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Electrocatalytic arsenite oxidation in bicarbonate solutions combined with CO₂ reduction to formate

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Graphical Abstract
Abstract

Sunlight-driven water-energy nexus technologies are receiving increasing attention. This study presents a hybrid electrochemical system that catalyzes the oxidation of As(III) to As(V) with a nanoparticulate TiO$_2$ electrocatalyst (Ti/Ir$_{1-x}$Ta$_x$O$_y$/TiO$_2$; denoted as an n-TEC) while simultaneously converting CO$_2$ to formate on a Bi electrode in aqueous bicarbonate solutions at circum-neutral pH. Linear sweep voltammograms of n-TEC exhibit a specific As(III) oxidation peak ($E_{p,As}$), at which the Faradaic efficiency (FE) of As(V) production is $\sim$100%. However, the application of a potential higher than the peak ($E > E_{p,As}$) leads to a significant decrease in the FE due to water oxidation. Upon the addition of chloride, the oxidation of water and chloride occur competitively, producing reactive chlorine species responsible for mediating the oxidation of As(III). The Bi electrodes synthesized via the electrodeposition of Bi$^{3+}$ typically show high FEs of $>$80% for formate production in bicarbonate solution purged with CO$_2$. The addition of chloride significantly enhances the current while maintaining the FE. The n-TEC catalyst and Bi electrodes are paired in a single device equipped with a membrane, and significant effort is made to achieve the same FEs in both the anodic and cathodic reactions as in their half-reactions. Finally, the optimized n-TEC/Bi pair is coupled with a low-cost, commercially available photovoltaic (PV). Various technical factors that drive the overall reactions with the PV are considered, and maximum FEs of $\sim$95% are achieved for the production of both As(V) and formate.

Keywords

Redox reactions; Electrocatalysis; Water treatment; Solar fuels; Water-energy nexus
1. Introduction

Solar CO₂ reduction reactions (CO₂RRs) have received growing attention not only for CO₂ mitigation but also for the production of value-added building-block chemicals [1-3]. There are several types of CO₂ conversion photosystems, including photocatalysis, photoelectrocatalysis, and photovoltaic (PV)-assisted electrocatalysis (Table S1). Among them, the last has been considered the most promising due to the easy fabrication, high efficiency, and scalability. Whereas many CO₂ conversion products (C1-C6) are possible, formate is considered one of the most-cost effective ($0.4 \text{ kWh}^{-1}$), and hence, significant efforts have been made to achieve the selective production of formate at low overpotential and high current [4-9]. Most CO₂RRs have been evaluated in terms of the yield and selectivity of formate [1], particularly in aqueous solutions containing bicarbonate (HCO₃⁻), which enhances the CO₂ conversion rate [10,11] and buffering effect, at circum-neutral pH (6.5–8.5). Unfortunately, bicarbonate is not a suitable electrolyte for the oxygen evolution reaction (OER) via water oxidation, which is a counter half-reaction that supplies CO₂ with proton and electron (H⁺/e⁻) pairs [6,12,13]. It is strongly adsorbed onto the OER electrodes via mono- and bidentate modes [14,15], interfering with the molecular interaction of H₂O with the electrode surface. Furthermore, bicarbonate can be oxidized into carbonate radical anions (CO₃²⁻) by hydroxyl radicals (OH⁺), which are a transient intermediate during the water oxidation, at a relatively high bimolecular rate constant (~10⁷ M⁻¹ s⁻¹) [16].

Other value-added anodic reactions can be employed as alternatives to the OER to supply CO₂ with H⁺/e⁻ pairs (Table S1). One example is the electrocatalytic advanced oxidation reaction of toxic and recalcitrant aquatic substrates via direct and mediated charge
transfer [17-19]. In the former charge transfer, transiently oxidized electrocatalysts withdraw electrons directly from the substrate. In the latter, reactive oxygen and chlorine species (ROS and RCS, respectively) produced via one- or multi-electron oxidations of water (e.g., free OH• and O1, respectively) [19,20] and two-electron oxidation of chloride (2Cl− → Cl2; Cl2 + H2O → HClO + Cl− + H+) mediate the oxidation of the substrate in the bulk solution [21,22]. OH• is preferred to RCS in terms of reactivity [16]; however, its production requires greater energy (2.7 V) than that of RCS (1.3–1.5 V). Coupling of the advanced oxidation reaction with the hydrogen evolution reaction (HER) is well established, as the HER is rather straightforward and proceeds well under various conditions [22-26].

With this in mind, we attempted to simultaneously drive the advanced oxidation reaction and CO2RR with nearly 100% faradaic efficiencies (FE) for both (Scheme 1). To achieve this, Ir and Ta mixed oxides (Ir1-xTaxOy) were coated onto Ti foils as an underlayer for an ohmic contact, onto which TiO2 nanoparticles were coated as an electrocatalytic overlayer [23,27]. As-synthesized nanoparticulate TiO2 electrocatalysts (denoted as n-TECs) were utilized for the oxidation of As(III) to As(V) in aqueous bicarbonate solutions purged with CO2 at circum-neutral pH (e.g., H3AsIII O3 + H2O → H2AsV O5− + 2e− + 3H+). Compared to As(III), As(V) is less toxic to humans, while the latter is more removable with conventional techniques (e.g., coagulation, ion exchange, membrane, and adsorption [28-31]. In bicarbonate-free aqueous solutions, As(III) can be oxidized by several ROSs (e.g., *OH, *HO2/O2•−, O3, and H2O2) via advanced oxidation processes [32] including Fenton [33], photolysis [28], and photocatalysis [34]. In the presence of chloride, RCSs can play a primary role in the As(III) oxidation particularly in electrocatalysis [26]. In groundwater with (bi)carbonate, pre-adsorbed As(III) onto metal oxides is released via the formation of arseno-carbonate complexes,
increasing the As(III) concentration in groundwater [35-37]. For the CO$_2$RR, Bi electrodes were quickly synthesized via the electrodeposition of Bi$^{3+}$ onto Ti foils, and the performance of the Bi electrodes for formate production was optimized. The addition of naturally abundant chloride to the bicarbonate solutions not only mediated the As(III) oxidation via RCS but also enhanced the formate production. Finally, n-TEC and Bi electrode pairs were separated in a two-compartment cell divided by a proton-exchange membrane and coupled with a PV cell to drive the overall redox reactions using sunlight.

