Hybrid bioinorganic approach to solar-to-chemical conversion

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Natural photosynthesis harnesses solar energy to convert CO2 and water to value-added chemical products for sustaining life. We present a hybrid bioinorganic approach to solar-to-chemical conversion in which sustainable electrical and/or solar input drives production of hydrogen from water splitting using bio-compatible inorganic catalysts. The hydrogen is then used by living cells as a source of reducing equivalents for conversion of CO2 to the value-added chemical product methane. Using platinum or an earth-abundant substitute, α-NiS, as biocompatible hydrogen evolution reaction (HER) electrocatalysts and Methanosarcina barkeri as a biocatalyst for CO2 fixation, we demonstrate robust and efficient electrochemical CO2 to CH4 conversion at up to 86% overall Faradaic efficiency for >7 d. Introduction of indium phosphide photocathodes and titanium dioxide photoanodes affords a fully solar-driven system for methane generation from water and CO2, establishing that compatible inorganic and biological components can synergistically couple light-harvesting and catalytic functions for solar-to-chemical conversion.

artificial photosynthesis | solar fuels | photocatalysis | carbon dioxide fixation | water splitting

Methods for the sustainable conversion of carbon dioxide to value-added chemical products are of technological and societal importance (1–3). Elegant advances in traditional approaches to CO2 reduction driven by electrical and/or solar inputs using homogeneous (4–16), heterogeneous (17–26), and biological (27–31) catalysts point out key challenges in this area, namely (i) the chemoselective conversion of CO2 to a single product while minimizing the competitive reduction of protons to hydrogen, (ii) long-term stability under environmentally friendly aqueous conditions, and (iii) unassisted light-driven CO2 reduction that does not require external electrical bias and/or sacrificial chemical quenchers. Indeed, synthetic homogeneous and heterogeneous CO2 catalysts are often limited by product selectivity and/or aqueous compatibility, whereas enzymes show exquisite specificity but are generally less robust outside of their protective cellular environment. In addition, the conversion of electrolysytic systems to photosynthetic ones is nontrivial owing to the complexities of effectively integrating components of light capture with bond-making and bond-breaking chemistry.

Inspired by the process of natural photosynthesis in which light-harvesting, charge-transfer, and catalytic functions are integrated to achieve solar-driven CO2 fixation (32–35), we have initiated a program in solar-to-chemical conversion to harness the strengths inherent to both inorganic materials chemistry and biology (36). As shown in Fig. 1, our strategy to drive synthesis with sustainable electrical and/or solar energy input (37) interfaces a biocompatible photo( electro)chemical hydrogen evolution reaction (HER) catalyst with a microorganism that uses this sustainably derived hydrogen as an electron donor for CO2 reduction. Important previous reports have shown the feasibility of electrolysytic (38–42) but have not yet established solar-driven processes. We selected methane as an initial target for this approach owing to the ease of product separation, the potential for integration into existing infrastructures for the delivery and use of natural gas (of which CH4 is the principle component), and the fact that direct conversion of CO2 to CH4 with synthetic catalysts remains a formidable challenge due to large overpotentials and poor CH4/H2 selectivity. Two of the most active and selective direct electrocatalysts for CO2 to CH4 conversion reported to date produce methane with ≥ 61% (43) and ≥ 76% (44) Faradaic efficiencies, but require overpotentials of η = 1.28 V and η = 1.52 V, respectively. Promising advances in photothermal reduction of CO2 to CH4 also have been recently reported (45). In comparison with fully inorganic catalysts, a distinct conceptual advantage of this hybrid materials biology approach, where the materials component performs water splitting to generate hydrogen and the biological component uses these reducing equivalents for CO2 fixation, is that one can leverage the fact that biological catalysts operate at near thermodynamic potential (46). As such, the only overpotential involved is associated with hydrogen evolution from water, a more facile reaction to catalyze via sustainable electrochemical and photochemical means compared with CO2 reduction. Coupled with the diversity of potential chemical products available via synthetic biology, the marriage between artificial and natural platforms can create opportunities to develop catalyst systems with enhanced function over the individual parts in isolation.

