Nickel-Borate Oxygen-Evolving Catalyst that Functions under Benign Conditions

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Nickel-borate oxygen-evolving catalyst that functions under benign conditions

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Contributed by Daniel George Nocera, April 8, 2010 (sent for review December 26, 2009)

Thin catalyst films with electrocatalytic water oxidation properties similar to those of a recently reported Co-based catalyst can be electrodeposited from dilute Ni2+ solutions in borate electrolyte at pH 9.2 (B). The Ni-B films can be prepared with precise thickness control and operate at modest overpotential providing an alternative to the Co catalyst for applications in solar energy conversion.

Global energy need will roughly double by midcentury and triple by 2100. Most of that demand is driven by 3 billion low-energy users in the nonlegacy world and by 3 billion people yet to inhabit the planet over the next half century (1). Research targets that meet the energy need of these 6 billion new energy users provide global society with a direct path to providing a solution for a sustainable and carbon-neutral energy future. Moreover, because energy use scales with wealth (2), geopolitical and economic stability is greatly enhanced by abundant and clean energy as it enables individuals in the smallest village in the nonlegacy world and in the largest city of the legacy world to coexist more equitably. However, the task of supplying energy to the nonlegacy world will not be an easy one. To achieve the target of a carbon-neutral energy supply for the nonlegacy world will require the delivery of energy on a scale commensurate with, or larger than, the entire present-day energy supply from all sources combined (3).

The success in meeting the energy challenge in the nonlegacy world will largely depend on the design, research, and development of new technologies that are at odds with energy systems of the legacy world. Energy systems of the past and present operate at large scale, they are centralized, and energy is distributed to the masses. Such infrastructure is not compatible to the near-term needs of the nonlegacy world where it is cost prohibitive to build centralized energy and distribution systems. An alternative strategy that is better adapted to making energy available to the 6 billion new energy users is highly distributed energy systems for the individual—a personalized energy (PE) (4, 5)—that places a premium on low cost, even if efficiency must be sacrificed up front. Low cost in a manufacturing environment is most profoundly affected by materials goods of the system (most generalized by the weight of the system) and the production volume (6, 7). Here again, the nonlegacy and legacy worlds diverge. Large, centralized, and efficient energy systems come with significant balance-of-system (BOS) costs. Down-scaling such technology does not scale economically because the BOS costs do not scale commensurately. Thus, off-the-shelf technologies will be difficult to adapt to low-cost energy systems. Rather, the disruptive energy technologies of the future will be those that conform to the message of lightweight and high-volume manufacturing. Simply put, new research and development are needed to provide the nonlegacy world with the “fast food” equivalent of energy systems.

Solar PE is particularly well adapted to meeting the energy needs of the nonlegacy world. It is highly distributed and available to all. And low-cost and large-scale manufacturing are already an emerging trend in the deployment of photovoltaics (8). However, low-cost photovoltaics are not enough. Because local insolation is diurnal, solar energy cannot be used as a large-scale energy supply unless it can be stored. Most current methods of solar storage are characterized by low-energy densities and consequently they present formidable challenges for large-scale solar implementation. This includes batteries, which have low-energy density (0.5 MJ/kg with a ceiling of ~3.0 MJ/kg). Conversely, the energy density of fuels is ≥102 larger and H2 is ≥103 larger than the upper limit of battery storage capacity. Indeed, society has intuitively understood this disparity in energy density as all large-scale energy storage is in the form of fuels. But these fuels are carbon based. An imperative for our research has been to develop fuel storage methods that are carbon-neutral, sustainable, easily scalable, and inexpensive.

We have taken important steps towards meeting this imperative by developing a cobalt oxygen-evolving catalyst (Co-OEC) (9, 10) that can use an energy input from a photoanode or photovoltaic to split water into hydrogen and oxygen. Co-OEC is unique because it (i) is a functional model of the oxygen-evolving complex of Photosystem II (11), (ii) operates safely with high activity under benign conditions (room temperature and pH 7) (9, 10), (iii) is comprised of inexpensive, Earth-abundant materials and is easy to manufacture (9, 10), (iv) is self-healing (12), (v) is functional in natural, waste, and salt water (10, 11), (vi) can form on diverse conducting surfaces of varying geometry and be easily interfaced with a variety of light-absorbing and charge-separating materials (11), and (vii) may be activated by solar-derived electricity or directly by sunlight mediated by a semiconductor (13, 14). The simple operation of the catalyst from impure water sets a path to providing distributed solar and clean drinking water (5).

