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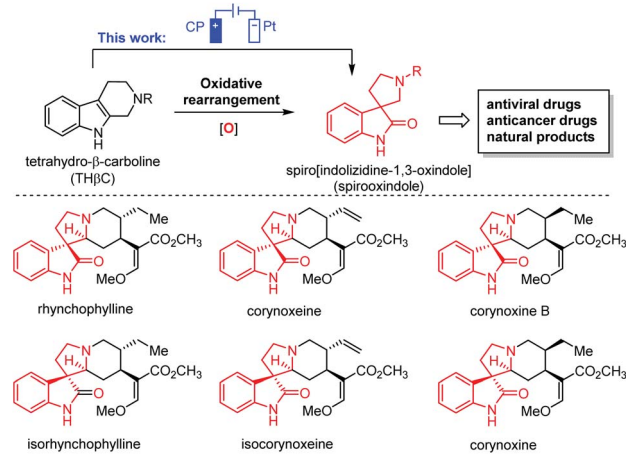
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Introduction

Oxidative rearrangement of tetrahydro- β -carbolines (TH β Cs)¹ (Fig. 1) was proposed as an enzyme-mediated reaction^{2–4} responsible for biosynthesis of many biologically active spiro [indolizidine-1,3-oxindole] (*i.e.*, spirooxindoles) natural products such as mitraphylline, rhynchophylline, (iso)corynoxine, corynoxine, corynoxine B, spirotryprostins A and B, and alstonisine (Fig. 1). With chemical oxidants [Pb(OAc)₄, OsO₄, *t*-BuOCl, *N*-bromosuccinimide (NBS), *etc.*] identified for such oxidative rearrangement, it has become a widely employed method for the construction of spiro[indolizidine-1,3-oxindole]^{5–11}, which is associated with significant biological activities (antiviral,¹² anticancer,¹³ and antibacterial¹⁴) and regarded as a pharmaceutically privileged structural motif.^{15–18} However, these chemical oxidants are hazardous and toxic and would generate stoichiometric amounts of harmful chemical waste. Recently, two green approaches using Oxone-halide¹⁹ and Fenton-halide²⁰ for the oxidative rearrangement of TH β Cs were

reported to reduce the impact on the environment and human health. While Oxone as an inorganic triple salt (KHSO₅·0.5KHSO₄·0.5K₂SO₄) has admirable bench stability, unique reactivity towards halides, and safe/easy operation, it generates two equivalents of potassium sulfate as a byproduct with an often-criticized high *E*-factor. On the other hand, hydrogen peroxide is widely considered to be an ideal terminal oxidant in organic synthesis since the only byproduct is water. However, hydrogen peroxide (typically 30 wt% in water was used in laboratory) is unstable at room temperature with an inconsistent concentration and quality and might decompose explosively. Herein, we report the first electrochemical method for



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Fig. 1 Oxidative rearrangement of tetrahydro- β -carboline and spirooxindole natural products.



the oxidative rearrangement of TH β Cs without using any chemical oxidant.

Electrochemical synthesis, a potentially sustainable and atom-economic means towards organic compounds, has received increasing attention in the past decade.^{21–24} The use of mediators for oxidation/reduction is one of the most effective strategies to achieve high regio- and chemo-selectivity through minimizing the direct non-selective oxidation/reduction of substrates on electrodes.^{25,26} Halides have emerged as one of the most successful mediators for selective electrochemical organic synthesis.²⁷ Mechanistically, the halide mediators NaX, KX, NH₄X or *n*-Bu₄NX (X[−] is Cl[−], Br[−] or I[−]) are oxidized at the anode into reactive halogenating species (RHS), which then react with substrates to generate an intermediate that can undergo further transformation(s) to yield a product. Inspired by this general mechanism, we envisioned that the electrochemically generated RHS could be used as an oxidant for the oxidative rearrangement of TH β Cs (Fig. 2(A)) because RHS generated *in situ* from chemical oxidation of halide (Oxone/halide or Fenton/halide) was successfully used for this reaction.^{19,20} The challenge is to minimize the competing nonselective anodic oxidation of TH β Cs over a halide because related direct electro-oxidation of indoles was reported by Oliveira-Brett,²⁸ Mount,²⁹ Vincent,^{30,31} Lei,³² Fang,³³ etc. Therefore, the selectivity between a halide and indole (*i.e.*, TH β Cs) is critical to the success of the oxidative rearrangement of TH β Cs.