In developing hybrid bioinorganic platforms for solar-to-chemical conversion of CO2, we drew inspiration from both

Significance

Natural photosynthesis, a process of solar-to-chemical conversion, uses light, water, and carbon dioxide to generate the chemical products needed to sustain life. Here we report a strategy inspired by photosynthesis in which compatible inorganic and biological components are used to transform light, water, and carbon dioxide to the value-added product methane. Specifically, this solar-to-chemical conversion platform integrates photo( electro)chemical hydrogen evolution reaction (HER) catalysts with microorganisms that consume this sustainably derived hydrogen to drive the transformation of carbon dioxide to methane with high efficiency. This system establishes a starting point for a broader materials biology approach to the synthesis of more complex chemical products from carbon dioxide and water.


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tandem organometallic–microbial systems (47, 48), in which products of microbial metabolism are further transformed by organometallic catalysts, as well as biological electrocatalysis, in which organisms accept reducing equivalents from an electrode either in the form of soluble electron carriers (for example, H₂ or formate) (41, 49, 50) or via direct electron transfer (36, 51–53). Engineered strains of Ralstonia eutropha have been used for the aerobic production of isobutanol and 3-methyl-1-butanol (41), and isopropanol (42). However, owing to the oxygen requirements of this organism and the relative inefficiency of its carbon fixation pathways (54), product titers and production efficiencies are relatively modest, and generation of reactive oxygen species is a serious concern. In addition, to our knowledge, no photosynthetic systems of this type have been reported. As such, we turned our attention to the use of a pure culture of Methanosarcina barkeri, an obligately anaerobic archaean that fuels its metabolism via the 8-proton, 8-electron reduction of CO₂ to CH₄ (55). Prior studies have reported methanogenic electrocatalysis (51, 53, 56); however, a fully light-driven system remains to be realized. Additionally, mixed cultures and multiple possible sources of reducing equivalents have complicated Faradaic efficiency measurements in previous studies (51, 53, 56). Through the design of our hybrid system, we sought to surmount some of these aforementioned challenges.

Here we report an integrated bioinorganic catalyst platform for solar-to-chemical CO₂ conversion using sustainable inorganic hydrogen generators in conjunction with CO₂-fixing archaea. Under electrosynthetic conditions with a platinum cathode, a culture of M. barkeri shows chemoselective conversion of CO₂ to CH₄ with high Faradaic efficiencies (up to 86%) and low overpotential (η = 360 mV). The system is also capable of high yield production, cumulatively generating 110 mL (4.3 mmol) of methane over 7 d. Isotope labeling with ¹³CO₂ establishes that CH₄ is uniquely derived from CO₂ for cultures in both rich media and minimal, carbon-free media. Replacement of Pt with an earth-abundant α-NiS electrocatalyst allows for CH₄ generation at similar titers. Moreover, using a photoactive silicon cathode reduces the overpotential to 175 mV upon irradiation with 740-nm light. Unassisted light-driven methane generation was achieved using tandem solar absorption by a photoactive n- TiO₂ anode and p-InP cathode assembly. Taken together, the results demonstrate the feasibility of combining compatible inorganic and biological systems to achieve solar-to-chemical conversion from light, H₂O, and CO₂, affording a starting point for the realization of sustainable fixation of CO₂ to value-added molecules.

Results and Discussion
Selection of Biological Catalyst. Careful organism selection is critical to the successful realization of an integrated bioinorganic system. The autotrophic obligate anaerobe M. barkeri (55) is amenable to integration with inorganic catalysts for a variety of reasons. M. barkeri can use H₂ as a source of reducing equivalents for the reduction of CO₂ to CH₄; the cathode of a water-splitting device could serve as a potential source of this H₂. Owing to the anaerobic metabolism of the organism, oxygen is not required at the cathode, thereby improving Faradaic efficiency for the product of interest, simplifying gas delivery to the culture, and preventing generation of potentially harmful reactive oxygen species. Furthermore, CH₄ is generated with high efficiency as a byproduct of normal metabolism. Finally, M. barkeri requires no added sources of reduced carbon and can produce CH₄ in minimal media containing only supplemental vitamins and minerals.

