

COMMUNICATION

Electrochemical Syntheses of Aryl-substituted Benzothiophenes and Phenanthrenes Using Benzenediazonium Salts as the Aryl Radical Precursors

Li-Yuan Lan ^{a,#}, Yang-Ye Jiang ^{a,#}, Raymond Daniel Little ^b, Cheng-Chu Zeng ^{a,*}

^a Faculty of Environment & Life, Beijing University of Technology, Beijing 100124, China

^b Department of Chemistry & Biochemistry, University of California, Santa Barbara, CA 93106, USA

Abstract

Aryl-substituted benzothiophene and phenanthrene are important structural units in medicinal chemistry and materials science. Although extensive effort has been devoted to prepare these compounds and a variety of approaches have been developed to construct the 2-substituted benzothiophene core structure, environmental-friendly and efficient synthetic means are still desired. Based on our previous electrochemical Minisci-type arylation reaction with aryl diazonium salt as the aryl precursor, as well as the work from König's group, herein, we described the use of paired electrolysis to achieve 2-aryl benzothiophenes and 9-aryl phenanthrenes employing benzenediazonium salts as the aryl radical precursors. Initially, 2-methylthiobenzendiazonium salt 1a and 4-methylbenzene ethyne 2a were chosen as the model substrates to optimize the reaction conditions by examining solvent, supporting electrolyte, electrode material and current density. After extensive efforts, it was found that an 89% yield of the desired product 3a was afforded in an undivided cell equipped with a graphite felt anode and a Ni plate cathode, using $n\text{-Bu}_4\text{NBF}_4$ as the supporting electrolyte and DMSO as the solvent, while operating at a constant current density of $4\text{ mA}\cdot\text{cm}^{-2}$. Under the optimal conditions, the generality of the electrochemical protocol and substrate scope were then examined. The results showed that both alkyl acetylene and aryl acetylene could be applied in this method, and a series of aryl-substituted benzothiophene derivatives were obtained successfully. Considering the wide range of application of phenanthrene molecules in medicinal chemistry and materials science, we then applied this protocol to the synthesis of phenanthrene derivatives, and succeeded in obtaining the corresponding 9-arylphenanthrene derivatives. Finally, cyclic voltammetric (CV) measurement was conducted to analyze the possible mechanism. It was found that 2-methylthiobenzene diazonium salt 1a gave a significant irreversible reduction peak at $-0.4\text{ V vs. Ag/Ag}^+$ in CH_3CN , whereas no signal was detected for phenylacetylene 2a in the scanning potential window. In addition, the presence of 2a did not alter the peak potential of 1a, albeit the peak current increased slightly. These results indicate that the reduction of 1a is easier than that of 2a. Based on our CV analysis and previous photocatalytic results, a sequential paired electrolysis mechanism is proposed, that is, the electrochemical reduction of benzenediazonium salt 1a at the cathode produces aryl radical 5a, which is then added to phenylacetylene to produce vinyl radical 6a and sulfonyl radical 7a following an intramolecular cyclization. Finally, the anodic oxidation of 7a, followed by demethylation with DMSO, generates the target product 3a. In summary, we have developed a paired electrolysis method for the syntheses of 2-arylbenzothiophene derivatives and 9-arylphenanthrene derivatives. The protocol features wide substrate scope and functional group tolerance, which further demonstrates that the practicability of aryldiazonium salts as versatile aryl radical sources to generate aryl radicals through electrochemical reduction.

Keywords: Benzothiophenes; Phenanthrenes; Paired electrosynthesis; Aryldiazonium salts; Aryl radicals

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[#] These authors contributed equally to this work.

* Corresponding author, Cheng-Chu Zeng, Tel: (86-10)67391609, E-mail address: zengcc@bjut.edu.cn.