We planned to address this selectivity challenge by (1) evaluating different electrodes and supporting electrolytes and (2) devising a flow cell. As we illustrate in Fig. 2(A), the oxidative rearrangement of TH β Cs should occur preferably at the diffusion layer but not at the double layer, where the choices of the anode and even the supporting electrolyte can determine the selectivity: the generation of RHS or the undesired nonselective oxidation. Secondly, a high-rate divided flow cell would

considerably reduce the substrate diffusion into the double layer for possible direct oxidation on the electrode, while it could achieve a desirable rapid transport of the halide towards the electrode for an adequate local concentration of RHS necessary for the rearrangement and at the same time avoiding concentration polarization that may result in side reactions.^{34,35} In this article, we disclose the results of our research efforts to achieve the first electrochemical oxidative rearrangement.

Results and discussion

With TH β C **1a** as the model compound, we designed the electrochemical synthesis as follows. The electrolyte contained two common solvents: water for high salt solubility and high conductivity, and acetonitrile (MeCN) for dissolving organic molecules. We added acetic acid (AcOH), which was shown to facilitate the generation of RHS in aqueous media.³⁶ The volume ratio of MeCN, AcOH, and H₂O was optimized to be 15 : 2.4 : 2 (Fig. S1†). The anode was a commercial carbon paper (CP, SGL-39AA) and baked in air at 400 °C for 24 hours to improve the wetting with the electrolyte and further suppress the oxygen evolution reaction (OER).³⁷ The cathode was Pt due to its catalytic activity towards the hydrogen evolution reaction (HER). The electrodes (1 cm² in area) were then assembled in a zero-gap flow cell as shown in Fig. 2(B) with two PEEK endplates and sealed with PTFE gaskets (0.5 mm thick) as shown in Fig. S3 and S4.† The term zero-gap refers to the design wherein the anode, the membrane, and the cathode were pressed using a torque wrench together with no gap among them, so the cell resistance can be minimized, and the flow can be accelerated with a flow field. Through a serpentine flow channel engraved in the endplate, as presented in Fig. S3†, a peristaltic pump circulated the electrolyte at a relatively fast flow rate (>1 mL min^{−1}). The membrane was Nafion® 117, whose effectiveness in preventing the reduction of product **2a** at the counter electrode was obvious as compared with an undivided beaker cell under otherwise the same conditions.

Under the optimal conditions (Table 1), the oxidative rearrangement of **1a** achieved 97% yield of **2a** and 96% faradaic efficiency (FE). Under a fixed applied cell potential, the yield was attained with the full conversion as monitored by thin layer chromatography (TLC). The amount of the product was also converted to charge *via* Faraday's law, the ratio of which over the charge input during the synthesis was the FE (calculated from eqn (1) in the ESI†). As shown in Table 1, the amount of bromide ion was determined to be two equivalents: 1.2 equivalents of LiBr led to significant OER, while 3.0 equivalents resulted in bromination of spirooxindoles (~10%) (entries 2 and 3). The use of a Pt mesh anode in place of carbon paper (entry 6) drastically diminished the yield and FE probably due to Pt-catalyzed side reactions (*i.e.*, C1 oxidation/ring-opening and/or the OER).³⁴ Chloride and iodide (Cl[−] and I[−]) (entries 7 and 8) were also examined but were not effective. This was not unexpected given that the oxidation potential of the chloride ion might be higher than that of nonselective indole oxidation, while the easily generated iodonium ion [I⁺] at the low oxidation potential promoted sluggish oxidative rearrangement.³⁸

