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High-Yielding One-Pot Synthesis of Diaryliodonium Triflates from Arenes and Iodine or Aryl Iodides†

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Unsymmetric and symmetric diaryliodonium triflates are synthesized from both electron-deficient and electron-rich substrates in a fast, high yielding, and operationally simple 10 protocol employing arenes and aryl iodides or iodine.

Hypervalent iodine compounds have recently received considerable attention as mild, non-toxic and selective reagents in organic synthesis.^{1, 2} Iodine(III) reagents with two heteroatom ligands, e.g. (diacetoxyiodo)benzene and iodosylbenzene, are successfully 15 employed in oxidations of alcohols, alkenes and α-oxidations of carbonyl compounds.³ The properties of iodine(III) reagents with two carbon ligands resemble those of metals such as Hg, Pb and Pd, allowing for reaction pathways similar to metal-catalyzed reactions while avoiding the drawbacks of cost, toxicity and 20 threshold values in pharmaceutical products. 2 Diaryliodonium salts are the most studied compounds in this class; they are versatile electrophiles in α-arylation of carbonyl compounds^{4, 5} and metalcatalyzed cross-couplings,6 and are frequently employed as photo initiators in polymerizations.⁷

Synthetic routes to diaryliodonium salts typically involve 2-3 steps, with initial oxidation of an aryl iodide to iodine(III) followed by ligand exchange with an arene to obtain the diaryliodonium salt.8 Preformed inorganic iodine(III) reagents, such as iodosyl fluorosulfate, have been employed to shorten the the route. These 30 reagents are, however, unstable and not commercially available. Reported reactions with oxidation and ligand exchange in the same pot suffer from narrow substrate scope 10, 11 or long reaction times, 11 need excess reagents¹² or employ toxic chromium reagents.¹³ In many cases a subsequent anion exchange step is necessary, as the 35 anion influences both the solubility and reactivity of the iodonium salt. Non-nucleophilic anions, such as triflate or tetrafluoroborate, have proven superior to halide anions in many applications. ^{6, 14}

The lack of general and efficient methods for the synthesis of diaryliodonium salts is cumbersome, and clearly limits their scope 40 as environmentally benign reagents in organic chemistry. 15 The development of an operationally simple protocol with broad substrate scope would greatly facilitate the application of these efficient, non-toxic arylation agents. Herein we present our preliminary results on a fast, high-yielding, one-pot synthesis of the 45 title compounds.

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An atom efficient and simple one-pot synthesis of diaryliodonium salts would involve treatment of an aryl iodide with a commercially available oxidant in the presence of an arene and a suitable acid, the anion of which would end up in the iodonium salt 50 (eqn. 1).

$$Ar^{1}-I + Ar^{2}-H \xrightarrow{\text{oxidant}} Ar^{1} \xrightarrow{Ar^{2}} X^{-} \qquad (1)$$

mCPBA has recently been reported to oxidize iodobenzene to (diacyloxyiodo)benzene, 16 which encouraged us to investigate whether this oxidant could be employed also in the direct synthesis of iodonium salts. Suitable reaction conditions for the reaction of 55 iodobenzene (1a) and benzene (2a) with mCPBA in organic were thus screened. Boron trifluoride trifluoromethanesulfonic acid were deemed as interesting acids, as they could give rise to iodonium salts with tetrafluoroborate and triflate anions, respectively, without a subsequent anion exchange 60 step. Initial attempts to combine mCPBA with boron trifluoride etherate in dichloromethane at room temperature were encouraging, as a diphenyliodonium salt was formed in moderate yield. Due to problems with purification and anion identification, the use of triflic acid instead of BF₃·OEt₂ was next investigated.

Gratifyingly, the treatment of iodobenzene with mCPBA, 17 TfOH and excess benzene in dichloromethane at room temperature resulted in a clean reaction to give iodonium triflate 3a in 82% isolated yield (Table 1, entry 1). The reaction time could be considerably reduced by increasing the temperature, delivering 3a 70 in 89% yield after 1 h at 40 °C (entry 3) or in 85% yield after only 10 min at 80 °C (entry 4). In this fast and operationally simple onepot reaction, 3a is formed directly from commercially available substrates and can be isolated with a suitable anion in high yield after a simple workup.

