Mechanism of electrochemical capture of CO₂ via redox cycle of chlorinated 1,4-naphthoquinone in BMIMBF₄: An in-situ FT-IR spectroelectrochemical approach

Xuejiao Qiao, Dan Li, Longjiu Cheng, Baokang Jin

Department of Chemistry, Anhui University, Hefei 230601, People’s Republic of China

ABSTRACT

The mechanisms of electrochemical capture of CO₂ via redox cycle of 1,4-naphthoquinone (1,4-NQ), 2,3-dichloro-1,4-naphthoquinone (DCNQ) and 2-chloro-1,4-naphthoquinone (CNQ) in 1-butyl-3-methylimidazolium tetrafluoroborate (BMIMBF₄) have been investigated by cyclic voltammetry (CV) and in-situ FT-IR spectroelectrochemistry techniques. In the absence of CO₂, 1,4-NQ and DCNQ undergo a reversible two-step one-electron process while CNQ not. In addition, their two successive electron transfer processes in ionic liquid (IL) are different from that of molecular solvent. There is only a small difference in peak potential between two-step reduction, which possibly due to the ion-pair effects. Meanwhile, the reduction potentials of DCNQ and CNQ shift positively due to the electron-withdrawing effects of chlorine groups. The radical anion (Q⁻) and dianion (Q₂⁻) formed during the electrochemical reduction can be used as nucleophiles to attack the electrophilic carbon center of CO₂ and form a stable CO₂ adduct. For 1,4-NQ, 2 equiv. of CO₂ can be captured during the electrochemical reduction of 1,4-NQ. Whereas, 1 equiv. of CO₂ is captured by 1 equiv. of dianion during the electrochemical reduction of DCNQ and CNQ. Furthermore, CO₂ adducts should be the carbonate type products according to the theoretical calculation results by Gaussian 09 package.

1. Introduction

The excessive CO₂ emission caused by fossil fuel consumption has driven tons of attention to CO₂ separation and reduction [1–5]. Therefore, strategies to decrease the CO₂ emission, such as CO₂ capture, storage, and reduction, have attracted worldwide attention [6–10]. Nucleophiles and alkalines, such as amine, KOH, ammonia, and potassium, are normally used in post-combustion CO₂ capture [11–14]. However, these reagents are limited in practical applications due to their corrosion, energy-intensive to regenerate, easy oxidation, and low CO₂ loading capacity [14]. Therefore, numerous separation approaches, including physical absorption, membranes, ionic liquids, and electrochemical reactions have been employed in CO₂ capture research area [15–18]. Among these new methods, electrochemical reactions, using electrons as strong redox reagents for transformation of target compounds in aqueous or organic solutions, are recognized as a promising way to capture CO₂. Electrochemical methods are more energy-efficient, which can be conducted under relatively mild conditions (20–60 °C) [19].

Quinones and their derivatives, such as 1,4-naphthoquinone (1,4-NQ), 5-hydroxy-1,4-naphthoquinone (HNQ), and p-benzoquinone (BQ), have been widely studied due to their vital roles in electrochemical redox systems [20–23]. Normally, when hydrogen atoms on quinone ring are replaced by electron-withdrawing groups, their electrochemical behavior will also be changed [24]. The radical anion (Q⁻) and dianion (Q²⁻) formed during the electrochemical reduction are good nucleophiles for CO₂ capture, which has attracted lots of attention, recently [19,24,25]. However, these electrochemical reductions of quinones and their derivatives are usually conducted in buffer solution, aqueous solution and acetonitrile solvent [22,23]. As we all know, how to fix CO₂ in large scale with minimum energy has always been the goal of researchers. Therefore, traditional molecular organic solvents are not favorable option due to their evaporation losses. Ionic liquids, types of molten salts composed of organic cations and organic/inorganic anions, are desirable electrolytes for CO₂ capture due to their negligible vapor pressure, high thermal and electrochemical stability, nonflammability, and high solubility of CO₂ [26–28]. Also, cyclic voltammetric experiments can be readily performed in ionic liquids without supporting electrolyte [29]. Therefore, the application of ionic liquids as electrolytes avoids the use of harsh solvents, catalysts, and...
supporting electrolytes. In addition, BMIMBF₄ (1-butyl-3-methylimidazolium tetrafluoroborate) or BMIMPF₆ (1-butyl-3-methylimidazolium hexafluorophosphate) type IL is as the most effective and promising of substitute for classical solvents [26,30].

Recently, efforts have been made to electrochemically capture CO₂ in ionic liquids. Hatton has studied the electrochemical capture and release of CO₂ in 1-ethyl-3-methylimidazolium tricyanomethanide ([Emim][TCM]) [19]. The CO₂ electrochemical capture in superbasic [P66614][124Triz] has been studied by Hardacre [31]. The role of anions of 1-butyl-3-methylimidazolium ionic liquids during CO₂ electrochemical capture in N-heterocyclic carbene has also been investigated in detail [32]. The electrochemical capture and reduction of CO₂ in ionic liquids gels have been demonstrated by Wallace [33]. Though the electrochemical capture of CO₂ in ionic liquids have been investigated in detail in the past few years, few works related to electrochemical capture of CO₂ in ILs via redox cycle of quinone have ever been presented. Also, the mechanism of CO₂ electrochemical capture and release via redox cycle of quinone and its derivatives in ionic liquids is not clear. Therefore, the stoichiometry of CO₂ electrochemical capture deserves to be further investigated.