Electrochemical Reduction of Carbon Dioxide to Methane with a Hybrid Platinum/Archaean Catalyst Platform. Initial experiments were performed using a platinum cathode to electrochemically generate H₂, which was subsequently used in situ by M. barkeri to reduce CO₂ to CH₄. Fig. 24 shows a general schematic of the gas-tight, two-compartment electrochemical cell that was specially fabricated for batch-mode electrolysis and subsequent headspace analysis by gas chromatography (GC). (See Fig. S1 for electrolysis cell photo.) Separation of the cathodic and anodic chambers with an ion-permeable membrane prevented any noticeable diffusion of O₂ into the culture. After inoculation of the cathodic chamber with M. barkeri (130-mL final volume, OD₆₅₀ = 0.35) and saturation of the carbon-free catholyte with pure CO₂.

Fig. 1. General scheme depicting a hybrid bioinorganic approach to solar-to-chemical conversion. Sustainable energy inputs in the form of electrical potential or light can be used to generate hydrogen from water using inorganic HER catalysts; biological hydrogen-driven CO₂ fixation can subsequently generate value-added products such as methane. This materials biology interface can be generalized to other chemical intermediates and end products by mixing and matching different compatible inorganic and biological components.

Fig. 2. Electrocatalytic reduction of carbon dioxide to methane with a hybrid platform M. barkeri platform. (A) Schematic of generalized electrolytic setup showing in situ generation of hydrogen at the cathode followed by hydrogen-driven reduction of carbon dioxide to methane by the M. barkeri biocatalyst. (B) Cumulative methane generation and associated average Faradaic efficiency in minimal media. (C) Cumulative methane generation and associated average Faradaic efficiency at various currents in rich media. (D) Cumulative long-term methane generation and associated average Faradaic efficiency in rich media. (E and F) High-resolution mass spectrometry of headspace gases after electrolysis under an atmosphere of (E) ¹³CO₂ and (F) ¹²CO₂ in rich media. Error bars represent SD with n = 3 independent experiments in all cases.
galvanostatic electrolysis was performed at a current of 2.5 mA ($j = 0.29 \text{ mA/cm}^2$, $\eta = 360 \text{ mV}$) for 11.5 h, followed by 0.5 h of sampling. (See the Supporting Information for calculation of overpotentials.) The headspace was analyzed by GC and replaced by sparging with fresh CO$_2$ before restarting the electrolysis. (See Fig. S2 for GC calibration curves.) Methane production was linear under these conditions and cumulatively resulted in 16.8 ± 0.6 mL CH$_4$ (0.660 ± 0.024 mmol) over 3 d (all volumes reported in this work assume 1-atm pressure and a temperature of 310 K) with an average Faradaic efficiency of 81 ± 3% ($n = 3$). (All errors represent standard deviation unless otherwise noted.) (Fig. 2B).

Electrolysis was subsequently conducted in rich media containing yeast extract and casitone; variation of the applied current from 1 mA ($j = 0.12 \text{ mA/cm}^2$) to 7.5 mA ($j = 0.88 \text{ mA/cm}^2$) shows that CH$_4$ generation is proportional to applied current, suggesting that the system is operating in a hydrogen-limited regime (Fig. 2C). The highest Faradaic efficiency, 86 ± 3%, was observed with an applied current of 2.5 mA ($j = 0.29 \text{ mA/cm}^2$). A slight decrease in Faradaic efficiency was observed at higher current densities. We speculate that increased rates of HER cause H$_2$ to be less effectively delivered to the culture solution with loss to the headspace.

To test the limits of CH$_4$ production of the system, the organisms were electrolyzed at 7.5 mA for 7 d (Fig. 2D). The headspace of the cathodic chamber was sampled daily and subsequently exchanged with fresh CO$_2$. During each 24-h period (23 h electrolysis and 1 h sampling), 15.6 ± 0.7 mL CH$_4$ (0.613 ± 0.028 mmol) were produced; this value corresponds to cumulative methane production of 109 ± 5 mL (4.28 ± 0.20 mmol) over 7 d at an average Faradaic efficiency of 75 ± 4% ($n = 3$). Moreover, the hybrid bioorganic system showed no loss in efficiency over the course of the experiments, which were terminated due to time constraints rather than a decrease in performance. Taken together, these data establish that there are no viability concerns on the timescale tested and presage the possibility of extended operation with a single inoculation of biomass.