2. Experimental section

2.1. Synthesis and characterization of the electrodes

For As(III) oxidation, double-layered electrodes consisting of an Ir$_{1-x}$Ta$_x$O underlayer and a nanoparticulate TiO$_2$ overlayer (denoted as n-TEC) were synthesized following a typical coating-annealing process [21,27,38]. In brief, Ti substrates (Grade 2, ~1 mm thick) were polished with sandpaper and washed with acetone, ethanol, and finally deionized water (>18 MΩ·cm, Human Corporation) in an ultrasonic bath (DAIHAN Scientific Co., Ltd). Then, they were coated with Ir (H$_2$IrCl$_6$) and Ta (TaCl$_5$) at a molar ratio of 73/27, followed by annealing at 500 °C for 1 h [38]. The XPS analysis of Ir and Ta for the as-synthesized Ir$_{0.73}$Ta$_{0.27}$O$_x$ sample confirmed the composition ratio (Fig. S1). A suspension of TiO$_2$ nanoparticles synthesized via a typical sol-gel process (Nanopac, Korea) was sprayed onto both sides of the substrates as an overcoat, followed by calcination at 425 °C for 5 h. Our previous studies on the electrodes showed that the underlayer was essential for ohmic contact and durability [17,21,27], and that the TiO$_2$ overcoat required metal dopants (e.g. Bi and Nb)
[27,39]. However, a further study revealed that the dopant-free TiO\textsubscript{2} electrode exhibited essentially the same performance as the doped one [40].

For the CO\textsubscript{2}RR, Bi electrodes were synthesized on the Ti substrates via an electrodeposition process [41]. After drying, the substrates were treated at 500 °C for 6 h in an air atmosphere. The Bi precursor solution was prepared by dissolving bismuth(III) nitrate pentahydrate (50 mM, Junsei) in an aqueous solution containing lactic acid (3 M, Sigma Aldrich, ≥85%) and polyvinylpyrrolidone (5 g L\textsuperscript{-1}, Sigma Aldrich, M.W. 360,000) at pH 3. The electrodeposition was performed in the Bi precursor solution using a typical three-electrode system with the as-prepared Ti substrates as the working electrode, a saturated calomel electrode (SCE) as the reference electrode, and a Pt wire as the counter electrode. A negative potential of −0.847 V vs. SCE was applied to the working electrodes for varying lengths of time (3–60 s). The electrodes were then washed with an ethanol/water mixture and dried [41].

The morphology of the as-synthesized n-TEC and Bi electrodes was examined using a scanning electron microscope (SEM, Su-8230, Hitachi) at an accelerating voltage of 15.0 kV. An energy dispersive spectrometer (EDS, X-MaxN80, Horiba) and an X-ray diffractometer (XRD, Max-2500 V, Rigaku) with Cu K\textalpha radiation (λ =1.5406 Å) were employed to examine elemental composition and crystalline structure, respectively. An n-TEC sample was cut with focused ion beam (FIB, Versa3D LoVac, FEI Company) and its cross-sectional surface was characterized using field-emission electron probe microanalysis (EPMA, JXA-8530F(5CH), JEOL).
2.2. Electrolysis and chemical analysis

The electrochemical behaviors of the n-TEC and Bi electrodes were examined in a customized, single-compartment reactor using a potentiostat/galvanostat (Ivium). Either n-TEC (1 × 1 cm² for each side) or Bi electrodes (1 × 1 cm²; a single side only), and SCE and Pt foil were immersed in aqueous solutions of KHCO₃ (0.1 M, Sigma Aldrich, 99.7%) in the absence or presence of CO₂ (99.9999%) purging. The CO₂ purging decreased the electrolyte pH from ~8.5 to ~6.8. For the n-TEC electrodes, the potential was swept from −0.5 to +1.4 V vs. SCE at a rate of 5 mV·s⁻¹ in the absence or presence of As(III) (1 mM, NaAsO₂, Sigma Aldrich, ≥90%). For the Bi electrodes, the potential was swept from −0.4 to −2.0 V vs. SCE at the same scan rate. If necessary, NaCl or NaClO₄ (0.1 M, Sigma-Aldrich, >99.0%) was added to the electrolyte.