It has been demonstrated the advantages of cyclic voltabsorptometry (CVA) and derivative cyclic voltabsorptometry (DCVA) in electrochemical research [34,35]. Therefore, the mechanism of electrochemical capture of CO₂ via redox cycle of 1,4-NQ, DCNQ and CNQ in BMIMBF₄ were explored by CV and IR spectroelectrochemistry in this work. The results indicate that electroreduction product can be well used as a nucleophile to capture CO₂. And, the electrochemical capture of CO₂ via redox cycle of chlorinated 1,4-naphthoquinones exhibit different capture mechanisms. In addition, the stoichiometry of CO₂ electrochemical capture was evaluated. Furthermore, combined theoretical computation technique, the possible structure of CO₂ adduct was analyzed.

The stoichiometry of CO₂ electrochemical capture deserves to be further investigated.

### 2. Experimental section

#### 2.1. Chemicals

1,4-Naphthoquinone (Sigma–Aldrich, > 98.0%), 2,3-dichloro-1,4-naphthoquinone (Sigma–Aldrich, 98%) and 2-chloro-1,4-naphthoquinone (Sigma–Aldrich, 98%) were used to capture CO₂ during their redox reactions in 1-butyl-3-methylimidazolium tetrafluoroborate (BMIMBF₄) (Linzhou Keneng Materials Technology Co. Ltd., 99.99%). High purity carbon dioxide (Nanjing special gas Co. Ltd., 99.99%) is used for electrochemical capture reactions directly. The solution was degassed with high purity nitrogen (Nanjing special gas Co. Ltd., 99.99%) for 10 min before experiment.

#### 2.2. Voltammetry

A three-electrode system with an electrochemical analyzer (Shanghai Chenhua Instrument Co. Ltd., CHI630E potentiostat) was used for electrochemical experiments. A 4 mm diameter gold disk was used as a working electrode. The working electrode was polished with 0.05 µm alumina slurry on a polishing cloth and cleaned with double distilled water and ethanol respectively before experiments. The Ag/AgCl and Pt wire were used as reference electrode and counter electrode, separately.

### 2.3. In-situ FT-IR spectroelectrochemistry

In-situ FT-IR spectroelectrochemistry experiments were performed in a homemade thin-layer spectroelectrochemical cell [36]. Spectroscopic apparatus (Nicolet Company of the United States, Nicolet iS50) equipped with a specular reflectance accessory and a HgCdTe/A detector cooled with liquid-nitrogen. Each spectrum was composed of 32 interferograms with 0.9 s as sampling intervals. And the spectral resolution was 16 cm⁻¹. The experimental results were processed with Grams/3D software.

#### 2.4. Theoretical computation

The structures were relaxed at the B3LYP [37]/6-311 + + G** level of theory. The energies and NBO (Natural Bond Orbital) charges of the anion radical, dianion were also given at this level of theory. All theoretical calculations were performed by the Gaussian 09 package [38].

### 3. Results and discussion

#### 3.1. Q in BMIMBF₄ without CO₂

The CVs of 1,4-NQ, DCNQ, CNQ in BMIMBF₄ are shown in Fig. 1 (recorded at a gold disk electrode, room temperature). There are two unwell-defined couples of anodic and cathodic peaks observed in their electrochemical redox processes. Also, the reduction potential of quinone shifts positively with the increased number of chlorine groups, which is due to the electron-withdrawing effects of chlorine. The two unwell-defined couples of CV peaks are quite different with two well-defined CV waves observed in dimethyl sulfoxide (DMSO), which might be attributed to the ion-pairing effects, especially the interaction between di-anion (Q²⁻) and IL [19,21].

In-situ FT-IR electrochemical experiments were performed in IL. The three-dimensional (3D) spectra are in-situ recorded in the wavenumber range from 1100 cm⁻¹ to 1800 cm⁻¹ to track the concentration variation of reactant, intermediate and product during the electrochemical reaction (Fig. 2). The downward bands at 1671, 1684 and 1680 cm⁻¹ show the concentration of 1,4-NQ, DCNQ and CNQ decrease in the reduction process while recover in the oxidation process. The upward bands at 1370, 1356, and 1408 cm⁻¹, etc. indicate that the concentrations of 1,4-NQ²⁻, DCNQ²⁻ and CNQ²⁻ increase during the reduction process while decrease during the oxidation process. Specially, the upward bands at 1515, 1510 and 1511 cm⁻¹ (assigned to 1,4-NQ⁻, DCNQ⁻ and CNQ⁻ respectively) appear and disappear periodically in both the reduction and oxidation processes [21,22]. During the reduction process, Q is first reduced to Q⁻ and as the reduction goes on, the intermediate Q⁻ is further oxidized to Q²⁻. Also during the oxidation process, Q²⁻ is oxidized to intermediate Q⁻ and subsequently Q⁻ is further oxidized to Q. Therefore, the concentration trend of Q⁻ in the reduction and oxidation process is similar. The assignment of IR spectra
3.2. Electrochemical capture of CO2 via redox cycle of quinone derivatives in BMIMBF4