A series of isotopic labeling experiments was conducted to show that the observed methane was derived from carbon dioxide. Electrolysis cells containing nitrogen-sparged rich media were inoculated with *M. barkeri*. The headspace of one cell was filled with $^{13}$C$_2$O$_2$ and another with $^{12}$CO$_2$ and both were subjected to galvanostatic electrolysis at a current of 2.5 mA. High-resolution GC-mass spectrometry was used to show that only $^{13}$CH$_4$ was observed when electrolyzed under an atmosphere of $^{12}$CO$_2$ (Fig. 2F), whereas only $^{12}$CH$_4$ was observed in the $^{13}$CO$_2$ control experiment (Fig. 2E). Based on the limit of detection and amount of methane generated, <4% of the observed methane could potentially come from sources other than CO$_2$. Similarly, only $^{13}$CH$_4$ was observed when the electrolysis was performed in minimal media (Fig. S3B). Based on the limit of detection and amount of methane generated, <7.5% of the observed methane could potentially come from sources other than CO$_2$ under these conditions.

**Synthesis and Characterization of $\alpha$-NiS HER Catalyst and Its Use in Carbon Dioxide to Methane Conversion.** After demonstrating CO$_2$-to-CH$_4$ conversion by *M. barkeri* during in situ electrolysis using a platinum HER cathode, we sought to replace the precious metal catalyst with an earth-abundant alternative. Several HER catalysts containing only first-row transition metals have been recently reported (57–60); among these examples, metal chalcogenides have featured prominently (61–67). In addition to low overpotential and long-term stability, another requirement for a catalyst in a biomaterials hybrid system is that it must operate in aqueous media within a biologically relevant pH range (pH 5–8) and be nontoxic to the organism. Relatively few published examples meet all of these criteria. As such, we developed nanoparticulate $\alpha$-NiS as a biocompatible HER catalyst for integration with *M. barkeri* cultures.

Nanoparticulate nickel sulfide was prepared by microwave irradiation (250 °C, 30 min) of an aqueous solution of nickel chloride, thiourea, and ammonium hydroxide (Fig. 3A). The resulting black powder was rinsed with water and isopropanol and dried under vacuum overnight. The newly synthesized catalyst was stored in a capped vial in air and retained activity for several months. Characterization by powder X-ray diffraction (pXRD) confirmed that the product is crystalline and identified the primary phase as $\alpha$-NiS (Joint Committee on Powder Diffraction Standards number 77–1624) (Fig. 3B). Transmission electron microscopy (TEM) images show polydisperse hexagonal particles of 20–100 nm diameter (Fig. S4). The nickel-to-sulfur ratio was verified to be 1:1 by energy-dispersive X-ray (EDX) spectroscopy based on the nickel L and sulfur K peaks (Fig. 3C), a result which was further corroborated by inductively coupled plasma-optical emission spectroscopy (ICP-OES) measurements. High-resolution TEM images taken along the [010] axis confirm the assignment of single-crystalline $\alpha$-NiS based on the (100) and (001) lattice planes (Fig. 3D). Thermogravimetric analysis (TGA) shows negligible loss of mass until 600 °C, suggesting the absence of surface-bound organics (Fig. S5).

Electrocatalytic properties of the $\alpha$-NiS particles were evaluated using rotating disk electrochemistry and chronopotentiometry. As shown in the polarization curve (Fig. 3E), crystalline $\alpha$-NiS is an active HER catalyst in pH 7 potassium phosphate buffer and achieves current densities of 1 and 5 mA/cm$^2$ at overpotentials of $\eta = 275$ and 350 mV, respectively. The observed Tafel slope of 111 mV/dec is similar to the predicted 118 mV/dec for a process in which the Volmer step (formation of an adsorbed H intermediate) is rate-determining (68). The exchange current density, $j_0$, is 3.5 × 10$^{-2}$ mA/cm$^2$ based on extrapolation of the Tafel plot (Fig. 3F). These parameters compare favorably to other heterogeneous, first-row transition metal HER catalysts, especially given the ease of synthesis (Table S1).

A suspension of $\alpha$-NiS powder in 3:1 ethanol:water with Nafion binder was deposited on conductive carbon cloth to
fabricate larger-scale electrodes for hydrogen generation and long-term stability measurements. Galvanostatic experiments at a current of 2.5 mA were run for 23 h; the Faradaic efficiency for hydrogen generation was found to be 95 ± 4% (n = 3) (Fig. S6). As shown in Fig. S7B, the potential applied during these experiments does not significantly decay over time, indicating that the catalyst is stable under such conditions.