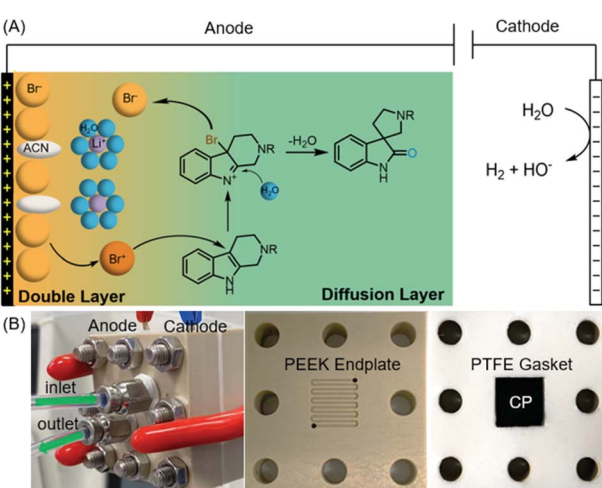


Fig. 2 Reaction mechanism and cell design. (A) The proposed bromide-mediated electrochemical oxidative rearrangement of tetrahydro- β -carboline to spirooxindole; (B) photos of the screw assembled flow cell (top), the PEEK endplate (bottom left), and the cathode (10 mm × 10 mm × 0.6 mm) fitted into the PTFE gasket (bottom right).



Table 1 Selected conditions for bromide-mediated electro-oxidative rearrangement of TH β C **1a**

Entry	Deviation from optimized conditions	Yield ^b (%)	
		FE (%)	
1	None ^a	97	96
2	LiBr: 1.2 eq.	67	67
3	LiBr: 3.0 eq.	88	88
4	Undivided cell (1.7 V) ^c	80	68
5	Undivided cell (1.2 V) ^c	25	11
6	Pt anode	24	9
7	No halide	0	0
8	LiCl instead of LiBr	7	5
9	LiI instead of LiBr	32	7.5
10	E: 0.85 V	55	55
11	E: 1.0 V	71	71
12	E: 1.6 V	50	50

^a The optimal conditions were room temperature with an anolyte of TH β C (0.2 mmol), and LiBr (0.4 mmol, 2.0 eq.). The reaction was performed in a zero-gap (Nafion 117) flow cell as per General Procedure A in the ESI. ^b NMR yield was obtained. ^c The potentials were calibrated vs. RHE from Ag/AgBr in an undivided cell with IR compensation.

Table 1 clearly shows the impact of key variables on the reaction: the cell type (H-cell vs. flow cell), electrode, electrolyte, working voltage, and mediator (LiBr), some of which are further elucidated below. A variable unique to the electrochemical synthesis is the metal cation in the electrolyte. Despite its lack of direct involvement in the reactions, we observed substantial differences in the FE in the order of $\text{Li}^+ > \text{Na}^+ > \text{K}^+ > \text{Bu}_4\text{N}^+$ (as Fig. 3(A) shows). We speculated that the cations acted as spectators, as broadly discussed in the literature on the HER and OER where the catalytic activities were correlated to the hydration energies of the cations, an indicator of the degree of blockage of active sites at the electrode by the hydrated cations.^{39–41} A similar correlation was found between FE and the hydration energy in our case. The cation was less likely to affect the bromide oxidation since the current density increased along with the hydration energy, as shown in Fig. 3(A). Instead, it was more likely to reduce the side reactions (direct electrode oxidation of indoles) through minimizing the diffusion of the substrate into the double layer. In the case of the lithium ion (Li^+), the high hydration energy resulted in water aggregation around the double layer^{39,41} and thus prevented the hydrophobic TH β C from diffusing into the double layer near the electrode (illustrated in Fig. 2(A)).