3.2.1. Electrochemical capture of CO2 via redox of 1,4-NQ in BMIMBF4

The mechanism of CO2 capture via electrochemical redox of 1,4-NQ in BMIMBF4 has been investigated by the CV experiments (Fig. 5). As shown in Fig. 5, there is no redox peaks in saturated CO2 solution without 1,4-NQ when the scan range is 0.7 to −1.6 V (recorded at a gold disk electrode, room temperature). With the addition of CO2, two reductive waves are merged into one peak. The first reduction peak increases correspondingly, and the current of the single reductive wave C1 (voltammogram a) is much larger (almost doubled) than that of reductive wave C3 (voltammogram c). The results indicate that the complexation of CO2 with the reduced quinone caused the second reduction to shift positively, this positive shift is large enough for both reductions to overlap at the reductive wave C1. Notably, in voltammogram b, although there is only one reduction peak, two oxidation peaks (A1 and A2) are observed. Indeed, with 30% CO2 in the system, the free reduced 1,4-NQ will exist because there was insufficient CO2 dissolved in the IL available to react with all of the reduced 1,4-NQ. In the oxidation process, the free reduced 1,4-NQ (Q−) will be oxidized at A1 position, so there are two oxidation peaks (A1 and A2) in voltammogram b.

To further explore the mechanism of 1,4-NQ with saturated CO2 in IL, in-situ FT-IR spectrroelectrochemistry experiments of 1,4-NQ with saturated CO2 in IL were performed in a homemade thin-layer [36]. The CV of 1,4-NQ with saturated CO2 in scan range of 0 to −1.3–0.7 V is shown in Fig. 6A (recorded at a gold disk electrode, room temperature). The corresponding 3D IR spectra presented in Fig. 6B is significantly different from the 3D IR spectra of 1,4-NQ without CO2 (Fig. 2A). There are only two types of IR absorption bands in Fig. 6B, while the characteristic absorption peak of intermediate-type (Q•−) disappears. However, there is a downward band at 2342 cm−1 appears, which can be used to track the concentration changes of CO2. Another new IR absorption peak at 1634 cm−1 may be attributed to the product CO2 adduct.

To further explore the reaction behavior, the IR spectra are processed to CVA curves (Fig. 6C). The absorption bands at 1671, 2342 and 1634 cm−1 are used to track the concentration changes of CO2. Moreover, the IR absorption bands basically return to the initial value at the end of the sweep potential (time). Consecutive scan results show that the electrochemical capture of CO2 is a reversible process (see Fig. S1, Supporting information for the details).

The DCVA curves are shown in Fig. 6D, the peak potential of 1671, 2342 and 1634 cm−1 is basically the same, which further indicates the rapid chemical reaction between 1,4-NQ•− or [1,4-NQ-CO2]2− and CO2.

The results of CV and IR spectrroelectrochemistry indicates that 1,4-NQ and DCNQ undergo a reversible two-step one-electron process (reaction (1) in Scheme 2). While CNQ undergoes a non-reversible two-step one-electron process (reaction (2) in Scheme 2).

3.2. Electrochemical capture of CO2 via redox cycle of quinone derivatives in BMIMBF4

obtained in BMIMBF4 of 1,4-NQ, DCNQ and CNQ are given in Table 1, respectively.

To further explore the electrochemical behavior, the IR spectra are processed to CVA curves (Fig. 3). The absorption bands of Q, Q− and Q2− increase and decrease periodically with the sweep potential (time). The IR absorption peaks of 1,4-NQ (Fig. 3A) and DCNQ (Fig. 3B) basically return to the initial values at the end of the sweep potential (time). However, IR bands of CNQ does not return to the original state, suggesting that the second step reduction of CNQ in IL is non-reversible, which can also be observed from its CVA curves (Fig. 3C). This non-reversibility was believed to be due to the much stronger interaction of IL with the reduced CNQ than that with the reduced 1,4-NQ and DCNQ as a result of the dipole induced by the ortho-positioned single chlorine group of the reduced CNQ.

DCVA curve is similar to CV wave in ideal state [39]. The DCVA curve greatly suppresses the influence of charging current, which shows two well-defined redox couples of peaks (Fig. 4). The results of DCVA can provide more clear DCVA waveforms when compared with the CV waves in Fig. 1. In addition, the DCVAs at 1515, 1510 and 1511 cm−1 correspond to Q•− which have two redox couples of peaks.