Having established the efficacy and stability of α-NiS as a catalyst for HER at biologically relevant pH, we sought to interface it with a *M. barkeri* culture for electrochemical methane production. A piece of conductive carbon cloth with 1.4 mg deposited α-NiS catalyst was used as an earth-abundant replacement for platinum. In a galvanostatic experiment at 2.5 mA (i_{avg} = 0.28 mA/cm², n = 695 mV), 5.1 ± 0.02 mL CH₄ (0.20 ± 0.01 mmol) were generated in 24 h, cumulatively generating 15.4 ± 0.8 mL CH₄ (0.605 ± 0.031 mmol) over 3 d with an average Faradaic efficiency of 73 ± 5% (n = 3) (Fig. S8). These values are comparable to those obtained for the corresponding platinum experiments. When the current was increased to 7.5 mA (i_{avg} = 0.83 mA/cm²), experiments using α-NiS/C were virtually indistinguishable from Pt in terms of daily and cumulative methane, producing 108 ± 4 mL CH₄ (4.24 ± 0.16 mmol) over 7 d with an average Faradaic efficiency of 74 ± 2% (n = 3) (Fig. 3G).

**Photoelectrochemical Generation of Methane from Carbon Dioxide.** Having performed electrochemical conversion of CO₂ to CH₄, we next sought to develop a photoelectrochemical system in which a portion of the potential required for water splitting is contributed by light. Indeed, transforming electrochemical systems to photochemical ones remains challenging. To achieve this goal, we sought to use semiconductor photocathodes coated with a thin-film HER catalyst. The overall performance of such an assembly is determined by a tradeoff between efficient catalysis and light absorption: thicker films confer superior electrocatalytic activity and stability, whereas thinner and more transparent films allow for greater photon capture. Although α-NiS proved effective as an earth-abundant HER electrocatalyst, previously published work from our laboratories on the electrodeposition of a related cobalt sulfide film on planar n+/p-Si showed that the current density under illumination drastically diminished as the thickness of the film increased (62). For this reason, we chose to use a nickel–molybdenum alloy, a previously characterized earth-abundant HER catalyst, which has shown favorable performance under photocatalytic conditions despite having a slightly larger overpotential than platinum (69). Photoactive cathodes were prepared by sputtering a thin layer of Ni-Mo alloy atop TiO₂-sputtered n+/p-Si (Fig. 4A). To confirm hydrogen evolution, abiotic galvanostatic experiments at a current of 2.5 mA (i_{avg} = 0.36 mA/cm²) were run for 23 h while illuminating the cathode with 740-nm light (20 mW/cm²); Faradaic efficiencies for H₂ were 103 ± 3% (Fig. S6). A light toxicity control experiment with *M. barkeri* showed that methane generation is not affected by illumination at this wavelength (Fig. S9). Biological galvanostatic electrolysis at 2.5 mA with the photocathode (740-nm illumination) generated 17.6 ± 2.1 mL CH₄ (0.692 ± 0.083 mmol) with a Faradaic efficiency of 82 ± 10% (n = 3) with only 175-mV overpotential (Fig. 4B).