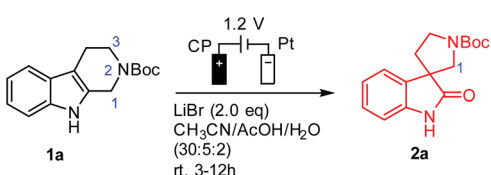
This rationalization was further supported by the current-time profile (Fig. 3(B)). The oxidation current was stable with LiBr at the beginning stage of the reaction but decayed quickly with KBr, which suggested passivation. When examining the carbon paper by scanning electron microscopy (SEM) (the inset

of Fig. 3(B)), we observed porous organic matter derived from TH β C when KBr was used as the mediator. In contrast, we did not detect similar organic matter in the carbon paper electrode for the LiBr-mediated reaction. In a control experiment, we also observed a much larger oxidation peak in LSV with KBr than that with LiBr in the supporting electrolyte (see Fig. S6(A)†). Although the complex CP did not permit full elucidation of the spectator effect, our experimental results reveal the great impact of the spectator cation on selectivity of halide *versus* TH β C oxidation, which facilitates future investigations.

The optimal potential was determined to be 1.2 V across the two electrodes. At a lower potential (0.85 and 1.0 V, Table 1, entries 9 and 10), insufficient RHS $[\text{Br}^+]$ was produced for the rearrangement; at a higher potential (1.6 V, entry 12), the OER and C1 oxidation/ring-opening were observed. This is consistent with linear sweeping voltammetry (LSV, Fig. 3(C)) performed in a three-electrode cell. The potential of LSV was converted to a scale *vs.* the relative hydrogen electrode (RHE) so that the value was comparable with the cell potential of the flow cell given the hydrogen evolution reaction in the positive side of the cell. We characterized three electrolytes, a blank supporting electrolyte of 0.8 M LiClO_4 , the supporting electrolyte with TH β Cs, and the electrolyte with LiBr. Although addition of the supporting salt, necessary for its ionic conductivity in the characterization, potentially complicated the reaction mechanism, we could estimate the onset potentials of the reaction, in good agreement with the optimal potential seen in the flow cell. The oxidation of the bromide ion was about 0.2 V earlier than a small oxidation peak, which was likely associated with the C1 oxidation/ring-opening reaction of TH β C and responsible for the low yield when no mediator was used in the flow cell (entry 7). The OER took place at a higher potential. Therefore, the highest efficiency was achieved in a \sim 0.2 V window.

Another key variable is the rate of electrolyte flow. The zero-gap design permits a high flow rate for rapid oxidation of the bromide ion despite its relatively low concentration, up to \sim 6 mA cm^{-2} at 40 mL min^{-1} , translating to \sim 10% of bromide oxidation per pass of the flow (Fig. 3(D)). A high current density led to an increase of productivity (calculated from eqn (2) in the ESI†). The higher the productivity, the shorter the time, the smaller the cell required, and the more the cost-effectiveness. In addition, the FE was increased slightly to 97% at the highest flow rate, which displayed the other key advantage of rapid flow. As the bromide oxidation was limited by mass transport (Fig. S6(B)†), the higher the flow rate, the more vigorous the convective transport, the higher the interface concentration of RHS $[\text{Br}^+]$, and the more selective the reaction towards the rearrangement. This advantage allowed the flow cell to deliver superior yields over both the undivided cell (Table 1, entry 4 and 5) and flow cells that operate at slow flow rates (<1 mL min^{-1} in Fig. 3(D)).