The bulk electrolysis for As(III) oxidation using the n-TEC was performed at various potentials (0.75−1.2 V vs. SCE). During electrolysis, aliquots were intermittently sampled and analyzed using the well-known molybdenum blue method for As(V) quantification [26,42]. In brief, aliquots of various volumes (0.1−0.5 mL) were mixed with the molybdate reagent solution (0.2 mL, Sigma-Aldrich) and L-ascorbic acid (0.1 mL, Sigma-Aldrich), and then diluted to a volume of 2 mL using deionized water (18 MΩ cm). After 1 h, the samples were analyzed using a UV-Vis spectrometer (T-60, PG Instruments) at λ = 870 nm. If necessary, As(III) was quantified using high performance liquid chromatography (HPLC, Waters 2695) with an Aminex HPX-87H ion exclusion column (Bio-Rad, 300 × 7.8 mm²) equipped with a UV detector (λ = 210 nm). 5 mM sulfuric acid was used as an eluent and flowed at 0.6 mL
The HOCl concentration was estimated using a colorimetric method with UV-Vis spectrometer (UV-2450, SHIMADZU) [21,43]. Formate, which was the primary product of the CO$_2$RR, was quantified using high performance liquid chromatography (HPLC, Waters Co.) with an Aminex HPX-87H column (BIO-RAD, USA) and a proton nuclear magnetic resonance (H-NMR; Avance III 500 MHz, Bruker) [7,8,44]. If necessary, H$_2$ in the reactor headspace was analyzed by gas chromatography equipped with thermal conductivity detector (GC-TCD, Agilent 7820). Details of the analytical method can be found elsewhere [6-8]. The concentrations of total chlorine were determined with $N,N$-diethyl-$p$-phenylenediamine (DPD) reagent (Hach methods). Hereafter, the unit for electrochemical potential is expressed as V only; “versus SCE” is omitted for simplicity, unless otherwise specified.

For As(III) oxidation and simultaneous CO$_2$RR, a two-compartment Teflon reactor separated by a proton exchange membrane (Nafion 117, Chemours) was designed. n-TEC and Bi electrodes of known area were placed in separate compartments containing 0.1 M KHCO$_3$ in the absence or presence of CO$_2$ purging. 1 mM As(III) was added to the anolyte. Various DC voltages (E$_{DC}$) were applied to the n-TEC (anode) and Bi (cathode) pairs, and the current ($I_{DC}$) was recorded using a DC power supply (E3647A, Keysight Technologies). To estimate the bias potentials at the n-TEC and Bi electrodes at a given E$_{DC}$, an SCE and Pt wire were placed in either the anode or cathode compartment, and the open circuit potentials of each electrode were measured using a potentiostat (Ivium). If necessary, a known potential was applied to the Bi electrode using the potentiostat. In this case, the n-TEC electrode acted as the counter electrode. For the stand-alone overall processes, a commercial, monocrystalline Si PV cell (Solarcenter Inc.) was connected to the n-TEC and Bi electrode pair and irradiated with simulated sunlight (LS-150-Xe, Abet Technologies, Inc., USA, AM 1.5; 100 mW-cm$^{-2}$). All the electrocatalytic
reactions were performed at least twice.

3. Results and discussion

3.1. Electrocatalytic arsenic oxidation using the n-TEC electrodes.

As-synthesized n-TEC samples were characterized with EPMA (Fig. 1a and b). The cross-sectional view of the samples obviously shows the sub-micrometer-thick interlayer between the underlying Ti substrate and ~5.5 \( \mu \text{m} \) thick TiO\(_2\) overlayer. The interlayer composed of Ir and Ta was uniformly distributed throughout the Ti substrate, whereas our previous attempt led to a localized distribution of Ta [45]. The energy dispersive X-ray spectroscopic analysis of the cross-section further reveals the fractional composition of 87%, 7.6%, and 5.4% for Ti, Ir, and Ta, respectively (Fig. S2). The XRD patterns exhibit that TiO\(_2\) was predominant, whereas IrO\(_2\) (2\( \theta \) = 34.46° and 58.30° for 101 and 220 planes, respectively) was trace and no Ta-oriented peaks were observed (Fig. 1c). Such the weak IrO\(_2\) and absent TaO\(_x\) peaks could be attributed to the thick TiO\(_2\) overlayer that inhibits the penetration of the X-ray beam. The primary phase of the TiO\(_2\) overlayer was anatase at ~82% (i.e., rutile of ~18%) [45] due to a relatively low annealing temperature (425 °C) (Fig. S3). The particle size of TiO\(_2\) was ~20 nm (Fig. S4), similar to a previous study [45].

Fig. 2a shows the linear sweep voltammograms (LSVs) of n-TEC electrodes in aqueous bicarbonate (BC, 0.1 M) solutions (without or with CO\(_2\) purging) in the absence and presence of As(III) (1 mM). In the pure bicarbonate solution, the onset potential (\( E_{on} \)) was ~0.8 V and \( I \) increased to ~2 mA at \( E = 1.1 \) V (versus SCE; hereafter omitted for simplicity unless otherwise specified). CO\(_2\) purging did not influence the LSV profile, although the solution pH
decreased slightly from ~8.6 (CO₂-free BC) to ~6.8 (CO₂-purged BC). The addition of As(III) to the CO₂-free bicarbonate solution generated a characteristic peak current ($I_{p,As}$) at ~0.8 V ($E_{p,As}$), followed by a current profile similar to that obtained in the As(III)-free solution. The same magnitude of $I_{p,As}$ with an $E_{p,As}$ shift of ~Δ50 mV was observed when CO₂ was purged into the bicarbonate/As(III) solution. The deviation from *Nernstian* behavior (ΔE = 59 mV/pH ~106 mV) might be attributed to the strong adsorption of carbonic species (HCO₃⁻ and/or dissolved CO₂) onto the TiO₂ surface and the consequent Fermi level pinning-like effect [46]. This interaction was altered by As(III), as shown by the appearance of $I_{p,As}$ in the presence of As(III). It has been postulated that aqueous As(III) (i.e., As$^{III}$(OH)$_{3-x}$O$_{x}$; $pK_{a1} = 9.23$ [47]) can act as a Lewis base to form the complexes with bicarbonate (e.g., As(OH)$_2$CO$_3$⁻, As(CO$_3$)$_2$⁻, and As(CO$_3$)$_3$⁺) [35,36,48,49]. Among these, As(OH)$_2$CO$_3$⁻ is considered to be the most abundant at circum-neutral pH (R1) [49,50], and is predominantly adsorbed onto the TiO₂ surface (R2).