The results of CV and IR spectroelectrochemistry indicate that 1,4-NQ and DCNQ undergo a reversible two-step one-electron process (reaction (1) in Scheme 2). While CNQ undergoes a non-reversible two-step one-electron process (reaction (2) in Scheme 2).

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**Fig. 1.** Cyclic voltammograms in BMIMBF 4 containing 20 mmol/L 1,4-NQ (a), DCNQ (b), CNQ (c) and only BMIMBF 4 (d), respectively. The potential scan rate was 10 mV/s.

**Fig. 2.** The corresponding rapid-scan IR spectra in BMIMBF 4 of 1,4-NQ (A), DCNQ (B) and CNQ (C), respectively.
an additional molecule of CO2 to yield [1,4-NQ-2CO2]2− (reactions (5) and (6) in Scheme 3). Therefore, the mechanism of electrochemical capture of CO2 via redox cycle of 1,4-NQ follows the electrochemistry-electrochemistry-chemistry (EECr) mode. It should be noted that two molecules of CO2 can be captured during the electrochemical reduction of 1,4-NQ, which will be further discussed in Section 3.3.

### 3.2.2. Electrochemical capture of CO2 via redox of DCNQ in BMIMBF4

The CV results of DCNQ with saturated CO2, unsaturated CO2 and without CO2 in IL are shown in Fig. 7 (recorded at a gold disk electrode, room temperature). Two reductive waves can still be observed when saturated CO2 is introduced (voltammogram a in Fig. 7). It means that DCNQ with CO2 still requires two reductive waves to form radical anion and dianion, which is obviously different from that of 1,4-NQ with saturated CO2 (Fig. 5).

However, the CV results obviously show that the second reductive potential of DCNQ with CO2 becomes more positive and the corresponding oxidative peak is insignificant. In Fig. 7, the second reductive wave shifts positively with the increase of CO2. However, two reductive waves can still be seen at 100% CO2 concentration. Moreover, the introduction of CO2 has no effect on the peak potential of the first reduction wave, indicating that there is no interaction between DCNQ− and CO2.

The IR spectroelectrochemistry experiments of DCNQ with CO2 were also carried out in order to understand its mechanism. The CV of DCNQ with saturated CO2 in the scan range of 0.1 to −1.1−0.5 V and the corresponding 3D IR spectra are shown in Fig. 8A and B. Three types of IR bands can still be observed in 3D IR spectra and the upward absorption band at 1510 cm −1 (assigned to DCNQ−) still exists.

In Fig. 8C, the bands at 1684, 2342, 1510, and 1640 cm −1 are employed to track DCNQ, CO2, DCNQ−, and [DCNQ-CO2]2− respectively. In the reduction process, the concentration of DCNQ (1684 cm −1) begins to decrease at about 21 s (−0.11 V). At the same time, the concentration of DCNQ− (1510 cm −1) begins to increase at 21 s (−0.11 V), and a maximum concentration is obtained at 72 s (−0.62 V). Subsequently, the DCNQ− is further reduced to DCNQ2−. DCNQ2− reacts with CO2 quickly, making the concentration of CO2 (2342 cm−1) decrease at 60 s (−0.50 V) and reach minimum at 119 s (−1.09 V). Meanwhile, the concentration of [DCNQ-CO2]2− (1640 cm−1) begins to increase at 60 s (−0.50 V) and reach maximum at 119 s (−1.09 V). The CVA shows that DCNQ2− reacts with CO2 (reaction 9 in Scheme 4). And the disappearance of CO2 (2342 cm−1) obviously lags behind DCNQ (1684 cm−1), which further proves that CO2 do not participate in the first reduction step of DCNQ. Moreover, the IR absorption bands basically return to the initial value at the end of the sweep potential (time). Consecutive scan results also prove that electrochemical capture of CO2 via redox of DCNQ is also a chemical reversible process (see Fig. S2, Supporting information for the details).

From Fig. 8D, the shape of DCVA at 1510 cm−1 corresponds to DCNQ− is similar to that of CV in Fig. 8A. DCVA curves of CO2 (2342 cm−1) and product (1640 cm−1) correspond to the chemical step, which obviously occurs in the second electrochemical step.

Based on the CV and IR spectral results, DCNQ with CO2 follows the electrochemistry-electrochemistry-chemistry (EECr) mode. The whole reaction process is described in Scheme 4.

### 3.2.3. Electrochemical capture of CO2 via redox of CNQ in BMIMBF4

The CV wave of CNQ with saturated CO2 is shown in Fig. 9A (recorded at a gold disk electrode, room temperature). Compared with Fig. 1c, only one pair of reedox waves is observed when saturated CO2 is introduced. This phenomenon is similar to 1,4-NQ with saturated CO2. However, the IR absorption band at 1511 cm−1 corresponding to the intermediate CNQ− still exists (Fig. 9B).