The electrochemical method was applicable to a broad scope of substrates as summarized in Table 2. Various N_2 protecting groups (NR^3) including $N\text{-CO}_2\text{Me}$, $N\text{-Ac}$, $N\text{-Bn}$ and $N\text{-allyl}$ (**2b–2e**) were tolerated under the electrochemical conditions with good to excellent yields (71–95%). Notably, $N\text{-Bn}$ and $N\text{-allyl}$ of TH β Cs were suitable substrates for this electrochemical oxidation,



Entry	Deviation from optimized conditions	Yield ^b (%)	FE (%)
1	None ^a	97	96
2	LiBr: 1.2 eq.	67	67
3	LiBr: 3.0 eq.	88	88
4	Undivided cell (1.7 V) ^c	80	68
5	Undivided cell (1.2 V) ^c	25	11
6	Pt anode	24	9
7	No halide	0	0
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9	LiI instead of LiBr	32	7.5
10	E: 0.85 V	55	55
11	E: 1.0 V	71	71
12	E: 1.6 V	50	50

^a The optimal conditions were room temperature with an anolyte of TH β C (0.2 mmol), and LiBr (0.4 mmol, 2.0 eq.). The reaction was performed in a zero-gap (Nafion 117) flow cell as per General Procedure A in the ESI. ^b NMR yield was obtained. ^c The potentials were calibrated vs. RHE from Ag/AgBr in an undivided cell with IR compensation.