\[
\text{As(OH)}_3 + \text{HCO}_3^- \rightarrow \text{As(OH)}_2\text{CO}_3^- + \text{H}_2\text{O}, \quad K_c = 0.22 \pm 0.04 \quad \text{(R1)}
\]

\[
>\text{Ti-OH} + \text{As(OH)}_2\text{CO}_3^- \rightarrow >\text{Ti-OH}--\text{As(OH)}_2 + \text{CO}_3^- \quad \text{(R2)}
\]

\[
>\text{Ti-OH} + \text{As(OH)}_3 \rightarrow >\text{Ti-O}--\text{As(OH)}_3^- + \text{H}^+ \quad \text{(R3)}
\]

where $K_c = [\text{As(OH)}_2\text{CO}_3^-] / (\gamma_0 [\text{As(OH)}_3] \times [\text{HCO}_3^-])$ and $>\text{Ti-OH}$ refers to the TiO₂ surface.

In $K_c$, $\gamma_0$ represents the activity coefficient for a neutral species. Assuming $\gamma_0 = 1$, only ~2% of the total As(III) is calculated to exist as complexes (i.e., [As(OH)$_2$CO$_3$⁻] ~2.2×10⁻⁵ M), and the free As(III) (i.e., not complexed with carbonic species) can be adsorbed onto the TiO₂ surface (R3). The changes in the LSV induced by the addition of As(III) to aqueous chloride and
perchlorate solutions free of carbonic species (Fig. S5) support the direct interaction of the free As(III) with the TiO₂ surface at circum-neutral pH, at which both TiO₂ and As(III) are neutral in charge. The addition of chloride to bicarbonate had an insignificant influence on the LSV profiles (Fig. 2a), suggesting that the chloride did not directly affect the As(III)-TiO₂ interaction at E_{p,As}.

The bulk electrolysis of As(III) was performed in aqueous bicarbonate solutions in the absence and presence of chloride (see Fig. S6 for the CO₂-purged condition). As shown in Fig. 2b, three bias potentials (E = 0.75, 1.0, and 1.2 V) were selected for the As(III) oxidation (E°(As^{V}/As^{III}) = −0.188 V at pH 8.6) [51]. At all three potentials, As(V) was produced linearly for 2 h, and its production accelerated with increasing E (i.e., I) (Fig. 2c). However, the FE did not exhibit a simple E dependency; the FE was ~100% at E = 0.75 V, and decreased to ~40 and 13% at E = 1.0 and 1.2 V, respectively (Fig. 2d). Considering that E_{p,As} is close to 0.75 V (Fig. 2a), the ~100% FE value at E = 0.75 V indicates that the direct oxidation of free or complexed As(III) to As(V) occurred efficiently. The FE values at ~100% in different supporting electrolytes further confirm that the direct oxidation of As(III) occurs efficiently as long as the interfacial charge transfer is not altered (Fig. S7). The XPS analysis of the n-TEC electrode confirmed the As(III) oxidation to As(V) after 2 h-electrolysis (Fig. S8). It is noteworthy that although the fraction of As(V) was only ~20% of a total As (i.e., As(III) = 0.8 mM; As(V) = 0.2 mM), the fraction of As(V) (~55%) on the n-TEC surface was greater than that of As(III) (~45%). This suggests that the adsorption of As(V) is favored over that of As(III). While As(III) oxidation and water oxidation (E°(O₂/H₂O) = 0.48 V at pH 8.6) occur competitively at E = 1.0 V, the latter becomes predominant at E = 1.2 V, significantly decreasing the FE of As(III) oxidation.
The presence of chloride did not influence the As(III) oxidation rate at \( E = 0.75 \) and 1.0 V; however, it enhanced the oxidation rate by a factor of two at \( E = 1.2 \) V. These changes could be attributed to \( I \), which was the nearly same at \( E = 0.75 \) and 1.0 V, but increased at \( E = 1.2 \) V upon the addition of chloride (Fig. 2c). Considering the chlorine evolution reaction (CER, \( E^\circ(\text{Cl}_2/\text{Cl}^-) = 1.12 \) V), the application of \( E = 0.75 \) V would be insufficient to directly drive the chloride oxidation. Hence, the FE was nearly the same (~100%) at \( E = 0.75 \) V, where the direct As(III) oxidation can occur [51]. This further indicates that the chloride interrupted neither the As(III) adsorption nor the direct charge transfer for the adsorbed As(III). However, a high potential (1.2 V) can oxidize chlorides into RCSs (e.g., \( \text{Cl}_2^- \) and \( \text{HClO}/\text{ClO}^- \)), effectively mediating As(III) oxidation (R4 and R5).

\[
\text{As(OH)}_3 + 2\text{Cl}_2^- + \text{H}_2\text{O} \rightarrow \text{H}_2\text{AsO}_4^- + 4\text{Cl}^- + 3\text{H}^+ \quad \text{(R4)}
\]

\[
\text{As(OH)}_3 + \text{HOCl} \rightarrow \text{H}_2\text{AsO}_4^- + 2\text{H}^+ + \text{Cl}^- \quad \text{(R5)}
\]