Table 1 Temperature influence on the synthesis of $3a^a$

	+	mCPBA, TfOH, CH	2Cl ₂) It	OTf
	1a	2a		3a	
Entry	2a (equiv)	TfOH (equiv)	T (°C)	Time	Yield $(\%)^b$
1	5	2	rt	21 h	82
2	2	3	rt	21 h	89
3	2	3	40	1 h	89
4	1.1	3	80	10 min	85

^a Reaction conditions: 1a (0.23 mmol), 2a and mCPBA (0.25 mmol) were dissolved in CH2Cl2 (2 mL), TfOH was added at rt and the reaction was stirred at the indicated temperature and time in a sealed tube. b Isolated yield.

[†] Electronic Supplementary Information (ESI) available: [Synthetic procedures, analytical data and NMR spectra for salts 3]. See http://dx.doi.org/10.1039/b000000x/

To determine the generality of this novel one-pot reaction, it was applied to the synthesis of various diaryliodonium salts 3 from aryl iodides 1 and arenes 2 (Table 2). Most previous protocols have been restricted to the synthesis of either electron-rich or electrondeficient iodonium salts, as the reactivity of the arenes varies 80 dramatically with the electronic properties. 8 Aryl iodides 1 and arenes 2 were thus selected to investigate the formation of both electron-rich and electron-deficient salts, including heteroaryl salts.

The use of iodobenzene both as aryl iodide (1a) and arene (2b) yielded 4-iodophenyl(phenyl)iodonium triflate (3b) as a single 85 regioisomer (entry 2). Likewise, the reaction of 1a with toluene (2c) was smooth, and unsymmetrical salt 3c could be isolated in good yield (entry 3). The electron-rich arenes anisole (2d) and thiophene (2e) were, as expected, very reactive under the standard conditions. By decreasing the temperature to -78 °C, salts 3d and 3e 90 could be obtained in excellent yields (entries 4, 5). Heteroaryl salt 3e has previously been synthesized in several steps via the corresponding stannane, 18 which further illuminates the efficiency and environmental friendliness of our procedure.

Table 2 Synthesis of substituted diaryliodonium salts 3^a

	Ar ¹ -I + Ar ²	TfOH, CH	2Cl ₂ Ar ¹ Ar ² Ar ²	Tf
Entr	y1 (Ar ¹ I)	2 (Ar ² H)	Salt 3 ^b	Yield
1^d	1a (PhI)	2a (PhH)	3a	(%) ^c 85
2	1a (PhI)	2b (PhI)	3b OTF	85
3	1a (PhI)	2c (PhMe)	3c OTf	85
4	1a (PhI)	2d (PhOMe)	3d OMe	87
5	1a (PhI)	2e (thiophene)	3e S OTH	82
6^d	1b (4-MePhI)	2a (PhH)	3c	71
7	1b (4-MePhI)	2c (PhMe)	3f OTf	52
8	1c (2-MePhI)	2a (PhH)	3g OTf	85
9	1c (2-MePhI)	2c (PhMe)	3h OTT	87
10	1d (4-NO ₂ PhI)	2a (PhH)	O ₂ N 3i OTf	85
11^d	1e (2-chloro-5-iodo-pyridine)	2a (PhH)	CI N 3j OTF	60
12 ^d	1e (2-chloro-5-iodo-pyridine)	2d (PhOMe)	CI N 3k OMe	53

^a Reaction conditions: 1 (1.0 equiv), 2 (1.1 equiv), mCPBA (1.1 equiv) and TfOH (2.0 equiv) in CH2Cl2, see Table 1 and ESI for details. ^b Formed with complete regioselectivity. ^c Isolated yield. ^d 3.0 equiv of TfOH was used.

Unsymmetrical salt 3c could also be formed by the reaction of 4-95 iodotoluene (1b) with benzene, as shown in entry 6. Aryl iodide 1b was also employed to form the symmetrically substituted bis(4methylphenyl)iodonium salt 3f (entry 7). The reason for the moderate yield of 3f is unclear, as 2-iodotoluene (1c) delivered salts 3g and 3h in high yields upon reaction with benzene and 100 toluene, respectively (entries 8, 9).

Generally, the synthesis of electron-rich salts was easier from iodobenzene and a substituted arene than from the "reverse" reaction of a substituted aryl iodide with benzene (compare entries 3 and 6). 4-iodoanisole and 2-iodothiophene both participated in the reaction with benzene, but salts 3d,e were contaminated with unidentified byproducts.