**Unassisted Light-Driven Synthesis of Methane from Carbon Dioxide.** Finally, we sought to construct a fully light-driven hybrid bio-inorganic system for CO₂-to-CH₄ conversion through the use of a tandem semiconductor assembly. In this setup, full-spectrum light first impinges on a large bandgap anode [nanowire n-ITO on fluorine-doped tin oxide (FTO)], where water oxidation generates oxygen (37). The filtered, lower-energy light subsequently illuminates a smaller bandgap cathode (p-InP coated with Pt), where water reduction generates hydrogen (Fig. 4C). The photoelectrochemical cell design described above required slight modifications for unassisted photochemical experiments: a 1-mm diameter quartz window was added to the anodic chamber to prevent initial filtering of the full-spectrum light by glass, and the anode and cathode compartments were separated by an anion exchange membrane to minimize pH changes. This linked two-electrode assembly generates nonzero photocurrent under illumination (i_{avg} = 0.17 mA, i_{avg} = 0.057 mA/cm² during the first hour) (Fig. S7D). The Faradaic efficiency for hydrogen generation in abiotic experiments is 100 ± 8% (Fig. S6 (n = 4)). Before introduction of *M. barkeri* into such a system, galvanostatic control electrolyses of a *M. barkeri* culture were performed using Pt electrodes under illuminated and nonilluminated conditions. When only the n-ITO/FTO anode was used as a light filter, lower than expected Faradaic efficiencies for methane were observed in the illuminated experiment (Fig. S9B), this is in agreement with literature concerning the photosensitivity of methanogenic archaea to blue light (70, 71). Installation of a 455-nm filter directly after the photoanode restored the Faradaic efficiency for methane to expected levels (Fig. S9B). Experiments to produce methane using this hybrid bioinorganic system were conducted similarly to those described earlier except headspace analysis was performed only once per experiment after 3 d. On average, 1.75 mL CH₄ (68.8 nmol) (n = 2) were produced in illuminated experiments, whereas only 0.58 mL CH₄ (22.8 nmol) (n = 3) were produced in identical nonilluminated controls (Fig. 4D). We note that higher background levels of methane production were observed in these unassisted photochemical experiments compared with the electrochemical and photoelectrochemical experiments and speculate that this may be due to biological corrosion of the multipurpose photocathode, a phenomenon previously documented in other systems (53, 72). Subtraction of the background methane results in an average Faradaic efficiency of 74%. This result clearly demonstrates the successful conversion of CO₂ to CH₄ using light as the sole energy input.

**Concluding Remarks**

In summary, we have established a hybrid bioinorganic approach to solar-to-chemical conversion by transforming carbon dioxide and water to the value-added chemical product methane. The present integrated system couples inorganic hydrogen generation
catalysts powered by sustainable electrical and/or solar energy inputs with a biological catalyst that can harvest reducing equivalents from H₂ to convert carbon dioxide to methane with up to 86% Faradaic efficiency. The data show that the low aqueous solubility and mass transfer rate of hydrogen do not preclude it from being an effective molecular redox carrier and offer a starting platform for potential integration with sustainable sources of electricity (e.g., solar, wind, hydrothermal, nuclear, etc.). Indeed, a solar-to-chemical efficiency of 10% and an electrical-to-chemical efficiency of 52%, which compare favorably to previously reported systems (73), are possible given ca- vities of materials and biological integration, assuming efficiency values of 20% for solar-to-electrical conversion using a commercially available photovoltaic and 70% for electrical-to- hydrogen conversion in an optimized system (74), as well as 86% energetic efficiency for H₂ to CH₄ based on thermodynamic values. Furthermore, we have developed an earth-abundant nanoparticulate nickel sulfide HER catalyst that operates effectively under biologically compatible conditions. At an applied current of 7.5 mA, use of the α-NiS catalyst results in 110 mL (4.3 mmol) of methane over 7 d with a Faradaic efficiency of 74%, comparable to Pt. Moreover, we have demonstrated that use of a photoactive silicon cathode results in an overpotential of only 175 mV, allowing for greater methane generation from a defined quantity of electrical energy. Finally, combining an n-TiO₂ photoanode with a p-InP photocathode allows for fully unassisted light-driven conversion of CO₂ to CH₄. By realizing an effective artificial photosynthesis platform for the production of a value-added chemical product from light, CO₂, and water, these results provide a starting point for achieving sustainable chemistry with materials biology.

Materials and Methods

Materials. Reagents were purchased from commercial sources as noted in the Supporting Information and used without further purification. Ultrahigh-purity gases purchased from Praxair were used for all anaerobic manipulations. 13C-labeled CO₂ was purchased from Cambridge Isotopes. Distilled water (dH₂O) was deionized to a resistivity of 18.2 MΩ cm using a Millipore Milli-Q UF Plus system (dH₂O).