In this case, the pre-adsorption of As(III) is not essential because the RCSs can diffuse into the bulk solution. Such heterogeneous reaction-induced homogeneous bulk reactions are often observed in CER-mediated processes [20,52]. Similarly, other halides (e.g., the \( I_3^-/I^- \) couple) have been shown to mediate As(III) oxidation in the bulk [28]. It is noteworthy that although the \( E^\circ \) for the CER is ~0.6 V greater than that of the OER at pH ~8.6, the former involving 2e\(^-\) transfer is kinetically faster than the latter, requiring a lower overpotential [53]. This should lead to a two-fold enhancement of the FE from 13% to 26%. Considering the FE of 30-35% for the CER during electrolysis in the absence of As(III) (Fig. S9), ~90% of the produced RCSs (i.e., \( \text{FE}_{\text{As(V)}} / \text{FE}_{\text{CER}} \)) in the presence of As(III) were estimated to be used to mediate the oxidation of As(III).
To further confirm the RCSs-mediated As(III) oxidation, pre-electrolysis with n-TEC at E = 1.2 V proceeded for 2 h in the absence of As(III) and As(V). Immediately after the electrolysis off, As(III) at 1 mM was spiked to bicarbonate solutions (with and without chloride) and the concentration changes in As(III) and As(V) were examined (Fig. 3). As a representative of RCSs, HOCl (i.e., HOCl/OCl⁻ couple with pKa of 7.46) was quantified because of the interference effect of bicarbonate on the quantification of total RCSs. In the bicarbonate solution with chloride, the pre-electrolysis led to the gradual production of HOCl to ~0.2 mM in 2 h (Fig. 3a). Immediately after the addition of As(III), the concentrations of As(III) and HOCl dropped to ~0.6 mM and zero, respectively, while As(V) at ~0.35 mM was produced (hence total As concentration of ~0.95 mM, nearly matching the spiked As(III) concentration at 1 mM). Considering the equimolar reaction (R5), the larger change in the As(V) concentration than that of HOCl during the post-electrolysis period suggests the involvement of other RCSs (e.g., Cl₂•⁻) in the As(III) oxidation [19-22,24,25,27,52]. The reactivity of RCSs significantly varies depending on the types of the species (e.g., Cl₂•⁻ vs. HOCl) and the substrates (e.g., aliphatic acids vs. aromatics) [21]. If Cl₂•⁻ is predominant, then the oxidation rate can be comparable to that with ROSs (e.g., OH•) [19,52]. However, direct comparison of Cl₂•⁻ and OH• for As(III) oxidation in the same electrochemical system is not straightforward, because i) the rate constant with OH• is >10⁹ M⁻¹ s⁻¹ [16] whereas that for Cl₂•⁻ is not available in literature, and ii) RCSs are readily produced at high efficiency whereas the production efficiency of ROSs is not high. Particularly in the latter, a chloride electrolyte favors RCSs regardless of the anode material [52], dramatically shifting the reaction mechanism from the ROSs-based to the RCSs-based in most electrochemical systems [19]. In the bicarbonate solution without chloride, the production of neither As(V) nor HOCl was observed and the
spiked As(III) was maintained without any change (Fig. 3b).

3.2. Electrocatalytic CO$_2$ reduction using Bi electrodes

The as-synthesized Bi electrodes fabricated via electrodeposition (ED) were composed of many Bi particle aggregates with a uniform size of ~170 nm (Fig. 4a and b; see Fig. S10 for more images). They were homogeneously distributed onto Ti substrate with an overall film thickness of 100-200 nm. The XRD analysis of the sample showed crystalline, rhombohedral structure (JCPDS# 85-1330) with predominant planes of (012), (104), (110), (116), and (214) at $\theta = 27.18^\circ$, $38.00^\circ$, $39.72^\circ$, $62.23^\circ$ and $70.62^\circ$, respectively (Fig. 4c). The LSVs of the as-synthesized Bi electrodes always exhibited a cathodic peak at $E = -0.84$ V in the first potential sweep, which disappeared in the second sweep (Fig. 4d). The Bi particles appeared to be partially oxidized while drying in air after ED. In the second cathodic scan, the oxidized species (e.g., BiO$_x$) should have been reduced to metallic Bi (Bi$^0$).

The electrocatalytic behavior of the Bi electrodes was compared under N$_2$ and CO$_2$ purging (Fig. 5a). The $E_{on}$ values were estimated to be $-1.6$ and $-1.4$ V, respectively; the difference between them ($-0.2$ V) was partially attributed to the reduction of the pH from ~9 under N$_2$ purging to 6.8 under CO$_2$ purging ($\Delta E = 59$ mV/pH $\sim 0.13$ V). It is noteworthy that the $I$ under CO$_2$ purging was significantly greater than that under N$_2$ purging. This suggests that Bi could effectively catalyze the CO$_2$RR while inhibiting the HER in bicarbonate solution. To confirm the shift from the HER to the CO$_2$RR, bulk electrolysis with N$_2$ purging was performed at $E = -1.6$ V in a single compartment cell; a stable $I$ of $\sim 3.5$ mA·cm$^{-2}$ flowed over 2 h with an FE of $\sim 40\%$ for H$_2$ (Fig. S11). On the other hand, under CO$_2$ purging, a trace of H$_2$
was evolved (FE < 1%) despite a similar \( I \) value at the same \( E \) (Fig. S11); instead, formate was produced linearly with time as the only carbon-containing product with \( \text{FE of } \sim 80\% \) (Fig. 5c). Such the formate production continued over 12 h with the same \( \text{FE} \) (Fig. S12). No other chemicals were found in the gas or liquid phases (Fig. S13); if any were produced, their concentrations were below the detection limit. As the \( \text{ED time} \) was increased from 3-60 s, \( \text{E}_{\text{on}} \) gradually shifted to positive potential and a larger \( I \) flowed (Fig. 5a inset) due to the greater coverage of the Ti substrate by Bi particles (Fig. S10). However, regardless of the \( \text{ED time} \), the \( \text{FE} \) remained nearly the same at 80\% (Fig. 5b). This suggests that the CO\(_2\)RR occurred only on the Bi surface and that the bare Ti surface was inactive for Faradaic charge transfers. In addition, the decreased \( \text{FE values} \) (60–70\%) at low \( E \) values (−1.4 and −1.5 \( V \)) indicate that a larger overpotential was required for formate production (\( \text{E}^\circ\)(CO\(_2\)/HCOOH) = ca. −1 \( V \) at pH 6.5). At \( E = −1.7 \) and −1.8 \( V \), a larger \( I \) flowed and consequently, the formate production was enhanced; however, the maximum \( \text{FE} \) was still ~80\% at both potentials. The “magic” \( \text{FE value} \) of 80\% was attributed to the fact that presumably, the produced formate was partially oxidized on the Pt in the single compartment cell. To avoid the possible unwanted reoxidation of formate, the Bi electrode was separated from the Pt counter electrode via a proton-exchange membrane. In this two-compartment cell, a larger amount of formate was produced with an \( \text{FE of } >95\% \) (Fig. 5c).