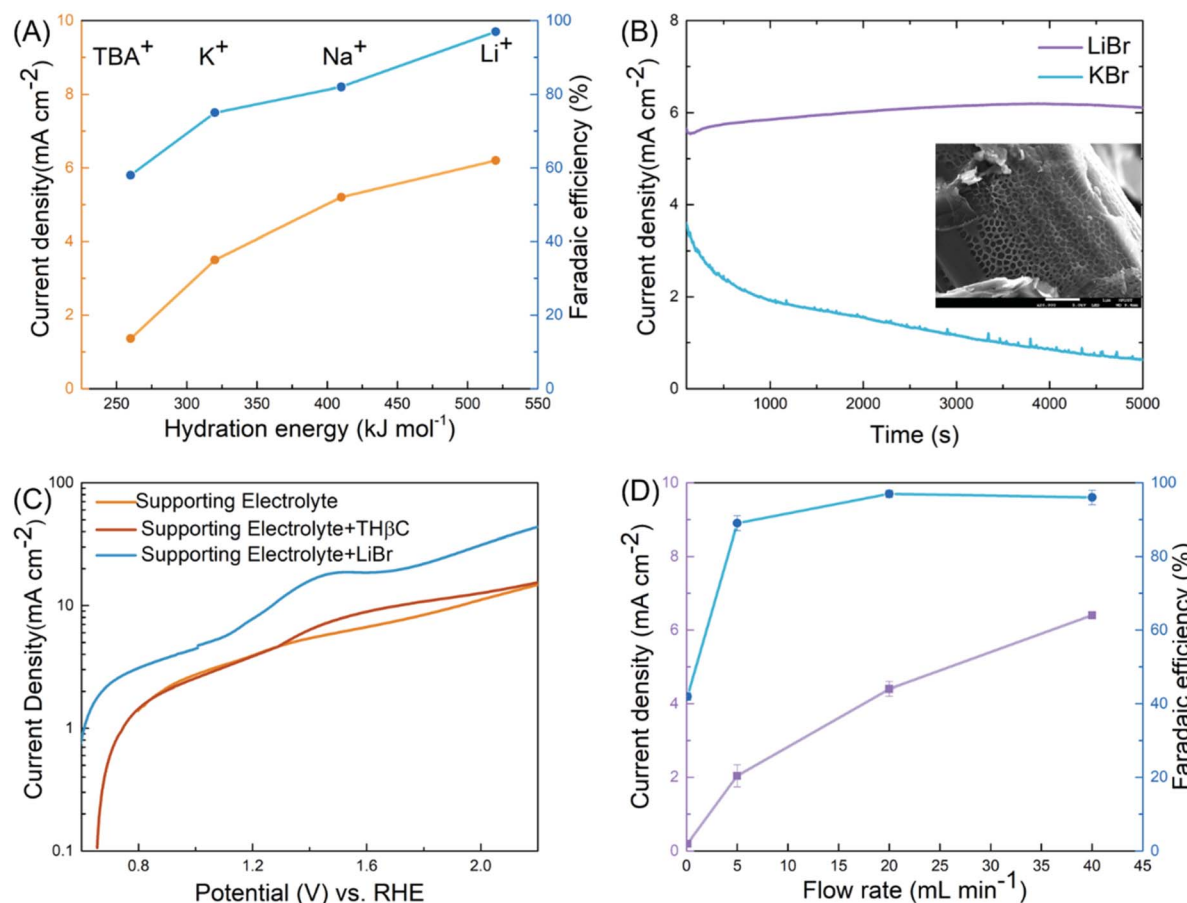


Fig. 3 Key variables of the electro-oxidative rearrangement of TH β C. (A) The effect of cations on the current density (orange) and the FE (blue). (B) Current–time profiles with LiBr (purple) and KBr (blue) as the salts. The inset is the SEM image of the carbon fiber in the carbon paper used in the synthesis reaction with KBr, on which a porous organic layer is likely the polymer product of the side reaction. (C) Linear sweep voltammetry (LSV) in a three-electrode cell (Fig. S6†) at 50 mV s⁻¹ of a supporting electrolyte (0.8 M LiClO₄, orange), the supporting electrolyte with TH β C (red), and the supporting electrolyte with LiBr (blue). (D) The impact of flow rates on the current density (purple) and the FE (blue).

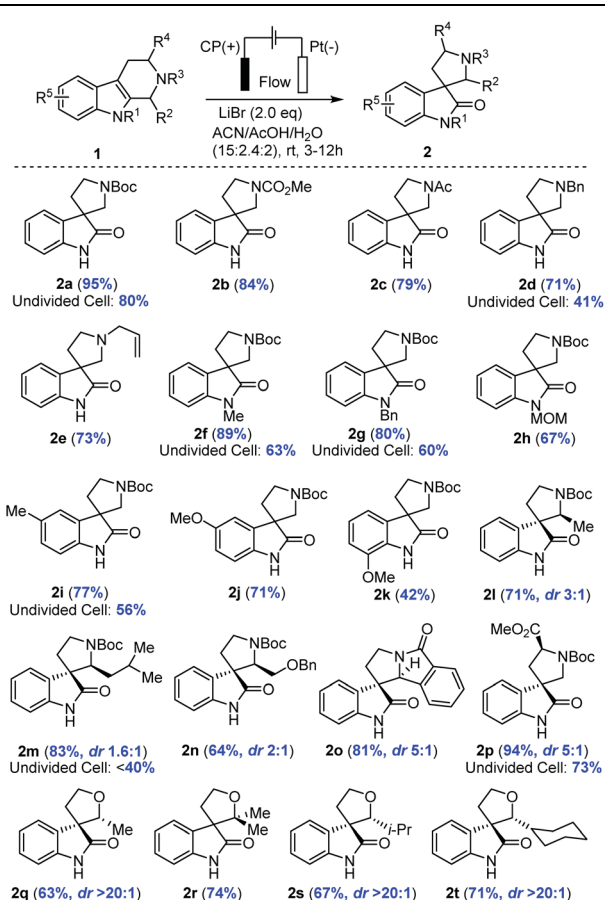
which was interesting in light of the well-established Shono oxidation^{42,43} that was the major pathway for tertiary amines or amides under conventional electrochemical oxidation conditions. Our studies (Table 2 and Fig. 3(C)) suggested that Li⁺ might impede the direct Shono oxidation of TH β Cs at the anode. This finding was important for mediator-promoted electrochemical oxidation/reduction in organic synthesis and for application of our electrochemical oxidative rearrangement of TH β Cs in synthesis of bioactive spirooxindole molecules containing an *N*-alkyl group, particularly tertiary amines.