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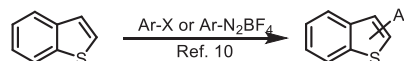
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1. Introduction

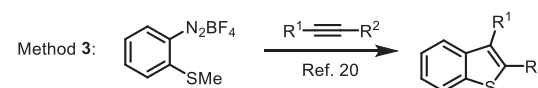
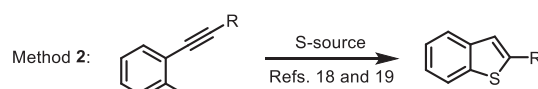
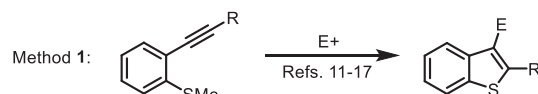
2-Substituted benzothiophenes are prominent structural features in natural products and organic molecules with diverse biological activities, including anti-tumor, anti-oxidant, anti-tubercular and antimicrobial activities [1–4]. For example, FDA-approved drugs, Raloxifene, Arzoxifene and Zileuton, contain these subunits (Fig. 1). It has also been of considerable interest in materials chemistry to introduce this subunit in organic solar cells, organic light-emitting diodes, and semiconductors [5–9]. Phenanthrenes are also important organic compounds with wide range of applications in medicinal chemistry and materials science [10–12].

Consequently, extensive effort has been devoted to prepare these compounds and a variety of approaches have been developed to construct the 2-substituted benzothiophene core structure. Intermolecular cross-couplings of benzothiophenes with halogenated hydrocarbon or aryl diazonium salts are frequently employed, although this method often generates C2- and C3-regioisomers (Fig. 2A) [13,14]. The cyclization approaches illustrated in Fig. 2B are more attractive since only the desired regioisomers are delivered. Among the cyclization approaches, the electrophilic cyclization of *o*-alkynyl thioanisoles has proven to be particularly effective (Fig. 2B, Method 1). For example, in 2016, Ingleson [15] and Blum [16] independently reported the electrophilic cyclization of *o*-alkynyl thioanisoles to give 2-aryl-3-Bpin substituted benzothiophenes using BCl_3 or ClBcat as the electrophilic reagents in the presence of pinacol. The uses of iodine, NBS, or NCS as the

A. Intermolecular coupling for the synthesis of 2-substituted benzothiophenes



B. Cyclization approaches for the synthesis of 2-substituted benzothiophenes



C. Electrochemical synthesis of 2-substituted benzothiophenes

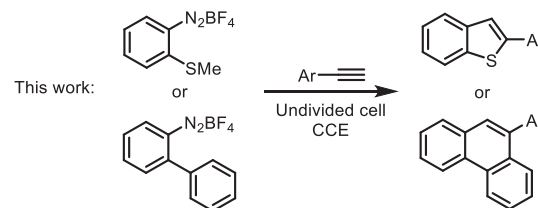
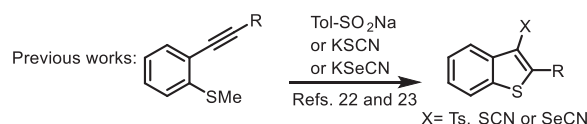
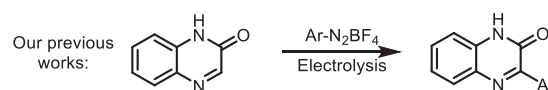


Fig. 2. The syntheses of 2-substituted benzothiophenes.

electrophilic reagents also afforded 2-amino benzothiophenes [17]. In addition, metal-catalyzed electrophilic cyclization of 2-alkylthio- or 2-tri-alkylsilyl-thio alkynyl benzene has also been used