The addition of chloride to the bicarbonate solution further enhanced the electrocatalytic performance of the Bi electrodes; \( \text{E}_{\text{on}} \) was nearly unchanged whereas \( I \) was significantly improved at \( E \) values active for CO\(_2\)RR (Fig. 5a). In bulk electrolysis at \( E = −1.6 \) \( V \), the \( I \) was approximately two-fold larger, and a proportional increase in the formate production was observed (hence, \( \text{FE } >95\% \) (Fig. 5c and S14). It has been postulated that
halides can be specifically adsorbed onto metal cathodes (e.g., Cu and Au) and mediate the charge transfer between the electron-deficient carbon atom of CO$_2$ (or H$^+$) and the metal cathodes [54-57]. Such enhanced van der Waals interactions usually enhance $I$, changing the product distribution and as a result the FE values. The same effect appeared to play a role in increasing $I$ without altering the product selectivity in this study. Compared to Cl$^-$, other halides (I$^-$ and Br$^-$) have been reported to exert greater effects (e.g., Cl$^- <$ Br$^- <$ I$^-$) due to stronger interactions [55]. However, in this study, they were found to have a negative effect on the formate production and FE (Fig. S15). Strong interaction between the halides and the metal surface can facilitate charge transfer from the metal to interfacial CO$_2$; however, at the same time, such strong adsorption can block and poison the active sites for CO$_2$RR. Chloride appears to have a moderate strength interaction with the Bi surface, efficiently transferring electrons to nearby CO$_2$ while being readily desorbed from the surface.

3.3. Designing an electrocatalytic device for arsenic oxidation and CO$_2$ reduction

To drive the arsenic oxidation and CO$_2$ reduction reactions simultaneously in a single cell, the LSVs of n-TEC and Bi electrodes (as an anode and a cathode, respectively) of various sizes were compared in bicarbonate solutions purged with CO$_2$ (Fig. S16). As the size was increased, more current flowed, and hence, the DC voltage required for cell operation ($E_{cell}$) was decreased. Based on the optimal anodic and cathodic half-reaction potentials (e.g., $E_a \sim 0.75$ V and $E_c \sim -1.6$ V), an $E_{cell}$ greater than 2.35 V$_{DC}$ must be applied to drive both reactions. However, the simple application of this $E_{cell}$ value based on the difference between $E_a$ and $E_c$ did not actually result in the same performance as those of the half-reactions. For example, n-
TEC and Bi pairs with area ratios of 16-64 at $E_{\text{cell}} = 2.35 \text{ V}_{\text{DC}}$ led to FEs of 70-90% and 50-70% for the production of As(V) and formate, respectively (Fig. S17). This suggested that a more rational design considering factors associated with the cell resistance (e.g., due to the membrane) and the different catalytic activity of the n-TEC and Bi electrodes was required for energy-efficient redox reactions.

After considering the potential drop across the employed membrane (~0.1 V), we focused on achieving a high FE value for As(III) oxidation rather than for CO$_2$RR, because the FE of the former is highly susceptible to $E$ (Fig. 2d vs. 5b). Then, various $E_{\text{cell}}$ values (1.5-3.0 V$_{\text{DC}}$) with different electrode area ratios were tested to find the optimal conditions for both the anodic and cathodic reactions (Fig. S18). The first optimal condition was an $E_{\text{cell}}$ value of 2.5 V$_{\text{DC}}$ with an n-TEC/Bi area ratio of 32 (8 cm$^2$/0.25 cm$^2$; denoted as EC-I pair), in which $E_a$ and $E_c$ were maintained at +0.88 V and −1.62 V, respectively. The choice of a large n-TEC electrode was attributed to the high FE at low current density (i.e., small bias) for As(III) oxidation compared to the large potential bias of CO$_2$RR. It is noteworthy that the obtained $E_a$ (0.88 V) was close to $E_{p,\text{As}}$ (0.75 V) (Fig. 2a), suggesting direct As(III) oxidation was possible. Accordingly, high FEs (~90%) were obtained for the As(III) oxidation as well as the CO$_2$RR (Fig. 6a and b). Unfortunately, $I_{\text{cell}}$ was small (max. ~1 mA) under these conditions (Fig. S19), and the overall redox reaction was slow. To increase the reaction rate, a larger $E_{\text{cell}}$ (3 V$_{\text{DC}}$) was applied to an electrode pair (EC-II pair) with an area ratio of 2. At this voltage, with $I_{\text{cell}} = 4$-5 mA (Fig. S19 an $E_a$ of 1.26 V and $E_c$ of −1.74 V were obtained. As expected from Fig. 2 and 5, the FE of formate production was nearly 100%, whereas the As(V) production was poor with an FE of ~10% due to unwanted water oxidation (Fig. 6a and b). To inhibit the water oxidation and enhance the As(III) oxidation while not affecting the CO$_2$RR, chloride was added to
bicarbonate in both compartments. Consistent with Fig. 2 and 5, the production of both As(V) and formate was enhanced, with the effect being more pronounced for the former. In the presence of chloride, the FE of formate production was nearly the same (~95%), while that of As(V) was doubled (~22%) due to the RCS-mediated As(III) oxidation (R4).

