacetoxy)iodo]benzene, DCE = 1,2-dichloroethane, DIB = (diacetoxyiodo)benzene, KPS = K₂S₂O₈, Piv = pivaloyl, TFA = trifluoroacetic acid.

place of BTI, even when using Pd(TFA)₂ as a catalyst (entry 10; see below for a mechanistic discussion). Furthermore, the reaction gives full conversion with 84 % yield in 9 h when the catalyst loading is lowered to 2 mol % (entry 11).

Less expensive inorganic oxidants, such as K₂S₂O₈ (KPS), have also been examined. The desired product was obtained in 4 % yield by simply replacing BTI with KPS (entry 12). The yield was improved to 17 % when 2 equiv of TFA was added (entry 13). Towards the end, complete conversion with 88 % yield was obtained when using TFA as the solvent and KPS (2 equiv) as the oxidant (entry 14). Note that the reaction proceeds at a lower temperature (50°C) with the less expensive Pd(OAc)₂ as a pre-catalyst, thus it is complementary to the conditions described in entry 11. Although it is well recognized that BTI is able to oxidize electron-rich aromatic compounds through a single-electron transfer mechanism,^[38] control experiments indicated that neither of the conditions in entries 16 or 17 showed significant reactivity in the absence of Pd.

With the optimized conditions in hand, we next investigated the substrate scope of this reaction (Scheme 2). A variety of aromatic ketones smoothly underwent the *ortho*-hydroxylation. Both aryl and alkyl-substituted ketones with different steric properties gave satisfying yields of the correspond *o*-acylphenols. Note that benzylic and enolizable hydrogens, which can participate in various oxidation reac-

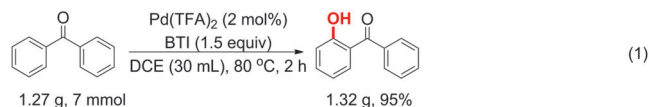


Scheme 2. Substrate scope. [a] 0.4 mmol scale. Condition A: Pd(TFA)₂ (5 mol %), BTI (2.0 equiv), DCE 2 mL, 80 °C. Condition B: Pd(OAc)₂ (5 mol %), K₂S₂O₈ (2 equiv), TFA 2 mL, 50 °C. [b] Yield of isolated product. [c] Pd(TFA)₂ 10 mol %, 100 °C. [d] with 1.3 equiv of BTI. [e] 70 °C. [f] K₂S₂O₈ 3 equiv [g] TFA 1 mL and DCE 1 mL. BTI = [bis(trifluoroacetoxy)i]d]benzene, DCE = 1,2-dichloroethane, TFA = trifluoroacetic acid.

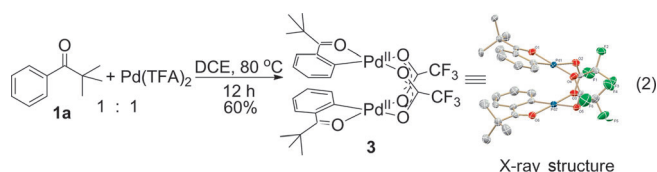
tions, are also tolerated under these conditions (**2c–2g**, **2k**, **2l**, **2n**, **2p**). Moreover, aryl bromides, methyl ethers, xanthenes, and nitro groups are also compatible (**2b**, **2j**, **2k**, **2m**, **2q**). Mono-oxidation is selectively achieved on mono-arylketones, most likely because further C–H activation/oxidation is inhibited by the strong inductive effect of the introduced trifluoroacetate group (which is cleaved upon workup). Notably, a single regioisomer (**2f**) was obtained in 82% yield using *m*-methyl-substituted ketone, which indicates that C–H oxidation is preferred at the less hindered site. For diarylketones, hydroxylation occurred on both phenyl rings in the presence of 2 equiv of BTI, but formation of mono-oxidized product **2h** (91% yield) can be controlled using 1.3 equiv of the oxidant. With excess oxidants, both electron-neutral and electron-rich aromatic rings could be smoothly oxidized without increasing catalyst loading (**2o** and **2q**). Furthermore, for unsymmetrical benzophenone substrates, the more electron-rich aromatic rings are selectively oxidized over the electron-poor ones, which is consistent with an electrophilic palladation mechanism (**2m**).^[14] The two conditions (**A** and **B**) employed in this study are complementary: generally, condition **A** (using BTI as an oxidant and DCE as a solvent) is milder and more selective, whereas condition **B** (using KPS/TFA) is more reactive. For example, condition **A** effectively circumvents oxidation on the enolizable protons;

on the other hand, sterically hindered or nitro-containing ketones react slowly under condition **A**, whereas high conversion was obtained under condition **B** (**2l** and **2m**).