We now report that thin catalyst films with O2 evolution properties can be electrodeposited from dilute Ni2+ solutions in borate electrolyte at pH 9.2 (B). The Ni-B films can be prepared with precise thickness control and operate at modest overpotential providing an alternative to the Co-OEC catalyst for applications in solar energy storage and water purification.

Results

Fig. 1 shows the onset of a catalytic wave at 1.2 V [vs, normal hydrogen electrode (NHE)] on the first anodic sweep of a glassy carbon electrode immersed in a 0.1 M H3BO3/H3BO3 electrolyte (B, electrolyte) solution at pH 9.2 containing 1 mM of Ni2+ ion. The cathodic return scan exhibits a broad feature at Ep,c = 0.87 V, attributed to the reduction of a surface adsorbed species formed during the initial sweep through the catalytic wave. Subsequent cyclic voltammetry (CV) scans display a sharp
served in the absence of the Bi electrolyte. Indeed, a CV of a electrodeposition. Neither film formation nor catalysis is ob-
duced after 20 scans, thus attesting to the controlled nature of the CV scan at estimate that a monolayer of catalyst is deposited after a single amplitude with scanning, suggesting the growth of a surface-
in Fig. 1, the anodic and the cathodic prefeatures increase in aqueous 1 mM Ni
1 mM aqueous solution of Ni
2+ Ni(OH)2/Ni(III)O(OH) (23–26). To examine if proton-coupled electron transfer is operational in our system, (27, 28) we measured the pH dependence of the anodic prefeature (see Fig. 2). Surprisingly, CV experiments conducted at a scan rate of 50 mV/s give rise to a plot of E
p,a vs. pH with a slope of –96 mV/pH unit. Similar slopes, –91 and –89 mV/pH unit, were obtained at scan rates of 10 and 5 mV/s, respectively (Fig. S2). These values are much higher than the theoretical value of –59 mV expected for the 1e–/1H+ oxidation of Ni(OH)2 to Ni(III)O(OH). Rather, a slope of –90 mV/pH unit corresponds to the loss of 1e– accompanied by the transfer of ~1.5 protons. This suggests that a 2e–/3H+ couple may be operative, as has been proposed for the dimerization of Ir(V) hydro-hydroxo species to form iridium oxide: 2[IrO2(OH)2]2− + 2e– + 3H2O ↔ [Ir2O3(OH)3]2+ + 2H2O3+ + 3OH– (29, 30), which exhibits an observed slope of ~88 mV/pH unit.

To further probe the morphology, composition, and structure of the electrodeposited material, thicker films of Ni-Bi were electodeposited onto indium-tin oxide (ITO) covered glass slides by subjecting 1 mM Ni2+ solutions in Bi electrolyte to bulk electrolysis at 1.3 V. During electrodeposition, the current density rapidly attains a plateau value of ~1.4 mA/cm2, as shown in Fig. 3. A 3–4-μm-thick film is grown by passing 10 C/cm2 at 1.3 V. Scanning electron microscopy and powder X-ray diffraction of this film reveal that it is relatively smooth and amorphous (see Fig. 3 Insert and Figs. S3 and S4, respectively). To interrogate the catalyst composition, thick films were grown on large surface area electrodes, and then dried in the air. Elemental analysis of the dried material revealed a formula of Ni2O3(OH)2/1(H2BO3)1/1 • 1.5H2O, which we understand may differ from that of the catalyst under wet and operational conditions. Nevertheless, the formula of Ni-Bi is reminiscent of NiO(OH), which exhibits O2 evolution under similar pH conditions and is an effective water oxidation catalyst in concentrated alkaline solution (24).