Based on the above experiments, the electrode pair was coupled to PVs to drive the overall reaction for off-grid operation. The LSVs of two anode/cathode pairs (EC-I and EC-II, with area ratios of 32 and 2, respectively) were compared to the I-V curves of the PVs (PV-I and PV-II, with power conversion efficiencies of 12.4 and 12.1%, respectively) (Fig. 6c). This comparison suggested that the coupling of EC-I and PV-I could drive direct As(III) oxidation and simultaneous CO₂RR at an $E_{\text{cell}}$ of 2.43 V and $I_{\text{cell}}$ of 1.47 mA with a maximum solar conversion system efficiency of ~9% (= $E_{\text{cell}} \times I_{\text{cell}} \times 100% / \text{PV area} / 100 \text{ mW-cm}^{-2}$). This efficiency is quite comparable to those of PV-electrocatalysis hybrid systems (Table S1). As expected, both As(V) and formate were produced linearly with time, leading to an FE of ~95% for both reactions (Fig. 6b and d). However, the actual $I_{\text{cell}}$ value (0.5-1 mA) was slightly lower than expected during the bulk electrocatalysis due to the inherent electrochemical polarization.

To further enhance the formate production, the coupling of EC-II and PV-II was considered ($E_{\text{cell}}$ ~2.8 V and $I_{\text{cell}}$ of 5.74 mA; solar conversion system efficiency of ~12%). Using this couple, the formate production was significantly enhanced ~4 times while maintaining an FE of ~95%. However, the As(V) production decreased due to the competitive oxidation of water, and hence, its FE was ~10%. This behavior was similar to that observed for the DC-powered systems. When chloride was added to the bicarbonate solutions, $E_{\text{cell}}$ decreased by 0.1 V, while $I_{\text{cell}}$ increased from 4 to 4.5 mA (Fig. S20). As a result, the formate production was slightly enhanced at the same FE of ~95%, while the As(V) production doubled, with an FE of ~22%.
due to the RCS-mediated oxidation of As(III).

Conclusions

This study demonstrated an off-grid hybrid electrocatalytic process for simultaneous As(III) oxidation and CO₂ reduction using a nanoparticulate TiO₂ anode and Bi cathode in aqueous bicarbonate solutions. Although the As(III) oxidation required a low potential bias for n-TEC, its Faradaic efficiency depended significantly on the applied potential, with a value of ~100% at Eₚ,As (~0.8 V) followed by a linear decrease with increasing potential. This dependency was attributed to specific interactions and charge transfer between free or complexed As(III) and TiO₂ at Eₚ,As. Potentials greater than the Eₚ,As gradually led to water oxidation, decreasing the Faradaic efficiency of As(V) production. However, in the presence of chloride, the chloride oxidation occurs competitively with water oxidation, effectively mediating the As(III) oxidation and enhancing the Faradaic efficiency. On the other hand, the CO₂ reduction reaction on Bi electrode occurred relatively efficiently as long as a sufficiently negative potential (< −1.5 V) was applied, leading to a Faradaic efficiency of >80% for formate production. In this cathodic reaction, chloride favorably enhanced the current and electron transfer to the Lewis-acidic carbon atom of CO₂. Although both the anodic or cathodic half-reaction could be optimized rather easily, driving both reactions in a single device at the same efficiencies as in the half-reactions was not straightforward due to the cell resistance and the different electrocatalytic activity of the anode and cathode. A well-designed device with a suitably matched photovoltaic was demonstrated to achieve both the anodic and cathodic reactions at high Faradaic efficiencies similar to those in the half-reactions. Despite being a
demonstration at the proof-of-concept level, this study should provide valuable information for designing platforms to realize two different redox reactions in a single device coupled with a photovoltaic.

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Author Contribution Statement

W. Choi: Methodology, Validation, Formal analysis, and Investigation

M. Kim: Methodology, Validation, Formal analysis, and Investigation

B.-j. Kim: Formal analysis

Y. Park: Resources, Funding acquisition

D.S. Han: Funding acquisition

M. R. Hoffmann: Resources

H. Park: Conceptualization, Methodology, Validation, Formal analysis, Resources, Writing – Original Draft, Writing – Review & Editing, Visualization, Supervision, Project administration, and Funding acquisition

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References


Scheme Caption

Scheme 1. Illustration of a PV-coupled electrocatalytic device for As(III) oxidation and CO2 reduction using an n-TEC anode and Bi cathode, respectively, in aqueous bicarbonate solution (0.1 M) purged with CO2. n-TEC represents nanoparticulate TiO2 films deposited on an IrxTa1-xOy-coated Ti substrate. In the absence of chloride, the oxidation of As(III) to As(V) and reduction of CO2 to HCOOH (formate) occurs via a direct charge transfer at high Faradaic efficiencies. In the presence of chloride, transiently formed reactive chlorine species (RCS) mediate the As(III) oxidation while CO2 reduction is facilitated due to the electron-donating ability of chloride. As3+ and As5+ represent H3AsO3 and H2AsO4− (or HAsO42−), respectively, at pH ∼6.8, while RCS include Cl2, ClO−, HClO, etc.