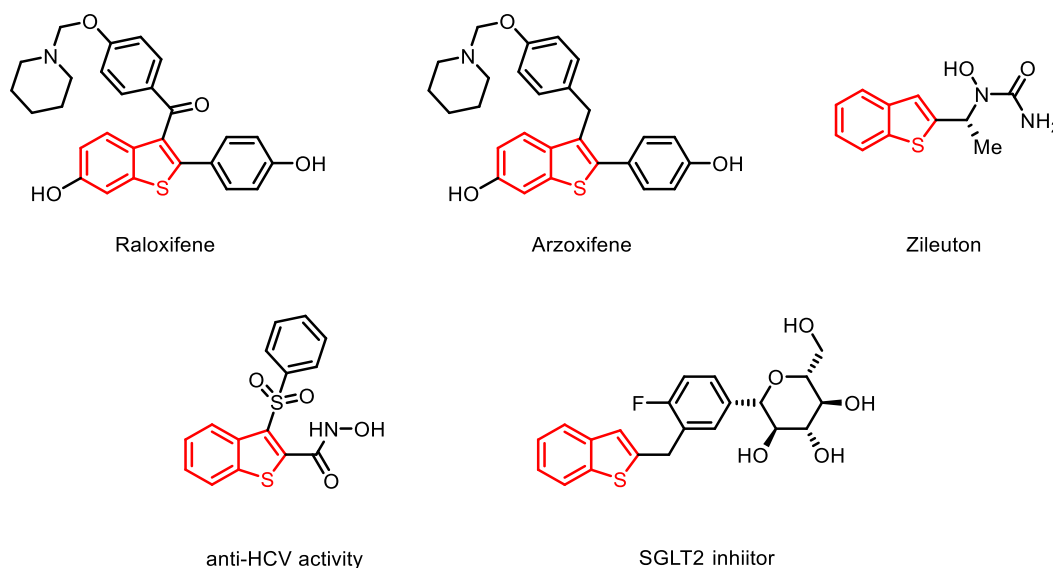


Fig. 1. Representative structures containing the benzothiophene framework.

as the starting materials to generate 2-aryl benzothiophenes [18–21].

Metal-mediated annulation of 2-halogenated alkylbenzenes in the presence of sulfur sources constitutes another approach to 2-benzothiophenes (Fig. 2B, Method 2). Paradies et al. [22] achieved the synthesis of benzothiophenes through a Pd-catalyzed reaction of 2-bromo arylolefinyl benzenes with thiourea. Using Na₂S as a source of sulfur and CuI as a catalyst, Zhang synthesized 2-aryl benzothiophenes in high yields [23].

In addition to these pathways, radical annulation of omethylthio-diazonium salts with alkynes also affords 2-substituted benzothiophenes (Fig. 2B, Method 3). In 2012, König [24] utilized Eosin Y as a photoredox catalyst to generate 2-substituted benzothiophenes via a sequence of intermolecular aryl radical addition to the alkyne, followed by intramolecular cyclization of an sp² C-centered radical to a pendant methylthio group and oxidative demethylation.

In order to demonstrate that a paired electrolysis provides an alternative to the visible light photoredox-based oxidation approach, we have previously developed an efficient electrochemical approach to the Minisci-type arylation reaction using aryldiazonium salts as the aryl radical precursors [25]; an example of our approach is illustrated in Fig. 2C. With these results in mind and

inspired by the work of König [24], we envisioned that 2-methylthio aryldiazonium salts may also undergo radical annulation under electrochemical conditions to construct 2-substituted benzothiophenes via a paired electrolysis.

Similar radical annulation reactions may also give 9-aryl phenanthrene when biphenyldiazonium salts are used as the radical precursor (Fig. 2C). For the research described herein, a paired electrolysis was carried out under galvanostatic conditions in a simple undivided cell without the need for an external supporting electrolyte. The chemistry is applicable to a wide range of substrates and the pathway is easy to be scaled-up. The results further demonstrate that photoredox catalyzed reactions may also be achievable under electrochemical conditions. Notably, while preparing the manuscript, the Ts-, NCS- or NCSe-substituted benzothiophenes have recently been synthesized from the electrochemical oxidations of o-alkynyl thioanisoles with TsSO₂Na [26], and KSCN or KSeCN [27], respectively (Fig. 2C).

2. Results and discussion

We began our search for optimal conditions by investigating the model reaction of 2-methylthiobenzendiazonium salt **1a** with 4-methylbenzene ethyne **2a**. As shown in Table 1, when a mixture of

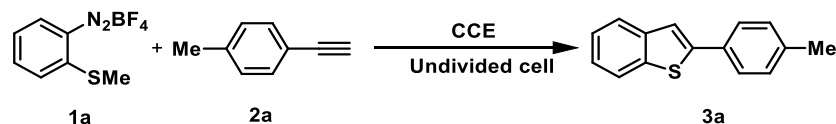


Table 1. Optimization of reaction conditions^a.