The bands at 1680, 2342, 1511 and 1636 cm−1 are selected to track CNQ, CO2, CNQ− and [CNQ-CO2]2−, respectively (Fig. 9C). In the reduction process, the concentration of CNQ (1680 cm−1) begins to decrease at about 22 s (−0.12 V). Meanwhile the concentration of CNQ

| Table 1 | The assignment of IR absorption bands of 1,4-NQ, DCNQ and CNQ with or without CO2 [21–23]. |
|----------------|------------------|------------------|
|Assignment Wavenumbers (cm−1) | Assignment Wavenumbers (cm−1) | Assignment Wavenumbers (cm−1) |
| C-O from [1,4-NQ-2CO2]2− | 1320 | C-O from [DCNQ-CO2]2− |
| C=O from [1,4-NQ-2CO2]2− | 1387 | C-O from DCNQ2−, [DCNQ-CO2]2− |
| C–O from 1,4-NQ2− | 1356 | C-O from CNQ2− |
| C=C from 1,4-NQ | 1439 | C-O from CNQ |
| C=O from 1,4-NQ | 1684 | C-O from DCNQ |
| C=O from CO2 | 1640 | C-O from DCNQ− |
| C–O from DCNQ2−, [DCNQ-CO2]2− | 1356 | C-O from CNQ− |
| C–O from CNQ2−, [CNQ-CO2]2−, | 1356 | C-O from CNQ− |
| C–O from CNQ2− | 1317 | C-O from CO2 |
| C=C from CNQ | 1318 | C-O from DCNQ |
| C=O from CNQ | 1640 | C-O from DCNQ− |
| C-O from CNQ2− | 1408 | C-O from CNQ− |
| C–O from CNQ2− | 1356 | C-O from CNQ− |
| C=C from CNQ | 1356 | C-O from CNQ− |
| C=O from CNQ | 1680 | C-O from CNQ |
| C-O from CNQ2− | 1511 | C-O from CNQ− |
| C–O from CNQ2− | 1408 | C-O from CNQ− |
| C=C from CNQ | 1318 | C-O from CNQ |
| C=O from CNQ | 1680 | C-O from CNQ |
| C-O from CNQ2− | 1511 | C-O from CNQ− |
| C–O from CNQ2− | 1408 | C-O from CNQ− |
| C=C from CNQ | 1356 | C-O from CNQ |
| C=O from CNQ | 1680 | C-O from CNQ |
| C-O from CNQ2− | 1511 | C-O from CNQ− |
| C–O from CNQ2− | 1356 | C-O from CNQ |

Fig. 3. The corresponding CVAs for 1,4-NQ (A) at 1671, 1515 and 1370 cm−1, for DCNQ (B) at 1684, 1510 and 1356 cm−1, for CNQ (C) at 1680, 1511 and 1356 cm−1, respectively.
− (1511 cm−1) starts to increase at 22 s (−0.12 V), and a maximum value is obtained at 77 s (−0.67 V). Subsequently, CNQ− is further reduced to CNQ2−. Then CNQ2− reacts with CO2 rapidly, making the concentration of CO2 (2342 cm−1) decrease at 39 s (−0.29 V) and reach its minimum at 120 s (−1.10 V). At the same time, the concentration of [CNQ-CO2]2− (1636 cm−1) begins to increase at 39 s (−0.29 V) and reach its maximum at 120 s (−1.10 V). The results indicate that CO2 is captured by CNQ2− rather than CNQ−. In addition, when compared to the CVAs in Fig. 3C, each absorption band in Fig. 9C returns to its original position to a large extent. The reason is that CO2 captured by CNQ2− is a reversible process (reaction (12) in Scheme 5). So the reversibility of the whole reaction is improved.

It can be clearly seen from Fig. 9D that the DCVA at 1511 cm−1 corresponding to CNQ− has two redox couples of peaks. Although the CV wave in Fig. 9A has only one pair of redox waves, the DCVA curve at 1511 cm−1 corresponds to a two-step one electron transfer process. The results further prove that the dianion CNQ2− reacts with CO2, rather than the radical anion CNQ− and CO2. In addition, DCVA curves of the absorbance at 2342 cm−1 and 1636 cm−1 correspond to the chemical step.

The results of CV and IR spectroelectrochemistry show that although CNQ undergoes a non-reversible two-step one electron transfer process in IL, CNQ2− with CO2 is a reversible chemical reaction. So, its mechanism can be described as electrochemistry-electrochemistry-chemistry (EECr) (Scheme 5).

Furthermore, the experiments with different scan rate from 1 to 10 mV/s are also conducted to depict its reaction mechanism (Fig. 10A). With the decrease of scan rate, the second reductive wave begins to appear step wisely. In addition, the CV experiments with different concentration of CO2 were also performed at a scan rate of 2 mV/s to confirm the reactions between CNQ2− and CO2 (Fig. 10B). It is clearly shown that the second reduction potential gradually shifts positively with the increase of CO2 concentration. The results further implied the EECr reaction between CNQ and CO2 (Scheme 5).