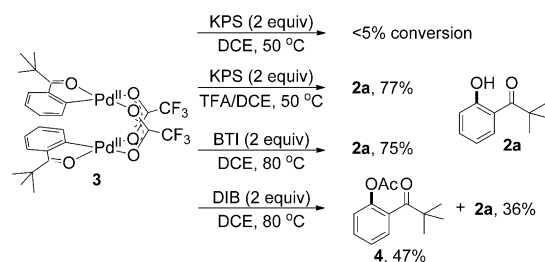
This reaction could also be run on a gram scale with only 2 mol % catalyst loading, affording product **2h** in 95% yield within 2 h [Eq. (1)].



Next, in order to understand the mechanism of this Pd-catalyzed ketone-directed C–H hydroxylation reaction, step-wise stoichiometric reactions were carried out. Compared to previous methods of preparing ketone-chelated palladacycles where additional strong acids (TfOH or TFA) are involved,^[28a,32] we found that the ketone is able to react directly with Pd(TFA)₂ in DCE to give the dimeric Pd complex (**3**) in good yield [Eq. (2)].^[39]



Moreover, although acid is not required for the C–H activation step, it was found to be critical in the subsequent KPS-mediated oxidation/C–O bond formation step (Scheme 3). No reaction occurred upon treating palladacycle

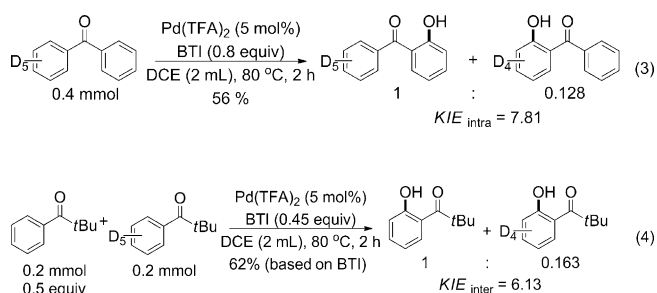


Scheme 3. Oxidation of Complex **3**.

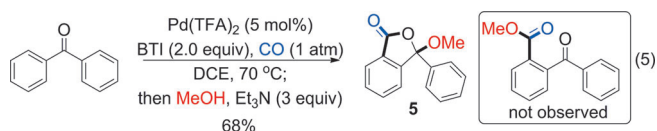
3 with KPS alone in DCE, whereas the addition of TFA promoted the formation of the product in 77% yield. When BTI was used as the oxidant (without adding acids), the corresponding arene oxidation products were also obtained in good yields. Moreover, the use of less reactive DIB as an oxidant led to the formation of *o*-OAc substituted ketone (**4**).

These studies suggest that trifluoroacetate anions play a critical role in maintaining the high electrophilicity of the Pd catalyst for the initial C–H bond cleavage, but are not required as a part of the oxidants. Furthermore, high KIE values were observed both in the intramolecular ($k_{H/D}$ = 7.81)

and intermolecular ($k_{\text{H/D}}=6.13$) competition experiments [Eq. (3) and (4)].^[40]



With this mechanistic knowledge in hand, we made further preliminary efforts to explore a ketone-directed Pd-catalyzed oxidative C–H carbonylation reaction [Eq. (5)].^[41,42] To our delight, under almost identical con-



ditions, except for the use of a CO atmosphere (1 atm), an interesting *ortho*-carboxylation product (5) was obtained in 68% yield when benzophenone was used as the substrate.^[43] Upon workup with methanol and Et₃N, the originally anticipated methyl ester was not observed; instead, the methyl ketal-lactone was formed as the major product.^[44] Although the mechanism of this reaction is still unclear, the mixed-anhydride was proposed as the initial product formed.^[42] It is noteworthy that the formation of phenol products were significantly suppressed in the presence of CO.

In summary, a ketone-directed Pd-catalyzed *ortho*-hydroxylation of arenes has been developed, which is an effective approach to access various *o*-acylphenol compounds from arylketones. As an exploratory study, a Pd-catalyzed *ortho*-carboxylation of simple aryl ketones to access an unusual ketal-lactone motif has also been disclosed. Both are the first examples of simple ketone-directed C–O bond formation and carboxylation of arenes by C–H bond activation. Because of the versatile nature of ketones, the methods described herein are expected to have significant value for practical applications, and should stimulate further research on simple ketone-directed C–H functionalizations. Expansion of the scope of the reaction and substrates, as well as more detailed mechanistic studies are ongoing.

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