Entry	Solvent	Ratio of 1a to 2a	Supporting electrolyte	<i>J</i> (mA·cm ⁻²)	Anode material	Yield ^b
1	DMSO	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	37
2	MeOH	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	Trace
3	CH ₃ CN	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	8
4	DCE	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	0
5	DMF	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	15
6	DMSO/MeOH	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	18
7	DMSO/HFIP	1:2	<i>n</i> -Bu ₄ NBF ₄	4	C felt	3
8	DMSO	1:3	<i>n</i> -Bu ₄ NBF ₄	4	C felt	89 (87) ^c
9	DMSO	1:5	<i>n</i> -Bu ₄ NBF ₄	4	C felt	69
10	DMSO	1:3	Et ₄ NBF ₄	4	C felt	39
11	DMSO	1:3	Me ₄ NBF ₄	4	C felt	49
12	DMSO	1:3	<i>n</i> -Bu ₄ NPF ₆	4	C felt	44
13	DMSO	1:3	Et ₄ NHSO ₄	4	C felt	49
14	DMSO	1:3	Et ₄ NCIO ₄	4	C felt	43
15	DMSO	1:3	LiClO ₄	4	C felt	9
16	DMSO	1:3	none	4	C felt	39
17	DMSO	1:3	<i>n</i> -Bu ₄ NBF ₄	6	C felt	54
18	DMSO	1:3	<i>n</i> -Bu ₄ NBF ₄	2	C felt	91 (6 hours)
19	DMSO	1:3	<i>n</i> -Bu ₄ NBF ₄	4	Graphite plate	25
20	DMSO	1:3	<i>n</i> -Bu ₄ NBF ₄	4	Pt wire	30

^a Conditions: **1a** (0.3 mmol) and **2a** in 3 mL solvent in a beaker-type undivided cell equipped with Ni plate cathode.

^b Yield determined by HPLC used 1,3,5-trimethoxybenzene as internal standard.

^c Isolated yield.