Strain Details and Culture Conditions. M. barkeri (ATCC 43241) was purchased from the American Type Culture Collection (ATCC). All culture manipulations were performed inside a Vacuum Atmospheres Nexus One glovebox with an atmosphere of 90% nitrogen and 10% hydrogen. Oxygen levels and humidity levels within the box were controlled using a STAK-PAK palladium catalyst and desiccant system in a Coy Laboratory Products unheated fan box. M. barkeri primary cultures were propagated in 18 × 150-mm Balb tubes with butyl rubber stoppers and aluminum crimp seals using ATCC medium 1043 with deoxygenated methanol (1% vol/vol) as the growth medium. Secondary cultures were started by diluting late exponential phase primary cultures 1:100 into 1 L of ATCC medium 1043 in a 2-L anaerobic media bottle with deoxygenated methanol (1% vol/vol) as the growth substrate. All cultures were grown at 37 °C with shaking (200 rpm). The primary culture was propagated by diluting 1:100 into fresh ATCC 1043 medium every 4 d for 3–4 mo, or until changes in growth patterns were observed. Glyceral stocks were used for long-term culture storage.

General Experimental Setup. A. barkeri secondary culture was harvested after 4 d of growth by centrifugation at 6,000 x g for 7 min at 4 °C using O-ring–sealed centrifuge bottles. After centrifugation, the pellets were combined and washed twice with methanol-free media (200 mL). The final washed pellet was resuspended in fresh methanol-free media (20 mL). At this point, the OD₆₀₀ nm of the culture was determined by diluting the cell suspension 1:100 into 10 mL of water (pH 8.5).

The electrolysis cells were assembled inside an anaerobic glovebox. The appropriate media was added to the cathodic (130 mL) and anodic (70 mL) compartments as specified in the Supporting Information. Additionally, a magnetic stir bar was added to the cathodic compartment. The resuspended cell pellet was used to inoculate the cathodic side of the electrolysis cell to a final OD₆₀₀ nm of 0.35. Appropriate electrodes were added to the cathodic and anodic compartments, and the tightly sealed electrolysis cells were removed from the anaerobic chamber. Once outside, the cathodic compartment was sparged with CO₂ for 5 min using the internal sparge line. Helium (1 mL) was injected into the cathodic headspace as a gaseous internal standard. The electrolysis cells were submerged to the horizontal flanges in water baths set to 37 °C. The cultures were stirred at 300 rpm for the duration of the experiment. In experiments with 24-h (12-h) timepoints, the experimental interval consisted of 23 h (11.5 h) of electrolysis followed by a 1-h (0.5-h) period for headspace analysis and sparging. GC analysis was performed at each timepoint by direct introduction of the headspace into a GC sampling loop as described in the Supporting Information. Immediately after sampling, the headspace of the cathodic chamber was exchanged by sparging with fresh CO₂ for 5 min via the internal sparge line, followed by injection of a (1 mL) internal standard.

Synthesis of α-NiS Catalyst. Freshly prepared solutions of NiCl₂·6H₂O (0.2 M), thioacetamide (0.4 M), and 5% (vol/vol) ammonium hydroxide were prepared using dH₂O. Nickel chloride solution (2 mL) and thioacetamide solution (2 mL) were combined in a 10-mL microwave reaction vial and sparged with N₂ for 10 min. Upon addition of ammonium hydroxide solution (40 mL), a small amount of yellow precipitate was observed. The reaction tube was then capped under N₂ and maintained at 30 min at 150 °C with a 5-min temperature ramp using a CEM microwave reactor. Following the reaction, the supernatant was decanted and the black particles were washed three times with dH₂O and once with isopropanol by centrifuging the sample, decanting the supernatant, and resuspending the particles. Drying under vacuum overnight yielded a fine black powder (22 mg, 61% yield). The product was characterized by rotating frame nuclear Overhauser effect (RNOE) and X-ray diffraction (XRD) experiments, pXRD, EDX, TEM, high-resolution TEM (HRTEM), ICP-OES, and TGA as described in the Supporting Information. The catalyst was stored in air for 3–4 mo, after which time it was found to be no longer active for HER.

Electrochemical Details. All electrochemical experiments were performed using a BASI Epsilon potentiostat. Pt gauze (electrochemical grade, purchased from Alfa Aesar or Sigma-Aldrich) was used as the anode in all experiments except the unassisted full-splitting, where an n-TiO₂-coated FTO anode was used instead. The reference electrode was an aqueous Ag/AgCl electrode (3.5 M KCl) purchased from BASI. All reference electrodes were externally calibrated to potassium ferrocyanide in pH 7 phosphate buffer (Fe₃⁺/Fe²⁺ couple is 0.436 V vs. standard hydrogen electrode) before and after each experiment. No iR compensation was applied.

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