Hypervalent iodine(III)-induced oxidative [4+2] annihilation of o-phenylenediamines and electron-deficient alkynes: direct synthesis of quinoxalines from alkyn e substrates under metal-free conditions†

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Hypervalent iodine(III)-induced oxidative [4+2] annihilation of o-phenylenediamines and electron-deficient alkynes under metal-free conditions has been developed. The reaction allows for direct access to quinoxalines bearing two electron-withdrawing groups in an efficient manner.

Quinoxaline derivatives not only constitute an important class of biologically active agents,1 but also find tremendous applications in materials science such as luminescent materials2 and low-band-gap polymers.3 Of the reported methods,4 the most widely used approach involves condensation of o-phenylenediamines with 1,2-dicarbonyl compounds bearing electron-rich or neutral substituents, which are generally prepared by oxidation of upstream alkynes [Scheme 1a]. On the other hand, the synthetic methods of the quinoxalines bearing electron-withdrawing groups (EWGs, e.g., –COR, –CO2R, –SO2R) have been poorly explored,5 although such compounds can serve as promising candidates for (opto)electronic materials6 and as versatile synthetic intermediates. Herein we present a hypervalent iodine(III) reagent-induced oxidative [4+2] annihilation of o-phenylenediamines and electron-deficient alkynes [Scheme 1b], which allows for direct access to electron-deficient quinoxalines from alkynes instead of diketo substrates in an efficient manner.

Recently, we have reported an oxidative dimerization of anilines through the agency of a unique and powerful iodinating reagent, tert-buty l hypiodide ([t-BuO]I), leading to aromatic azo compounds in an efficient and selective manner.7 The key to the success is an efficient two-fold iodination of nitrogen-center, forming ArNI2 species which then serves as an electrophile to form N–N bonds. Based on the findings, we envisioned that a tandem process consisting of (i) the Michael-addition of o-phenylenediamine to an electron-deficient alkyne and (ii) a subsequent nucleophilic attack on the highly electrophilic N-center (NL) by the resulting enamine could form a dihydro-quinoxaline skeleton. The subsequent elimination of HI would produce quinoxalines. At the outset, we attempted the oxidative annulation of o-phenylenediamine (1a) and DMAD (2a) as a model reaction (Table 1). However, contrary to our preliminary expectation, the results were disappointing: the treatment of an equimolar mixture of 1a and 2a with t-BuOI (4 equiv.) at –20 °C gave cis,cis-mucononitrile (5) as a major product, which should be formed through oxidative dearomatization and the following C–C bond cleavage of the benzene core (entries 1 and 2).8 These results clearly suggested that t-BuOI is not an appropriate oxidant for the aimed transformation, probably due to the rapid H–I exchange and dearomatization processes of 1a prior to Michael-addition. After extensive screening of iodine-containing reagents, we were delighted to find that the employment of phenyliodine diacetate (PIDA) was highly effective for the progression that leads to quinoxaline, has not been reported to date.12 Intriguingly, a significant solvent effect was observed (entries 3–6): as the polarity of solvent increased, the yield of 3a was enhanced while that of the by-product 4a13 was decreased.14 Other representative
hypervalent iodine(III) reagents were found ineffective for the annulation (entries 7–10), and PIDA was indispensable for the annulation (entry 11).

Having optimized the reaction conditions, the scope of the oxidative [4+2] annulation was investigated (Table 2). A wide variety of diamines bearing an electron-rich, -neutral, and -deficient substituent reacted with DMAD to give corresponding quinoxalines 3b–3j in high yields. Multiple-substituted diamines afforded 3k and 3l. Furthermore, sterically demanding diamines gave the corresponding product 3m in excellent yield. Although the reaction of naphthalene-2,3-diamine with DMAD required prolonged time, N-heteroacene 3n, which constitutes a family of electron-transporting materials,15 was obtained in 56% yield. Using the method, biquinoxaline 3o was prepared in good yield. In respect to alkyne substrates, dibutyl acetylenedicarboxylate was successfully applied to the reaction conditions to afford 3p and 3q in 77% and 44% yield, respectively. In addition, an unsymmetrical alkyne having an ester and a sulfonyl group also successfully underwent the annulation to give 3r in good yield.