Figure Captions

Fig. 1. Surface characterization of as-synthesized n-TEC samples. (a) Cross-sectional EPMA images for Ti, O, Ir, and Ta elements. (b) Depth-profiled elemental distributions of Ti, O, Ir, and Ta. 0 depth refers to the top surface. (c) XRD pattern.

Fig. 2. Electrocatalytic oxidation of As(III) (1 mM) to As(V) using n-TEC electrodes (1×1 cm2, bifacial) in aqueous potassium bicarbonate (BC) solution (0.1 M) under various conditions. (a) Linear sweep voltammograms (LSVs) in the absence (i.e., BC only) and presence of As(III) (1 mM), Cl− (0.1 M), and/or CO2 (purged through the solution). (b-d) Bulk electrolysis of As(III) at various potentials: (b) As(V) production, (c) time-profile current, and (d) Faradaic efficiencies of As(V) production.

Fig. 3. Changes in the concentrations of HOCl, As(III), and As(V) in bicarbonate solutions (a) with and (b) without Cl− (0.1 M). Prior to spiking As(III) at 1 mM, the solutions underwent pre-electrolysis with n-TEC electrodes at 1.2 V for 2 h. Immediately after the electrolysis off and the addition of As(III), the concentrations of As(III) and As(V) were estimated for the following 1 h. During the pre- and post-electrolysis periods, HOCl was quantified as a representative of RCSs.

Fig. 4. (a) Cross-sectional SEM image of an as-synthesized Bi sample (electrodeposited for 30 s). (b) Particle size distribution of the Bi sample. Inset shows the EDS (left) and top-view SEM image (right). See Fig. S10 for more images. (c) XRD pattern of the Bi sample. Unassigned peaks refer to Ti substrate. Inset shows the magnified region between 36° and 42° (indicated
with an asterisk), where the Bi peaks (104 and 110) were shown. (d) Linear sweep voltammograms of as-synthesized Bi electrodes (electrodeposited for 30 s) in aqueous bicarbonate solutions (0.1 M) purged with CO₂. The 1st scan represents the voltammogram obtained immediately after synthesis; after the 1st scan, the 2nd potential scan was applied.

Fig. 5. (a) LSVs of Bi electrodes synthesized for 30 s in aqueous bicarbonate (BC) solutions (0.1 M) purged with CO₂ (or N₂ for comparison). If necessary, chloride (0.1 M) was added to the bicarbonate solutions. The inset compares the voltammograms of Bi electrodes in the CO₂-purged bicarbonate solutions as a function of electrodeposition time (0-60 s). (b) Effects of Bi-electrodeposition time (3-60 s) (upper panel) and applied potential (−1.4 to −1.8 V for Bi electrodes synthesized for 30 s) on the Faradaic efficiencies of formate production from CO₂-purged bicarbonate solutions in a single compartment reactor. In the upper panel, an E of −1.6 V was applied for CO₂ reduction. (c) Effect of chloride (0.1 M) on the production of formate and its Faradaic efficiency in CO₂-purged bicarbonate solutions (0.1 M) using Bi electrodes (30 s-sample) at E = −1.6 V in a two-compartment reactor divided by a proton-exchange membrane.

Fig. 6. (a) Electrocatalytic As(III) oxidation and simultaneous formate production using n-TEC and Bi electrodes (EC-I pair with area ratio of 32 at 2.5 V DC; EC-II pair with area ratio of 2 at 3.0 V DC), respectively, placed in a two-compartment cell (divided by a proton-exchange membrane) containing aqueous bicarbonate (0.1 M) solutions purged with CO₂. Note that CO₂ was purged into both the anolyte and catholyte to avoid pH difference between the electrolytes. If necessary, chloride (0.1 M) was added to the both anolyte and catholyte. (b) Faradaic efficiencies (blue and red bars for As(V) and formate, respectively). (c) Comparison of the LSVs of the n-TEC and Bi electrode pairs (area ratios of 32 and 2; denoted as EC-I and EC-II pairs, respectively) and I-V curves of the photovoltaics. The specifications of the PVs were as follows: PV-I: Area = 0.4 cm², J_{SC} = 6.784 mA·cm⁻², V_{OC} = 2.64 V, Fill factor: 69.5%, Efficiency = 12.42%. PV-II: Area = 1.32 cm², J_{SC} = 4.173 mA·cm⁻², V_{OC} = 3.65 V, Fill factor: 70.5%, Efficiency = 12.13%. (d) PV-powered simultaneous As(III) oxidation and formate production. With PV-II, chloride (0.1 M) was added to the solution to examine the chloride effect. The other experimental conditions were identical to those of Fig. 6a. See Fig. 6b for the Faradaic efficiencies.
Scheme 1. Illustration of a PV-coupled electrocatalytic device for As(III) oxidation and CO₂ reduction using an n-TEC anode and Bi cathode, respectively, in aqueous bicarbonate solution (0.1 M) purged with CO₂. n-TEC represents nanoparticulate TiO₂ films deposited on an IrₓTa₁-xO₉-coated Ti substrate. In the absence of chloride, oxidation of As(III) to As(V) and reduction of CO₂ to HCOOH (formate) occur via direct charge transfer at high Faradaic efficiencies. In the presence of chloride, transiently formed reactive chlorine species (RCS) mediate As(III) oxidation while CO₂ reduction is facilitated by the electron-donating ability of chloride. As³⁺ and As⁵⁺ represent H₃AsO₃ and H₂AsO₄⁻ (or HAsO₄²⁻), respectively, at pH ~6.8, while RCS include Cl₂, Cl₂⁻, and HClO.
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