An Alternative to the Classical α-Arylation: The Transfer of an Intact 2-Iodoaryl from ArI(O₂CCF₃)₂**

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Abstract: The α -arylation of carbonyl compounds is generally accomplished under basic conditions, both under metal catalysis and via aryl transfer from the diaryl λ^3 -iodanes. Here, we describe an alternative metal-free α -arylation using $ArI(O_2CCF_3)_2$ as the source of a 2-iodoaryl group. The reaction is applicable to activated ketones, such as α -cyanoketones, and works with substituted aryliodanes. This formal C– H functionalization reaction is thought to proceed through a [3,3] rearrangement of an iodonium enolate. The final α -(2iodoaryl)ketones are versatile synthetic building blocks.

he transfer of an aryl group to a position α to a carbonyl is an important class of C–C bond-forming reactions, popularized with the introduction of the metal-catalyzed (mainly Pd and Cu) coupling of aryl halides (or equivalent) to enolates.^[1,2] Predating these advances, the metal-free α -arylation has been in use since the 1960s, following reports by Beringer et al. on the ability of the diaryl- λ^3 -iodanes (e.g. [Ph₂I]Cl) to transfer an aryl ligand to an enolate (Scheme 1).^[3,4] Recent studies showed that both the C- and the O-iodonium enolate intermediates can lead to the product through a [1,2] or [2,3] shift, respectively.^[5]

This methodology, including its asymmetric versions,^[6] has since then gained importance as complementary to the crosscoupling, in turn stimulating further research into diaryl λ^3 iodanes.^[7,8] Despite the attractiveness of the method, one of the two aryl groups must act as a "spectator" ligand extruded in the form of ArI. The choice of such a group (e.g. mesityl) is often the key to a selective arylation using asymmetric

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[**]	This work was supported by the ICIQ, MICINN (CTQ2011-22649), the MEC (Cons. Ing. CSD2007-00006), the Generalitat de Catalunya (2014SGR1192 and 2014SGR1105), and the China Scholarship

Council (fellowship to Z.J.). We thank MINECO for support through grant CTQ2013-46705-R and Severo Ochoa Excellence Accreditation 2014–2018 (SEV-2013-0319).

Supporting information for this article is available on the WWW under http://dx.doi.org/10.1002/anie.201405982.



Scheme 1. The Beringer-type arylation of $\beta\text{-ketoesters}$ using diaryliodonium salts.

diaryliodanes.^[7b] Although the use of the monoaryl iodonium species (e.g. PhIX₂) would thus be attractive, examples of such usage are scarce.^[9] As part of our own research on hypervalent iodine reactivity,^[10] we wish to report an α -arylation protocol that employs monoaryl iodonium species, exemplified by phenyliodine bis(trifluoroacetate) (PIFA, **2a**).

We found that exposing the β -ketoester **1** to **2a** in CH₃CN led to an unexpected *ortho*-iodoaryl species **4** in 17% yield (Scheme 2); in contrast, none of **4** was obtained using PhI(OAc)₂ or PhI(OH)(OTs) (entries 1–3, Table 1). The



Scheme 2. The outcome of treating the β -ketoester 1 with PhI(O₂CCF₃)₂ (2 a).

Table 1: Screening of conditions in the arylation of 1 with 2a (from Scheme 2).^[a]

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Entry	Solvent	PhIX ₂	Additive ^[b]	4 [%] ^[c]
1	CH₃CN	2 a	-	17
2	CH₃CN	PhI(OAc)₂	-	-
3	CH₃CN	PhI (OH) (OTs)	-	-
4	CH_2Cl_2	2a	-	14
5	CF ₃ CO ₂ H	2a	-	34
6	CH ₃ CN-CF ₃ CO ₂ H	2a	-	48
7	CH ₃ CN-CF ₃ CO ₂ H	2a	(CF ₃ CO) ₂ O	57 ^[d]
8	CH ₃ CN-CF ₃ CO ₂ H	2a	(CF ₃ SO ₂) ₂ O	< 5
9	CH ₃ CN-CF ₃ CO ₂ H	2a	H ₂ O	< 5
10	CH ₃ CN-CF ₃ CO ₂ H	PhI(OAc) ₂	(CF ₃ CO) ₂ O	23
11	CH ₃ CN-CF ₃ CO ₂ H	PhI (OH) (OTs)	(CF ₃ CO) ₂ O	26
12	CH ₃ CN-CF ₃ CO ₂ H	PhIO	(CF ₃ CO) ₂ O	52