3.3. An evaluation of the quantitative relationship on electrochemical capture of CO2

A series of experiments with different CO2 mole percent are carried out to explore the stoichiometry of CO2 capture during the redox reaction of Q in IL. The CVs of 20 mmol/L Q in IL with different concentrations of CO2 are shown in Fig. 11 (recorded at a gold disk electrode, room temperature). The saturated molarity of CO2 in BMIMBF4 is defined as 100% mole fraction of CO2 (≈99 mmol/L) [40,41]; the other percentages are relative concentration of CO2 with saturated concentration as a reference.

In Fig. 11A, voltammogram j is the CV curve with only CO2 dissolved in IL, from which it is obvious that the reduction wave appears at about −2.0 V. In fact, with the negative potential scanning, CO2 may undergo two reactions, that is, CO2 itself can be further reduced or may combine with the reduced IL [32,42]. Therefore, scanning to the more negative potential, the excess CO2 is further consumed, whether it is the reduction of CO2 itself or the capture of CO2 by the reduced IL. Using this method, the total amount of CO2 involved in the reaction can be observed, thus further obtaining the stoichiometry of CO2 capture as a reference.

As can be seen from Fig. 11 that the further reaction of CO2 does not occur at about −2.0 V when the concentration of CO2 is low (< 40% in Fig. 11A, < 20% in Fig. 11B, < 20% in Fig. 11C), which indicates that all CO2 will be completely captured by the reduced Q at lower concentration. However, the further reaction of CO2 occurs at about −2.0 V when the CO2 concentration is higher than 30% (> 55% in Fig. 11A, > 30% in Fig. 11B, > 30% in Fig. 11C).

Fig. 12 is the CVAs diagram corresponding to the CV process of Fig. 11. The absorption peak at 2342 cm−1 changes obviously when different concentrations of CO2 are introduced. As for 1,4-NQ, DCNQ and CNQ, at low CO2 concentration, the absorption value of 2342 cm−1 reaches the minimum at about −1.2 V and remains basically unchanged in the reduction process. The results imply that CO2 is completely captured by the reduced Q. But, the further reaction of CO2...
at the more negative potential will be observed when excessive CO₂ is introduced (> 55% in Fig. 12A, > 30% in Fig. 12B, > 30% in Fig. 12C). So, the CVA curves will drop again and reach the minimum value at about −2.4 V (Fig. 12). The CVA curves clearly show the periodic variation of 2342 cm⁻¹ in the redox process.

Significantly, the absorbance value of 2342 cm⁻¹ at −2.4 V can reflect the total amount of CO₂ involved in the reaction process. From Fig. 13, at −2.4 V, the mole percent of CO₂ (represented by x) and the corresponding absorbance value at 2342 cm⁻¹ are in a good linear relationship, respectively. The simulation results further prove that CO₂ participates in the reaction process.

Fig. 14 shows the relationship between CO₂ mole percent and its corresponding absorbance value of IR band at 2342 cm⁻¹ at −1.2 V. According to Fig. 14, the corresponding absorbance value decreases dramatically with increase of CO₂ concentrations. However, it decreases slowly at higher CO₂ concentration. At room temperature, the solubility of saturated CO₂ in BMIMBF₄ is 99 mmol/L. For 1,4-NQ (Fig. 14A), the intersection of the two lines is at about 40.43% (about 40 mmol/L), which means that 40 mmol/L CO₂ is captured. Therefore, 2 equiv. of CO₂ can be captured during the electrochemical reduction of 1,4-NQ, which is consistent with Scheme 3 proposed. Similarly, the molar ratio between CO₂ and DCNQ²⁻ or CNQ²⁻ is 1:1 (Fig. 14B and C). Therefore, their mechanism can be described in Schemes 4 and 5, respectively.

3.4. Theoretical computation

In order to explore the structure of [1,4-NQ-2CO₂]²⁻, [DCNQ-CO₂]²⁻ and [CNQ-CO₂]²⁻, we make some calculations at the level of B3LYP [37]/6-311++G^* with the help of Gauss software. The NBO charges of anion radical (Q⁻), dianion (Q⁻²) and the CO₂ adduct ([1,4-NQ-CO₂]²⁻) were calculated to know which parts possess higher charge densities. Simultaneously, the energies of product were calculated.