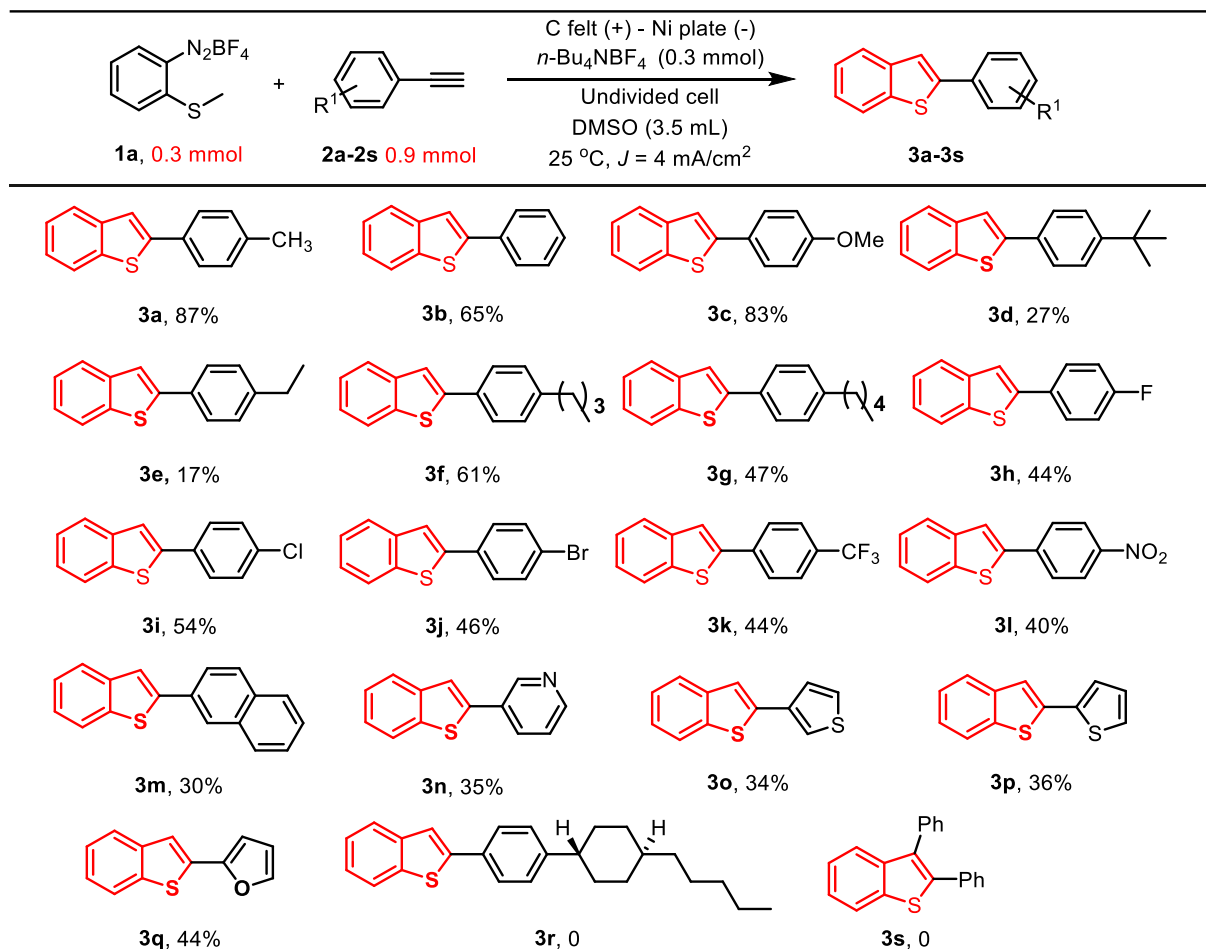
1a and 2 equiv of 2a in DMSO with *n*-Bu₄NBF₄ as the supporting electrolyte was electrolyzed in a beaker-type undivided cell equipped with a carbon felt anode and a Ni plate cathode, the corresponding benzothiophene 3a was produced in 37% yield (Entry 1). Replacing the solvent with MeOH, CH₃CN, DCE, and DMF, separately, or a mixed solution of DMSO with MeOH or HFIP gave inferior yields (Entries 2–7). Given the fact that 2a rapidly disappeared during the electrolysis, the amount of 2a was increased, this led to an 89% (87% isolated yield) or 69% yield of 3a in the presence of 3 or 5 equiv of 2a, respectively (Entries 8 and 9). Compared with the photocatalytic synthesis [24] of benzothiophene 2a, this electrochemical variation gave a higher yield (89% vs. 75%) and required fewer equivalents of 2a (3 equiv vs. 10 equiv).

The screening of supporting electrolyte indicated that *n*-Bu₄NBF₄ was superior, since other conducting salts, such as Et₄NBF₄, Me₄NBF₄, *n*-Bu₄NPF₆, Et₄NHSO₄, Et₄NClO₄ and LiClO₄ (Entries 10–15) gave less than 50% of yields. Notably, without external supporting electrolyte, 3a was still

produced although in a slightly lower yield. This is likely due to the conductivity of the starting material 1a (Entry 16). The reaction was less effective when the electrolysis was performed at 6 mA·cm⁻² (Entry 17). In the case of 2 mA·cm⁻², a nearly identical yield (91%) of 3a was observed, but twice as much time (6 hours) was required (Entry 18). Finally, we investigated the choice of anodic material on the reaction. When a graphite plate or a Pt plate was used as the anode, 25% or 35% yields of 3a were obtained, respectively (Entries 19–20). Based on the results described above, we concluded that the reaction was most efficient when conducted in an undivided cell equipped with a graphite felt anode and a Ni cathode at a constant current density of 4 mA·cm⁻² using *n*-Bu₄NBF₄ in DMSO as an electrolyte system (89% yield of 3a).

