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The electrosynthesis of highly encumbered biaryls from aryl *o*-iodobenzyl ethers by a radical to polar crossover sequence†‡

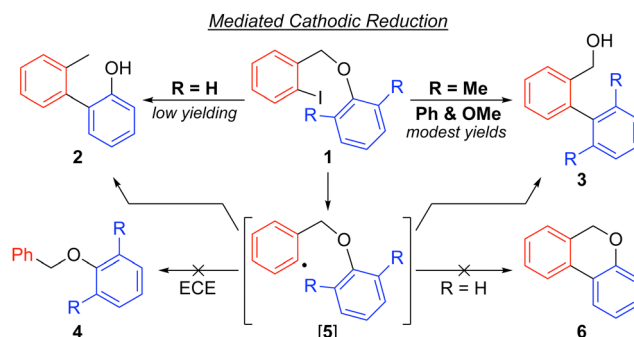
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Highly encumbered 2,2',6-tri- and 2,2',6,6'-tetra-substituted biaryls are readily prepared from aryl *ortho*-iodobenzyl ethers through mediated cathodic reduction under flow. The reaction proceeds via the stepwise transfer of two electrons: the first to induce loss of iodide and a radical cyclisation, and the second to induce a polar fragmentation.

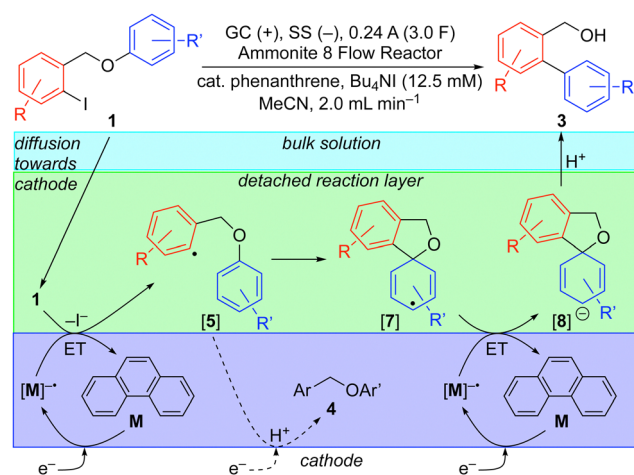
Benzyl *ortho*-iodoaryl ethers are commonly used as precursors to benzo[*c*]chromenes, *e.g.* **1** → **6**, through cyclisations induced by palladium catalysis,¹ or radical formation.^{2,3} Of these, the former usually proceed in high yield through *ortho*-cyclisation. In contrast, reactions *via* radical intermediates usually give cyclisation to **6** in low to modest yield due to competing processes such as iodide reduction to **4** or *ipso*-cyclisation and fragmentation (Scheme 1).² Cathodic reductions of aryl iodides also proceed *via* the corresponding radical intermediate.⁴ However, at the highly negative cathode potentials required, reduction of aryl radicals is extremely facile such that direct cathodic ECE reduction to the aryl anion is usually observed.^{5–7}

We recently showed how reductive radical cyclisation reactions of aryl halides could be effected electrochemically in flow using a strongly reducing catalytic mediator.^{5,6} The role of the mediator was shown to be key in ensuring that the generated aryl radical intermediate was formed away from the cathode, in a region where the flux of mediator radical anion [M]^{•–} leaving the cathode intercepts the flux of substrate coming toward it.^{5,6} By analogy, we envisioned an extension of the method to cathodic reductions of benzyl *ortho*-iodoaryl ethers **1** where a radical cyclisation [5] → [7] might be followed by reduction to

[8] and a polar fragmentation to biaryl **3** (Scheme 2).⁸ Herein, we describe our realisation of that sequence, its scope and limitations (Scheme 1).



Scheme 1 Summary of mediated cathodic reductions of benzyl *o*-iodoaryl ethers.



Scheme 2 Use of phenanthrene as a mediator to control the sequenced addition of two electrons from the cathode to the substrate.⁹

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† In memory of Prof. Pierre Duhamel.

‡ Electronic supplementary information (ESI) available: Experimental accounts with spectral details and copies of NMR spectra. CCDC 2359638 and 2363618. See DOI: <https://doi.org/10.1039/d4cc06061j>



Table 3 Isolated yields of biaryls **3j-x** from the mediated electrochemical radical-to-polar crossover reactions of 2-iodobenzyl aryl ethers **1j-x**⁹