In Fig. 15a, for 1,4-NQ⁻, the NBO charges located at the 1,4-position O-centered and the 2,3-position C-centered are −0.68 and −0.28, respectively. And for [1,4-NQ-O-CO₂]²⁻ (Fig. 15a), the NBO charges located at the 4-position O-centered and the 3-position C-centered are −0.82 and −0.34, respectively. It is similar to [1,4-NQ-C-CO₂]²⁻. For DCNQ²⁻ (Fig. 15b), the NBO charges located at the 1,4-position O-centered and the 2,3-position C-centered are −0.80 and −0.21, respectively. It is similar to CNQ²⁻. The calculation results

![Fig. 6. CV (A) and the corresponding 3D spectra (B) in IL containing 20 mmol/L 1,4-NQ with saturated CO₂. CVA (C) and DCVA (D) of 1,4-NQ with saturated CO₂ at 1671, 2342, and 1634 cm⁻¹. To make the DCVA data readily comparable to CV, the DCVA data of 1671 and 2342 cm⁻¹ were multiplied by −1.](image)

![Fig. 7. Reduction of 20 mmol/L DCNQ in BMIMBF₄ with 100% CO₂ (a), 50% CO₂ (b) and without CO₂ (c). The potential scan rate was 10 mV/s.](image)

Fig. 6. CV (A) and the corresponding 3D spectra (B) in IL containing 20 mmol/L 1,4-NQ with saturated CO₂. CVA (C) and DCVA (D) of 1,4-NQ with saturated CO₂ at 1671, 2342, 1634 cm⁻¹. To make the DCVA data readily comparable to CV, the DCVA data of 1671 and 2342 cm⁻¹ were multiplied by −1.

Significantly, the absorbance value of 2342 cm⁻¹ at −2.4 V can reflect the total amount of CO₂ involved in the reaction process. From Fig. 13, at −2.4 V, the mole percent of CO₂ (represented by x) and the corresponding absorbance value at 2342 cm⁻¹ are in a good linear relationship, respectively. The simulation results further prove that CO₂ participates in the reaction process.

Fig. 14 shows the relationship between CO₂ mole percent and its corresponding absorbance value of IR band at 2342 cm⁻¹ at −1.2 V. According to Fig. 14, the corresponding absorbance value decreases dramatically with increase of CO₂ concentrations. However, it decreases slowly at higher CO₂ concentration. At room temperature, the solubility of saturated CO₂ in BMIMBF₄ is 99 mmol/L. For 1,4-NQ (Fig. 14A), the intersection of the two lines is at about 40.43% (about 40 mmol/L), which means that 40 mmol/L CO₂ is captured. Therefore, 2 equiv. of CO₂ can be captured during the electrochemical reduction of 1,4-NQ, which is consistent with Scheme 3 proposed. Similarly, the molar ratio between CO₂ and DCNQ²⁻ or CNQ²⁻ is 1:1 (Fig. 14B and C). Therefore, their mechanism can be described in Schemes 4 and 5, respectively.

3.4. Theoretical computation

In order to explore the structure of [1,4-NQ-2CO₂]²⁻, [DCNQ-CO₂]²⁻ and [CNQ-CO₂]²⁻, we make some calculations at the level of B3LYP [37]/6-311++G^* with the help of Gauss software. The NBO charges of anion radical (Q⁻), dianion (Q⁻²) and the CO₂ adduct ([1,4-NQ-CO₂]²⁻) were calculated to know which parts possess higher charge densities. Simultaneously, the energies of product were calculated.

In Fig. 15a, for 1,4-NQ⁻, the NBO charges located at the 1,4-position O-centered and the 2,3-position C-centered are −0.68 and −0.28, respectively. And for [1,4-NQ-O-CO₂]²⁻ (Fig. 15a), the NBO charges located at the 4-position O-centered and the 3-position C-centered are −0.82 and −0.34, respectively. It is similar to [1,4-NQ-C-CO₂]²⁻. For DCNQ²⁻ (Fig. 15b), the NBO charges located at the 1,4-position O-centered and the 2,3-position C-centered are −0.80 and −0.21, respectively. It is similar to CNQ²⁻. The calculation results
Fig. 8. CV (A) and the corresponding 3D spectra (B) in IL containing 20 mmol/L DCNQ with saturated CO$_2$; CVA (C) and DCVA (D) of DCNQ with saturated CO$_2$ at 1684, 2342, 1510, 1640 cm$^{-1}$. To make the DCVA data readily comparable to CV, the DCVA data of 1684 and 2342 cm$^{-1}$ were multiplied by −1, and 1510 cm$^{-1}$ in the second reduction and the first oxidation were multiplied by −1. The potential scan rate was 10 mV/s.

Scheme 4. The mechanism of electrochemical capture of CO$_2$ via redox cycle of DCNQ.

DCNQ + e $\leftrightarrow$ DCNQ$^-$  \hspace{1cm} (7)

DCNQ$^-$ + e $\leftrightarrow$ DCNQ$_2^{2-}$ \hspace{1cm} (8)

DCNQ$_2^{2-}$ + CO$_2$ $\leftrightarrow$ [DCNQ-CO$_2$]$^{2-}$ \hspace{1cm} (9)

Fig. 9. CV (A) and the corresponding 3D spectra (B) in IL containing 20 mmol/L CNQ with saturated CO$_2$; CVA (C) and DCVA (D) of CNQ with saturated CO$_2$ at 1680, 2342, 1511, 1636 cm$^{-1}$. To make the DCVA data readily comparable to CV, the DCVA data of 1680 and 2342 cm$^{-1}$ were multiplied by −1, and 1511 cm$^{-1}$ in the second reduction and the first oxidation were multiplied by −1. The potential scan rate was 10 mV/s.