With the optimal reaction conditions in hand, we then examined the reaction generality and substrate scope using 2-methylthio-benzenediazonium 1a and a variety of phenyl acetylene derivatives. As shown in Table 2, phenyl acetylene and methoxy substituted phenyl acetylene worked well to give

Table 2. Syntheses of 2-Arylbenzothiophenes^{a,b}.



^a Reaction conditions: 1a (0.3 mmol), 2a–2s (0.9 mmol) in 3.5 mL of DMSO, undivided cell, 25 °C, *n*-Bu₄NBF₄ (0.3 mmol), current density of $J = 4 \text{ mA} \cdot \text{cm}^{-2}$, C felt anode and Ni plate cathode.

^b Isolated yield.

65% and 83% yields of corresponding products **3b** and **3c**, respectively. Alkyl-substituted phenyl acetylene derivatives were also useful, albeit in low yields (**3d** and **3e**) due to the messy reaction mixture, to moderate yields (**3f–3g**). Benzothio-phenene derivatives **3h–3l** were also formed in moderate yields. The presence of halogen or nitro groups provides a handle for late-stage functionalization. The compatibility of other aryl acetylenes with **1a** under the standard conditions was also examined. In the case of naphthyl-, pyridinyl-, thiophenyl- and furanyl acetylenes, the corresponding products **3m–3q** were delivered in 30%–44% yields, whereas **3r** and **3s** were not produced and the starting acetylenes **2r** and **2s** were consumed completely.

Considering the wide range of applications of phenanthrene containing molecules in medicinal chemistry and materials science, we next explored the paired electrolysis approach to the synthesis of phenanthrene derivatives. As listed in Table 3, when a mixture of biphenyldiazonium salt **1b** and aryl acetylene **2** was electrolyzed under the standard conditions, the corresponding 9-arylphenanthrene derivatives **4** were formed in moderate yields. For example, in the reactions with alkyl-substituted phenyl acetylenes, isolated yields of **4a–4d** ranged from 39% to 43%. Electron-withdrawing groups, such as F or CF₃ were also

tolerated, leading to **4e** and **4f**, although in a lower yield.

To gain mechanistic insights, cyclic voltammetry (CV) was employed. As shown in Fig. 3, phenylacetylene **2a** was not reducible in the scan range from 0.0 V to –1.0 V vs. Ag/Ag⁺ in CH₃CN, whereas 2-methylthio benzenediazonium salts **1a** gave an obvious irreversible reductive peak at –0.4 V vs. Ag/Ag⁺ in CH₃CN. When **2a** was present, the reduction peak potential **1a** did not