Table 3 Direct synthesis of salts 3 from arenes and iodine^a

	4 Ar-H •	+ l ₂	mCPBA, TfOH, Cl	→ 2 A / \	OTf
Entr	y 2 (ArH)	T (°C)	Time	Salt 3 ^b	Yield (%) ^c
1	2a (PhH)	80	10 min	3a	78
2	2a (PhH)	rt	22 h	3a	81
3	2c (PhMe)	rt	2 h	3h:3f ratio 3:1	52
4	2c (PhMe)	0	1 h	3h	31
5	2f (PhCl)	rt	21 h	CI 31 CI	57
6	2g (4-nitro- m-xylene)	rt	19 h	3m NO ₂	24

^a Reaction conditions: I₂ (1.0 equiv), **2** (4-10 equiv), mCPBA (4 equiv) and TfOH (4 equiv) in CH₂Cl₂, see ESI for details. ^b Formed with complete regioselectivity apart from entry 3. c Isolated yield.

Electron-deficient iodonium salts were, on the other hand, best made from the corresponding electron-deficient aryl iodide with benzene as the arene. This is exemplified by the reaction of iodo-4-110 nitrobenzene (1d) with benzene, which delivered salt 3i in 85% yield (entry 10). The reverse reaction with iodobenzene and nitrobenzene was sluggish and gave salt 3b as a byproduct, the result of competing reaction pathways of 1a between oxidation and reaction with the formed iodine(III) intermediate.

The generality of the developed protocol is especially evident from the successful reaction of 2-chloro-5-iodopyridine (1e) with benzene or anisole to give salts 3j and 3k, respectively (entries 11, 12). Iodonium salts containing this aryl moiety have recently been used in an efficient total synthesis of (-)-Epibatidine.⁴ Pyridyl iodonium salts have previously been inaccessible by acidic routes, and salt 3j was formerly obtained by a basic, atom-inefficient twostep procedure¹⁹ in moderate yield, which was still inadequate for the preparation of salt 3k.²⁰ The formation of iodonium salts 3 from substituted arenes 2 was in all cases highly regioselective, yielding 125 salts **3b-d,f,h,k** with complete *para*-selectivity. Likewise, the reaction of 1a with thiophene (2e) afforded only 2-substituted 3e.

Having succeeded in the direct synthesis of diaryliodonium salts from aryl iodides, we looked into possible extensions of this reaction. As aryl iodides are readily available but often expensive, we argued that it might be possible to form the aryl iodide in situ.²¹ Hence, the reaction of benzene (2a) and iodine with mCPBA and TfOH was studied, and indeed delivered iodonium salt 3a in high 190

yield in only 10 min at 80 °C or 22 hours at room temperature (Table 3, entries 1, 2).

This efficient synthesis of diaryliodonium salts was subsequently applied to other arenes. Toluene (2c) yielded a mixture of salts 3h and **3f** with 3:1 regioselectivity favouring *ortho*-iodination (entry 3). The regioselectivity was higher at lower conversions, and pure **3h** was obtained after one hour at 0 °C (entry 4). Chlorobenzene 140 (2f) gave symmetric salt 3l with complete para-selectivity (entry 5). Even highly functionalized, deactivated arene 2g participated in the reaction to give salt 3m (entry 6). As this one-pot reaction involves several consecutive steps and many possible sources of byproducts, it is surprising that salts 3 are easily isolated in 145 moderate to good yields.

In conclusion, a facile, direct synthesis of diaryliodonium triflates from the corresponding aryl iodide and arene has been realized. The method is fast, high yielding, operationally simple and has a large substrate scope. Electron-rich salts are conveniently 150 synthesized from iodobenzene and the corresponding arene, and electron-deficient salts are formed by the reaction of a substituted aryl iodide with benzene. The protocol can be extended to the synthesis of iodonium salts directly from iodine and arenes, conveniently circumventing the need for aryl iodides. A thorough 155 investigation of the scope and limitations of this reaction is underway, and will be reported separately.

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Chem., 1980, 45, 1543-1544.

Graphical abstract:

Unsymmetric and symmetric diaryliodonium triflates are synthesized from both electron-deficient and electron-rich substrates in a fast, high yielding, and operationally simple protocol employing arenes and aryl iodides or iodine.