Scheme 5. The mechanism of electrochemical capture of CO$_2$ via redox cycle of CNQ.

CNQ + e $\leftrightarrow$ CNQ$^-$  \hspace{1cm} (10)

CNQ$^-$ + e $\leftrightarrow$ CNQ$_2^{2-}$ \hspace{1cm} (11)

CNQ$_2^{2-}$ + CO$_2$ $\leftrightarrow$ [CNQ-CO$_2$]$^{2-}$ \hspace{1cm} (12)
show that O-centered has more electron localization and stronger nucleophilicity than C-centered.

In addition, CO$_2$ reacts with 1,4-NQ$^{-}$ instead of DCNQ$^{-}$ or CNQ$^{-}$. The reason maybe the electron-withdrawing effects of chlorine group, which making the charge densities of DCNQ or CNQ smaller. So the nucleophilicity of DCNQ$^{-}$ or CNQ$^{-}$ gets lower and it is difficult to attack the C-centered of CO$_2$. For this reason, the charges of DCNQ$^{-}$ and CNQ$^{-}$ are calculated (Fig. 15c). Compared the charges of 1,4-NQ$^{-}$. DCNQ$^{-}$ with CNQ$^{-}$, obviously 1,4-NQ$^{-}$ has higher charge densities.
Subsequently, the energies of product are calculated and the results are listed in Fig. 16. For 1,4-NQ, for its initial adduct [1,4-NQ-CO$_2$]$^-$ (Fig. 16a1), the calculation results show that carbonate adduct [1,4-NQ-O-CO$_2$]$^{2-}$ is 0.20 eV higher than carboxylate adduct [1,4-NQ-C-CO$_2$]$^{2-}$. The results of energy calculation are similar to the calculation results of [BQ-CO$_2$]$^-$ [24]. And for the product [1,4-NQ-2CO$_2$]$^{2-}$ (Fig. 16a2), the results of energy calculation show that carbonate-carboxylate type adduct ([1,4-NQ-O-C-2CO$_2$]$^{2-}$) has the lowest energy. Although the results of energy calculation which show carbonate-carboxylate type adduct [1,4-NQ-O-C-2CO$_2$]$^{2-}$ is more stable than bis(carbonate) adduct [1,4-NQ-O-O-2CO$_2$]$^{2-}$ adduct, bis(carbonate) adduct is reported as the product in the literature [19]. In addition, the chemical reaction of the electroreduction product with CO$_2$ is a nucleophilic reaction, so from the results of the NBO charge calculation (Fig. 15a), it can be speculated that its reaction mechanism may follow the EC$_2$EC$_2$ mode of the reaction (13) in Scheme 6 more reasonably.

For CNQ (Fig. 16b), carbonate adduct ([CNQ-O-CO$_2$]$^{2-}$) is 0.23 eV lower than carboxylate adduct ([CNQ-C-CO$_2$]$^{2-}$). Whereas for DCNQ (Fig. 16c), from the perspective of reversibility of the whole reaction, the chlorine atom cannot be replaced by CO$_2$. Therefore, it is impossible to produce carboxylate product ([DCNQ-C-CO$_2$]$^{2-}$), and carbonate product ([DCNQ-O-CO$_2$]$^{2-}$) is more reasonable.

The calculation results indicate that the CO$_2$ adduct should be carbonate-type structure. Therefore, their mechanisms can be elaborated in Scheme 6. In fact, carbonate-type product has also been reported as the main product [19,24,43].

4. Conclusions

In summary, CV and IR spectroelectrochemistry techniques were used to investigate the mechanism of electrochemical capture of CO$_2$ via redox cycle of 1,4-NQ, DCNQ and CNQ in BMIMBF$_4$, respectively. Our research shows that chlorine group plays an important role in electrochemical capture of CO$_2$ via redox cycle of quinones.

The redox mechanisms of 1,4-NQ and DCNQ in IL both follow the reversible EE mode, while CNQ follows the non-reversible mode.
However, in the presence of CO$_2$, 1,4-NQ, DCNQ and CNQ follow the modes of EC$_r$EC$_r$, EEC$_r$ and EEC$_r$, respectively. The results show that the reduced quinones can be well used as nucleophiles to capture CO$_2$. Quantitatively, it is found that both the molar ratio between CO$_2$ and 1,4-NQ$^-$ or [1,4-NQ-CO$_2$]$^{2-}$ is 1:1; whereas the molar ratio of CO$_2$ and DCNQ$^{2-}$ or CNQ$^{2-}$ is 1:1. CO$_2$ adduct should be carbonate-type products according to the theoretical calculation results. In addition, CO$_2$ can be released during the redox cycle of quinones since the chemical reaction of the electroreduction product with CO$_2$ is reversible. Therefore, 1,4-NQ, DCNQ and CNQ are promising adsorbents, which can be regenerated easily.

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Appendix A. Supplementary data

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References