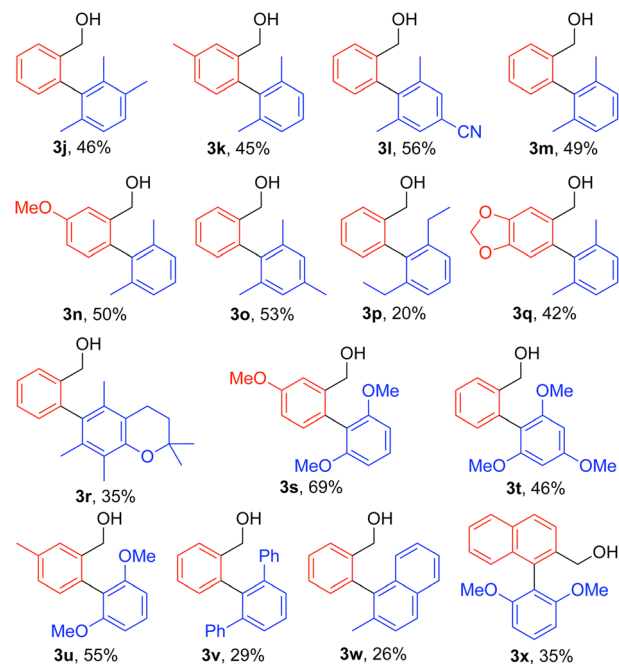
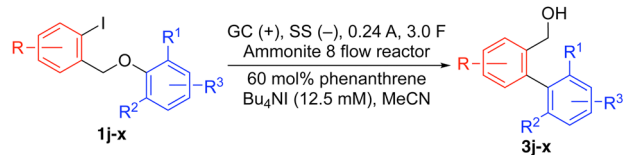
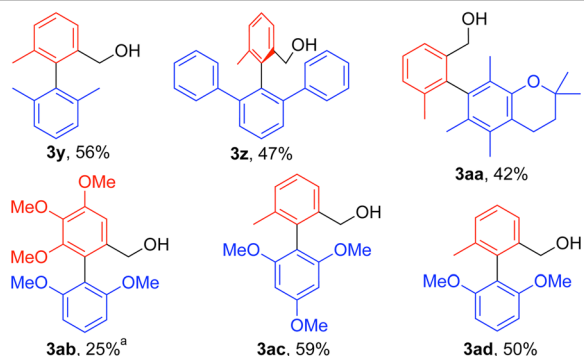
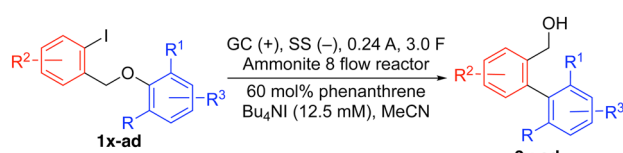


Table 4 2,2',6,6'-substituted biaryls formed by electrochemical radical-to-polar crossover reactions⁹



^a From the corresponding bromide.

3-fluoro-2-(*o*-tolyl)-phenol **19**, implicating an unprecedented reductive rearrangement of spirocyclic radical intermediate **7ae** to 6*H*-benzo[*c*]chromene **18**. Although biaryl **19** was formed as an oil, its identity was confirmed by X-ray analysis using the

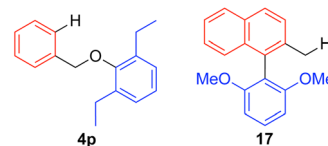
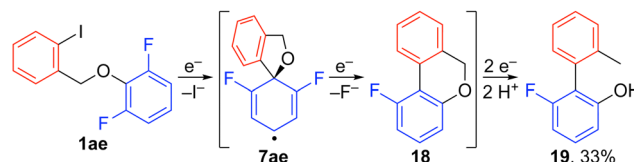


Fig. 1 By-products evidencing ECE reduction, product iodination and product reduction as competing side reactions.



Scheme 4 An unexpected rearrangement leading to 3-fluoro-2-(*o*-tolyl)-phenol **19**.

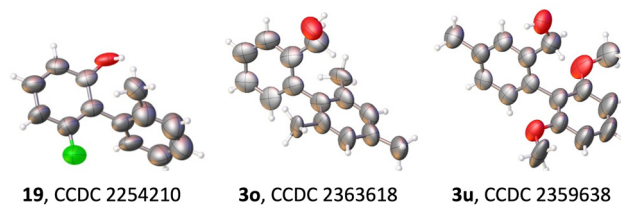


Fig. 2 Representative examples of X-ray analysis undertaken using the crystalline sponge method.^{12c}

recently introduced crystalline sponge technique,¹² which proved applicable to all of the biaryls surveyed (Fig. 2 and ESI†).^{12c}

In conclusion, mediated electrochemistry provides a means to control the rate at which sequential electron additions to a substrate occur. By slowing the transfer of a second electron to the substrate, the transient aryl radical has time to react with the proximal arene. From a synthetic perspective, the method provides rapid access to highly substituted biaryls, including 2,2',6,6'-tetrasubstituted biaryls, in modest yield. Notably, flow electrochemistry is widely seen as an emerging sustainable method and the required aryl *o*-iodobenzyl ethers are easy to prepare at low cost (as detailed in the ESI†). We are currently looking to develop further reductive electrochemical radical-to-polar crossover, and higher, cascade reaction sequences.

James E. Pearce conducted the bulk of the experimental work with support from Jack Hodgson, Ana Folguez-Amador, Johanna Fish and Philip Parsons. Crystalline Sponge X-ray analyses were conducted by Robert Carroll under the supervision of Simon Coles. The corresponding authors conceived of, and supervised, the project as a whole. We gratefully acknowledge financial support from EPSRC [EP/P013341/1, EP/W02098X/1 and EP/K039466/1], Pareon Chemicals Ltd and the European Regional Development Fund [ERDF Interreg Va programme (Project 121)].

Data availability

Experimental accounts and MP, IR, ¹H NMR, ¹³C NMR, LRMS, HRMS and X-ray data have been included, where applicable, in



the ESI[±] for both the products and starting materials detailed herein. These data include copies of the recorded ¹H and ¹³C NMR spectra.

Conflicts of interest

There are no conflicts of interest to declare.

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