[a] Using 1 (1.0 mmol) and 2a (1.3 mmol) in 4 mL of solvent for 4 h at rt;
[b] 1.5 equiv; [c] GC yield corrected versus the internal standard C₆H₁₁CN; [d] Yield of the isolated product.

Angew. Chem. Int. Ed. 2014, 53, 1-5

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formation of **4** was found to be solvent-dependent (entries 1 and 4–6), with a 48% yield achieved using a 1:1 CH₃CN/ CF₃CO₂H mixture. The addition of the trifluoroacetic anhydride (1.5 equiv) led to a 57% yield of **4** after 2 h at room temperature; other additives proved detrimental (entries 7–9). Under the new conditions, the use of other hypervalent iodine reagents was now possible (entries 10–12), likely through the in situ formation of **2a**. Oxidative degradation of **1** accounts for the reaction mass balance.

To probe the reaction scope, the cyclic β -ketoesters 5–7 were transformed into products 11–13 in 2 h at room temperature (Scheme 3, X-Ray structure of 13 shown). Similarly, α -



Scheme 3. The α -iodoarylation of β -dicarbonyl compounds using PIFA.

(2-iodoaryl)-diketones **14–16** were synthesized from the corresponding β -diketones **8–10**.

Particularly efficient was the arylation of α -cyanoketones (Table 2). Thus, the cyclic substrates **17–19** underwent a smooth reaction with PIFA to give a 76–80% yield of



[a] 1.3:1 ratio of **2/17**. Using **17** (1 mmol) in CH₃CN/CF₃COOH (1:1, 4 mL). [b] From *m*-Br-C₆H₄I(O₂CCF₃)₂ (**2**). [c] At 60 °C for 48 h.

20a, 21, and 22 after 6-8 h.^[11] Next, 17 was exposed to eight additional $ArI(O_2CCF_3)_2$ reagents (2b-2i) prepared by a method developed by Zhdankin et al.^[12] The use of the halo derivatives 2b-2e led to the formation of the dihaloaryl cyanoketones 20b (63%), 20c (65%), 20d (71%), and 20e (50%), with **20e** featuring the iodine flanked by a C-Br and a C-C bonds. A 76% yield of the carboxy-substituted 20 f was achieved and the p-NO₂ derivative **20** g was isolated in 68% yield. The transfer of a 2-iodo-3-methylphenyl group took place with a 49% yield (product 20h). The coupling at the two ortho CH sites of the meta-Br iodane 2i took place in a 3:1 ratio, with the minor isomer 20i' (17%) observed as two rotamers (70:30) at -20°C (see the Supporting Information). Interestingly, although the secondary cyanoketones, including benzoylacetonitrile, proved unsuitable, the 2-benzoylpropionitrile, which only differs by a 2-methyl group, gave the expected 23 in 60% yield. Finally, the protocol was used to prepare a 19 g batch of **20 a** (74%, Scheme 4).^[13]



Scheme 4. Gram-scale preparation of **20**a; α -arylation conditions as in Table 2.



Scheme 5. Two possible enolate rearrangement paths leading to 4.

Although a mechanistic study is currently underway, this formal C–H alkylation may arise from a [3,3] shift of an O-enolate **A** (Scheme 5) as seen in a related sulfoxide-mediated α -arylation.^[14] Alternatively, the selectivity could be explained by a [1,3] shift of a C-enolate **B**.