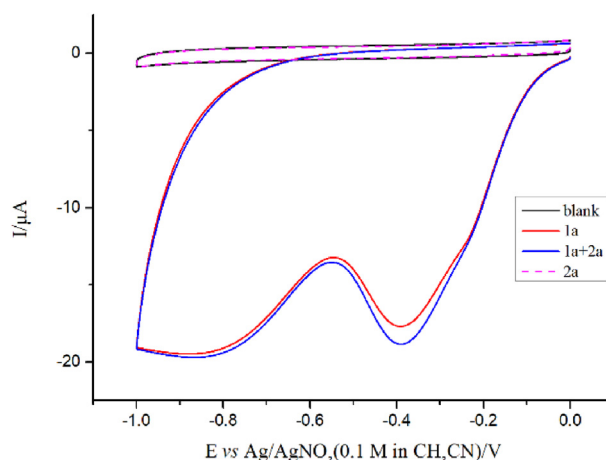
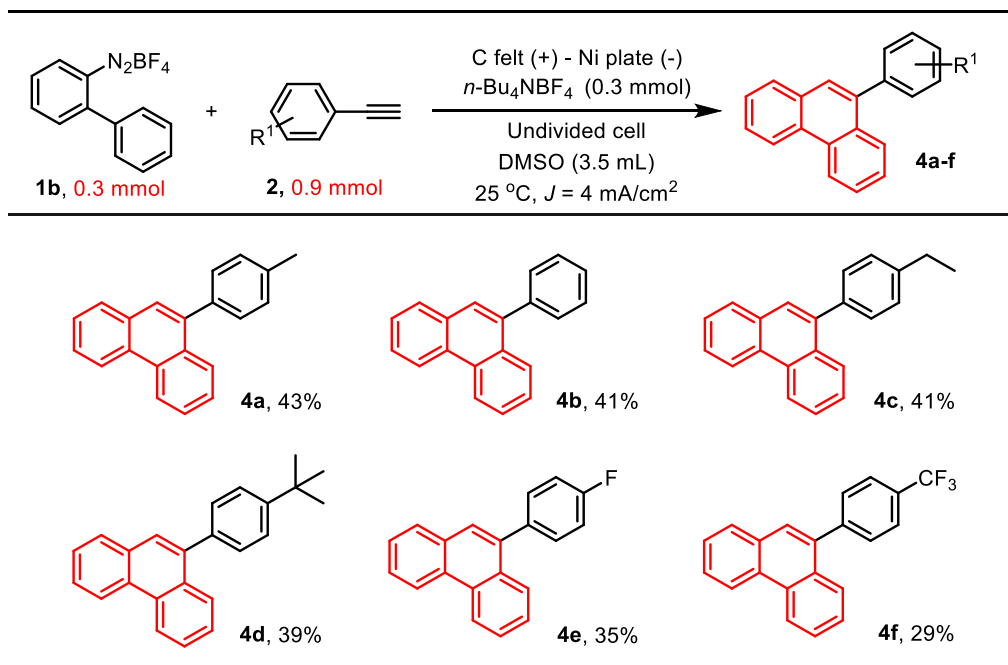


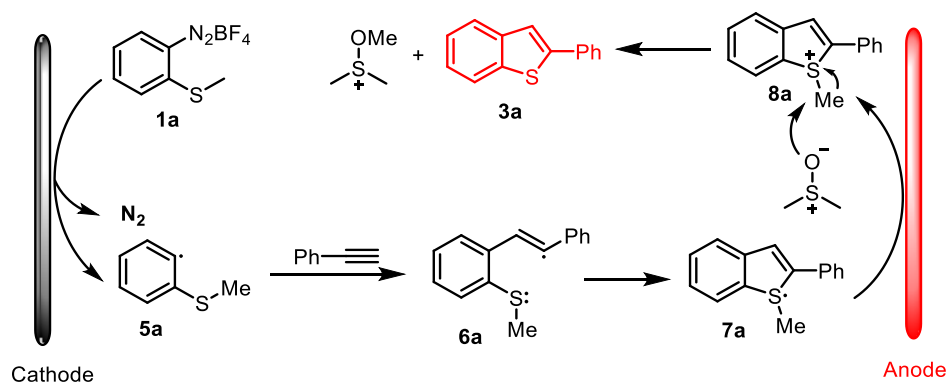
Fig. 3. Cyclic voltammograms of related compound (5 mmol·L⁻¹) in 0.1 mol·L⁻¹ Bu₄NBF₄/DMSO.

Table 3. Syntheses of 9-arylphenanthrene derivatives^{a,b}.



^a Reaction conditions: **1b** (0.3 mmol), **2** (0.9 mmol) in 3.5 mL of DMSO, undivided cell, 25 °C, *n*-Bu₄NBF₄ (0.3 mmol), current density of *J* = 4 mA·cm⁻², C felt anode and Ni plate cathode.

^b Isolated yield.



Scheme 1. Proposed mechanism for the paired electrochemical syntheses of benzothiophenes.

change, although there was a slight increase of peak current. These results indicate that electrochemical reduction of a mixture of **1a** and **2a** begins with the reduction of **1a**.

Based on our CV analysis and previous photocatalytic results [24], a sequential paired electrolysis pathway is proposed (Scheme 1). Taking the electro-syn-synthesis of **3a** as an example, the electrochemical reduction of benzenediazonium salt **1a** at the cathode affords the aryl radical **5a**, along with the release of nitrogen gas. The radical **5a** then adds to phenyl acetylene to generate vinyl radical **6a** and sulphydryl radical **7a** following an intramolecular cyclization and anodic oxidation of **7a**. Finally, the subsequent demethylation using DMSO leads to the target structure **3a**.