Next, we examined the protecting group on indole nitrogen (NR¹) and found that only electron-donating groups such *N*-Me (**2f**), *N*-Bn (**2g**), and *N*-MOM (**2h**) were suitable for the oxidative rearrangement with good yields (67–89%), not a surprise as an electron-withdrawing group (*N*-Ac, *N*-Ts, or *N*-Boc) would increase the oxidation potential to overlap with the direct indole oxidation and water oxidation. Electron-rich TH β Cs with a methyl or methoxy group on the benzene ring were good substrates for the electrochemical oxidative rearrangement to provide spirooxindoles with good yields (**2i–2k**: 42–77%). Different C1 substituents (*R*²) of TH β Cs did not reduce the

yields (64–83%), but a diastereomeric mixture (**2l–2o**) was often obtained with a ratio ranging from 1.6 : 1–5 : 1. Fortunately, these diastereomers could be separated by flash column chromatography on silica gel and assigned reliably by comparison of the NMR spectra with reported ones.¹⁹ When tryptophan-derived TH β C was used, the spirooxindole (**2p**) was isolated in excellent yield (94%) under the optimal conditions, although diastereoselectivity was only moderate (dr. 5 : 1). Lastly, we further extended our electrochemical oxidative rearrangement to tetrahydropyrano[3,4-*b*]indoles (THPIs).¹⁹ All THPIs with monosubstituted or disubstituted alkyl groups at the C2 position underwent the expected oxidative rearrangement to provide **2q–2t** in good yields. Interestingly, the diastereoselectivity (dr. >20 : 1) was excellent as compared to TH β C substrates.

The advantage of the zero-gap flow cell over an undivided beaker cell (**2a**, **2d**, **2f**, **2g**, **2i**, **2m**, and **2p**, Table 2) was significant in the extended substrate scope. In the undivided cell, we carried out the synthesis similarly under a constant potential but with a reference electrode (AgBr/Ag, calibrated to be stable at –0.26 V vs. RHE) to avoid inconsistency caused by the large

Table 2 Substrate scope of electrochemical oxidative rearrangement of tetrahydro- β -carbolines to spirooxindoles. The yields and the diastereoselectivity (dr, if present) are shown in blue^a

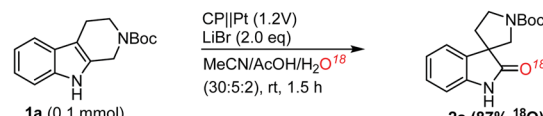


^a The reaction was carried out at room temperature as per General Procedure A using a zero-gap flow cell. Isolated yield was obtained.

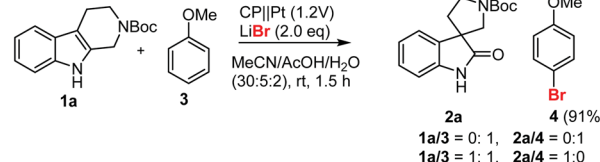
cell resistance (\sim 120 ohms). To attain a current at the same order of magnitude as that of the flow cell, the potential approached 1.7 V vs. RHE. Under these conditions, the undivided cell could merely deliver 40 to 60% of yield, inferior to the generally high yields ($>70\%$) in the flow cell. In addition to the use of the ion-selective membrane in the flow cell to separate the product from the cathode as discussed earlier, another key factor was mass transport. Take the transformation of 1m to 2m as an example (details in ESI, Section 6†). The main byproduct in the undivided cell was from arene bromination, likely because this large molecule cannot diffuse through the porous electrode. Concentration polarization thus led to a large overpotential and a low selectivity ($<40\%$ in the undivided cell). The polarization was minimized by the rapid electrolyte flow, resolving the trade-off between a high productivity and a high selectivity encountered in many previous studies on similar transformations as we summarize in Fig. S7.†