A priori, the [1,3] shift appears less likely. Indeed, while the C-enolates are intermediates in the formation of the iodonium ylides^[15] and can be isolated,^[16] the existence (or intermediacy) of the quaternary analogues (such as B in Scheme 5) is less frequent.^[5b] Furthermore, heating the isolated C-enolates typically leads to the formation of the α -C-O (e.g. C-OTs) bond.^[16] In our hands, the isolated phenyliodonium ylide PhCOC(=IPh)CN, expected to give a C-enolate upon protonation,^[15c] failed to undergo the aryl transfer under the reaction conditions. Thus, we favor a [3,3] shift of an iodonium O-enolate (A, Scheme 5), akin the iodonio-Claisen rearrangement introduced by Ochiai et al. in the 1990s.^[17,18] Despite our efforts, reaction intermediates have so far proven elusive, possibly due to the rearrangement proceeding faster than the I-O-enolate formation.^[19] Not even the o,o-disubstituted 2j allowed for the trapping of the Ienolate, leading instead to non-arylative oxidation processes (Scheme 6A). It should also be noted that the iodine-free species 24 (5%), isolated during the synthesis of 20b, proved

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Scheme 6. Additional observations in the arylation of 17.

to be the *para*-fluoro regioisomer, rather than the initially assumed *meta* (Scheme 6B); the steps leading to **24** remain to be investigated.

The cyanoketones **20a** and **21** were readily converted to the amides **25** and **26**. Although PIFA was unsuitable for the arylation of the parent cyclohexanone, the arylketone **27** could, nevertheless, be obtained through the decarboxylation of **25** (Scheme 7, top). The substrates also underwent the Suzuki (see the Supporting Information) and Sonogashira coupling reaction (Scheme 7, bottom, products **28–30**).

The reduction of **25** led to the alcohol **31** as a 4:1 *trans/cis* mixture (Scheme 8), with the solid state structures of both **25** and **31-***trans* showing an equatorial *o*-iodophenyl group.^[20] Preliminary tests showed that **31** can be converted to the hydroxy-spiroxindole **32** (X-ray structure shown for *trans*) using a Cu-catalyzed C–N coupling,^[21] with **32-cis** representing the spiroxindole portion of Gelsemine, a synthetically interesting natural product target (Scheme 8).^[22,23]



Scheme 7. Functional group manipulation in the α -(2-iodoaryl) derivatives.



Scheme 8. Some simple transformations of the ketoamide 25.

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In summary, the ArI(O_2CCF_3)₂ reagents have been used in the α -arylation of β -dicarbonyls and α -cyanoketones. The aryl transfer takes place with retention of the iodide *ortho* to the newly formed C–C bond. The method is complementary to metal-catalyzed arylations, and could overcome the issues of the aryl loss associated with the use of diaryliodonium salts. In a more general sense, the concept of a reversible formation of an iodonium-based Claisen precursor, shown here with Oenolates, might open the door to the development of a range of new synthetic methods.

Received: June 6, 2014 Revised: July 11, 2014 Published online:

Keywords: α -arylation \cdot C–H functionalization \cdot iodonium-Claisen rearrangement \cdot dehydrogenative C– C coupling \cdot hypervalent iodine

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Synthetic Methods

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An Alternative to the Classical α -Arylation: The Transfer of an Intact 2-Iodoaryl from ArI(O₂CCF₃)₂



[3,3] Iodonium enolate rearrangement: Activated ketone derivatives, including β -

dicarbonyl and α -cyanoketones, react with Arl(O₂CCF₃)₂ reagents to give an α arylated product with the iodine atom retained *ortho* to the new C–C bond. The -coupling *ortho* to iodine -likely iodonio-Claisen mechanism -complementary to metalcatalyzed arylation

reaction takes place under acidic conditions. This formal C–H functionalization reaction presumably proceeds through a [3,3] rearrangement of an iodonium enolate. The final α -(2-iodoaryl)ketones are versatile synthetic building blocks.