3. Conclusions

In summary, we have developed a paired electrolysis approach for the syntheses of 2-aryl benzothiophenes and 9-aryl phenanthrenes. The chemical principle is initiated from the cathodic reduction of benzenediazonium salts, and the synthesis is conducted in an undivided cell equipped with a graphite felt anode and a Ni plate cathode using $n\text{-Bu}_4\text{NBF}_4$ supporting electrolyte and DMSO solvent at the constant current density of $4\text{ mA}\cdot\text{cm}^{-2}$. The methodology features wide substrate scope and functional group tolerance. It further demonstrates the utility of the electrochemical reduction of aryldiazonium salts as a source of aryl radicals.

Acknowledgements

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以芳基重氮盐为芳基前体电化学合成芳基取代的苯并噻吩和菲

蓝丽媛^{a,#}, 蒋洋叶^{a,#}, R. Daniel Little^b, 曾程初^{a,*}

^a北京工业大学环境与生命学院, 北京 100124, 中国

^bDepartment of Chemistry & Biochemistry, University of California, Santa Barbara, California 93106, USA

摘要

芳基取代的苯并噻吩和菲是药物化学和材料科学中的重要结构单元。尽管已经投入了很多来制备这类化合物, 并且已经开发了多种方法来构建 2-取代的苯并噻吩核心结构, 但仍然需要环境友好且有效的合成方法。基于我们以前的以芳基重氮盐为芳基前体的电化学 Minisci 型芳基化反应, 以及来自 König 小组的工作, 本文使用成对电解以苯重氮盐为芳基前体来获得 2-芳基苯并噻吩和 9-芳基菲。首先, 选择 2-甲硫基苯重氮盐 1a 和 4-甲基苯乙炔 2a 作为模型底物, 通过考察溶剂、支持电解质、电极材料和电流密度来优化反应条件。经过大量的努力, 发现在装备有石墨毡阳极、镍片阴极的单室电解池中, 使用 *n*-Bu₄NBF₄ 作为支持电解质, DMSO 作为溶剂, 控制在 4 mA·cm⁻² 条件下恒流电解, 获得 89% 产率的所需产物 3a。在最佳条件下, 考察该电化学方案和底物范围的普适性。结果表明, 烷基乙炔和芳基乙炔都适用于该方法, 并成功地得到了一系列芳基取代的苯并噻吩衍生物。考虑到菲类化合物在药物化学和材料科学中的广泛应用, 将此方案应用到菲衍生物的合成中, 获得了相应的 9-芳基菲衍生物。最后, 通过循环伏安法分析了可能的机理。2-甲硫基苯重氮盐 1a 在 CH₃CN 中相对于 Ag/Ag⁺ 在 -0.4 V 处产生一个显著的不可逆还原峰, 而在扫描电位窗口中没有检测到苯乙炔 2a 的信号。此外, 尽管峰电流略有增加, 2a 的存在没有改变 1a 的还原峰电位。这些结果表明 1a 的还原比 2a 的还原容易。基于循环伏安分析和光催化结果, 我们提出了成对电解机制, 即芳基重氮盐 1a 在阴极的电化学还原产生芳基 5a, 然后芳基 5a 加成到苯乙炔上, 在分子内环化后产生乙烯基 6a 和磺酰基 7a。最后, 7a 的阳极氧化, 接着用 DMSO 脱甲基化, 生成目标产物 3a。总之, 我们发展了一种成对电解法来合成 2-芳基苯并噻吩衍生物和 9-芳基菲衍生物。该方案具有广泛的底物范围和官能团耐受性, 进一步证明了芳基重氮盐作为多用途芳基自由基来源通过电化学还原产生芳基自由基的实用性。

关键字: 苯并噻吩; 菲; 成对电合成; 芳基重氮盐; 芳基自由基