To shed light on the mechanism of this electrochemical oxidative rearrangement, we performed some mechanistic studies as shown in Scheme 1. First, an oxygen labelling

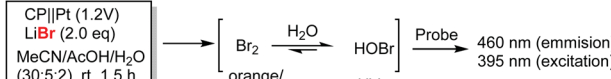
(a) H₂O¹⁸ labeling experiment



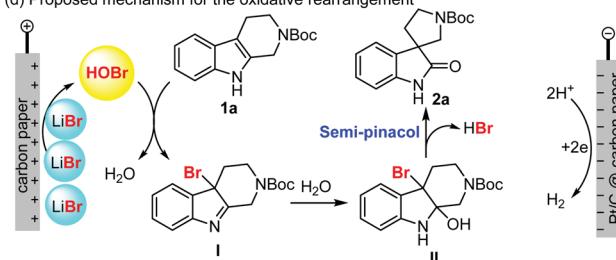
(b) Trapping the reactive brominating species



(c) Detection of reactive brominating species



(d) Proposed mechanism for the oxidative rearrangement



Scheme 1 Mechanistic studies and the proposed mechanism for electrochemical oxidative rearrangement of tetrahydro- β -carbolines. (A) Isotopic labelling experiment; (B) RBS trapping experiments; (C) detection of RBS; the (D) proposed mechanism for electrochemical oxidative rearrangement of TH β C.

experiment was performed to identify the source of oxygen on spirooxindoles (Scheme 1a). When H₂O was replaced with H₂¹⁸O as the reaction medium, more than 87% ¹⁸O (determination by mass spectrum analysis) was incorporated in the spirooxindoles (1a), which suggested that water was involved in the reaction and was the main oxygen source of spirooxindole. Secondly, we attempted to capture the reactive brominating species (RBS, Scheme 1b), which was believed to be responsible for the oxidative rearrangement. Anisole (3) was easily brominated in excellent yield (91%) in the absence of TH β C. When a 1 : 1 ratio of anisole and TH β C was used, spirooxindole was isolated as the sole product without detection of bromoanisole. These results suggest that RBS was generated in the reaction, and interestingly, oxidative rearrangement occurred faster than bromination. This also explained why bromination of spirooxindoles was rarely observed. Thirdly, we attempted to elucidate the possible molecular structure of RBS using UV absorption and fluorescence spectra of the HOBr probe⁴⁴ (Scheme 1c, also see Fig. S8 in the ESI†), which proved the generation of HOBr as the RBS for the rearrangement and bromination reactions. The obvious color change of the reaction mixture (brown-orange-clear) suggested the presence of a small amount of bromine, which could be converted into HOBr under acidic aqueous reaction conditions. Based on the above results and the well-established bromide-mediated electrooxidation, we proposed



a plausible mechanism of electrochemical oxidative rearrangement of indole (Scheme 1d). Electrooxidation of LiBr generated HOBr, which oxidized the TH β C into bromo indoline intermediate I. Water addition followed by semipinacol rearrangement furnished spirooxindole.

Conclusion

We have designed a zero-gap flow cell and demonstrated efficient, high-rate LiBr-mediated electro-oxidative rearrangement of tetrahydro- β -carbolines to spirooxindoles using commercial electrodes. The generality of this electrochemical oxidative rearrangement was demonstrated with 20 examples with good to excellent yields. Importantly, we discovered that the lithium ion played a key role as a spectator in the double layer to suppress the direct oxidation of the substrate and improve the yield. Compared to recent literature reports, our designed zero-gap flow cell delivers the highest FE and productivity. We expect that our catalytic system and cell design could be further applied to other electro-organic reactions.

Data availability

Experimental procedures and characterization data are available within this article and its ESI.† Data are also available from the corresponding author on request.

Author contributions

Y. Zheng and Y. T. Cheung contributed equally to this work.

Conflicts of interest

The authors declare no competing interests.

Acknowledgements

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