Taking advantage of the ester functionality, 3a was diversified derivatized into functionalized quinoxalines (Scheme 2). For example, diester of 3a easily underwent hydrolysis to give dicarboxylic acid 6 in high yield, which was further efficiently converted to 7 by dehydration. Moreover, anhydride 7 was successfully transformed into imide-fused quinoxaline 8 by condensation with p-toluidine, which is an N-analogue of triboluminescent material.16 It is noted that such compounds are quite difficult to prepare by traditional condensation methods.17

To investigate the reaction pathways, several experiments were conducted as follows: enamine 9, which was readily prepared by the Michael-addition of N-Boc-protected o-phenylenediamine to DMAD,13 was treated with PIDA in the presence of trifluoroacetic acid (Scheme 3).18 At −20 °C, 9 underwent oxidative cyclization to give N-Boc dihydroquinoxaline 10 in 45% yield,19 while 9 was quantitatively recovered in the absence of PIDA. In contrast, at room temperature, quinoxaline 3a was obtained in 54% yield. In conjunction with the fact that DMAD does not react with PIDA in the absence of o-phenylenediamine, the most likely intermediate of the annulation would be the deprotected counterpart of the Michael-adduct 9 as preliminary assumed.

On the basis of the experimental results and knowledge accumulated from the literature about hypervalent iodine(III)-mediated oxidative C–N bond forming reactions using enamine substrates,20–22 conceivable reaction pathways are illustrated in Scheme 4. The reaction would start with Michael addition

### Table 1 Summary of the screening of the reaction conditions

| Entry | Oxidant (equiv.) | Solvent | T [°C] | 3a [%] | 4a [%] | 5 [%] |
|-------|-----------------|---------|--------|--------|--------|-------|
| 1     | t-BuOI (4)      | THF     | −20    | 12     | 0      | 64    |
| 2     | t-BuOI (4)      | DME     | −20    | 2      | 0      | 33    |
| 3     | PhI(OAc) 2 (2)  | CH 2Cl 2| −20    | 5      | 40     | 0     |
| 4     | PhI(OAc) 2 (2)  | THF     | −20    | 63     | 18     | 0     |
| 5     | PhI(OAc) 2 (2)  | DME     | −20    | 60°    | 4c     | 0     |
| 6     | PhI(OAc) 2 (2)  | DMF     | −20    | 92°    | 4c     | 0     |
| 7     | PhI−O (2)       | DMF     | −20    | 3      | 0      | 0     |
| 9     | HO (2)          | DMF     | −20    | 0      | 0      | 0     |
| 10    | PhI(OH)OTs (2)  | DMF     | −20    | 0      | 0      | 0     |
| 11    | —               | DMF     | −20    | 21     | 0      | 0     |

* Reaction conditions: 1a (0.25 mmol), 2a (0.25 mmol), and iodine-containing oxidant (0.50–1.0 mmol) were mixed in a solvent (3 mL) at the temperature in the column and stirred for 24 h. | (Isolated yield. | NMR yields. |

### Table 2 Scope of the oxidative [4+2] annulation

| R       | Product          | Yield [%] |
|---------|------------------|-----------|
| N-H     | 3a               | 92%       |
| N-Me    | 3b               | 94%       |
| N-Boc   | 3c               | 92%       |
| N-OMe   | 3d               | 90%       |
| N-CF 3   | 3e               | 87%       |
| N-CN   | 3f               | 87%       |
| N-CN   | 3g               | 85%       |
| N-CN   | 3h               | 83%       |
| N-CN   | 3i               | 80%       |
| N-CN   | 3j               | 80%       |
| N-CN   | 3k               | 80%       |
| N-CN   | 3l               | 80%       |
| N-CN   | 3m               | 77%       |

* Reaction conditions: 1 (0.25 mmol), 2 (0.25 mmol), and PhI(OAc) 2 (0.50 mmol) were mixed in DMF (3 mL) at 20 °C and stirred for 24 h. | The values in parentheses indicate the yields of quinoxaline products. | Reaction time: 48 h. | Yield [%] | NMR yields. |

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**Scheme 2** Derivatization of 3a.
of o-phenylenediamine to DMAD, forming the enamine intermediate A, which has three possible reactive points when reacting with PIDA, namely, β-carbon of enamine (a),\(^{20}\) enamine nitrogen (b),\(^{21}\) and nitrogen on the benzene moiety (c).\(^{22}\)

Accordingly, three species should be extrapolated as intermediates prior to cyclization: (i) α-iodo(III) imine B (route a); (ii), (iii) enamines C and D (routes b and c, respectively). Successive cyclizative nucleophilic substitution on the iodine-attached sp³-carbon (from B), on the enamine carbon in a pseudo-S_{N}2’ manner (from C), or on the electrophilic N-center (from D) would provide a common intermediate E. Oxidative aromatization of E with another equivalent of PIDA should lead to quinoxaline F. Ma and Lei reported an oxidative dimerization of aromatic amines using PIDA to give azobenzenes.\(^{23}\) Noazo compounds were detected in our system, suggesting that the pathway via intermediacy of D (route c) might be excluded. On the one hand, according to the reactivity of carbonyl-conjugated CF₃CH₂I(OH)(OTs),\(^{24}\) pseudo-S_{N}2.\(^{22}\)

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