The potential-dependent O2 evolution activity of Ni-Bi, evaluated in Ni2+-free Bi electrolyte at pH 9.2, is in line with that observed for NiO(OH) under highly alkaline conditions. Faradaic efficiency for O2 evolution, as measured by fluorescence detec-

**Fig. 1.** CVs using a glassy carbon working electrode, 50 mV/s scan rate, of aqueous 1 mM Ni2+ in 0.1 M B, electrolyte, pH 9.2. Successive scans show an increase of the anodic peak current for the redox event centered at ~1.0 e. (Inset) First (—) and second (--) CV scans using a glassy carbon working electrode, 50 mV/s scan rate, of aqueous 1 mM Ni2+ solutions in 0.1 M B, electrolyte, pH 9.2. CV trace in the absence of Ni2+ is shown as a black trace.

**Fig. 2.** CVs scans of a Ni borate catalyst film at pH values of 7.82, 8.30, 8.68, 8.88, 9.16, 9.42, 9.72, and 9.98 in. going from left to right, respectively. The inset shows a linear fit for the 8.7–10.0 pH region, with a slope of 96 mV/pH unit. The CVs were taken at scan rates of 50 mV/s.

**Fig. 3.** Current density trace for bulk electrolysis at 1.30 V in 0.1 M B2O3, electrolyte, pH 9.2, 1 mM Ni2+ using an ITO anode. Irregularities are due to bubble formation and to stirring. The inset shows an SEM photograph of a film obtained by passing 10 C/cm2 at 1.30 V.
tion, is quantitative (Fig. S5). The current density for O₂ evolution, j, obtained for a thin film grown by passing 300 mC/cm² was measured as a function of the overpotential, η. A plot of log(j) vs. η (Tafel plot, Fig. 4) produces a slope of ~59 mV/decade for current densities ranging from 1 mA/cm² to 7 μA/cm².

Notably, preconditioning the electrode by subjecting it to 12 h of preelectrolysis at a current density of ~1 mA/cm² is necessary to obtain reproducible values of the Tafel slope. Indeed, measurements performed on fresh Ni-Bi films without preconditioning produced Tafel slopes of up to 120 mV/decade, outside the range of 40–80 mV/decade reported for typical Ni oxides (31, 32).

Very thin films of Ni-Bi are stable to cathodic bias, as demonstrated by cycling an electrodeposited film through 100 CV scans with a switching potential below the catalytic wave (1.1 V). Catalysis was still operative after these cycles, and the peak current of the anodic prefeature drops by only ~16% during the cycling experiment (Fig. S6). The stability of the Ni-Bi film to a cathodic bias on the CV timescale suggests that this material may also exhibit activity for O₂ reduction, a possibility that is being explored currently.

In summary, a Ni-based oxide catalyst can be deposited in a highly controllable fashion under mild conditions from dilute NiO₂ solutions in the presence of borate. Deposition routes mimicking those employed for the formation of a related Co OEC give rise to a Ni-based catalyst that evolves O₂ in near-neutral conditions with an activity of 1 mA/cm² at an overpotential of ~425 mV. The catalyst exhibits long-term stability in water with no observed corrosion. The results reported here contribute to an expanding group of water splitting catalysts that operate under mild conditions and at current densities commensurate with non-concentrated solar photovoltaic outputs (5–20 mA/cm²). The simple operating conditions may permit energy storage to be performed with devices that are inexpensive and highly manufacturable.

**Experimental Methods**

**Materials.** Ni(NO₃)₂ • 6H₂O 99.9985% was used as received from Strem. H₂BO₃ 99.5%, NaNO₃, and KOH 85% were used as received from Sigma-Aldrich. NaOH 99% was used as received from Mallinckrodt. All buffers were prepared from reagent-grade water (Rica Chemical, 18 MΩ-cm resistivity). ITO-coated glass slides (8–12 Ω/sq surface resistivity) and fluorine-tin-oxide (FTO) coated glass slides were purchased from Sigma-Aldrich and from Hartford Glass, respectively.

**Electrochemical Methods.** All electrochemical experiments were performed at ambient temperature in air-saturated solutions with a CH Instruments 760C potentiostat and a BASI Ag/AgCl reference electrode. All electrode potentials were converted to the NHE scale using E(NHE) = E(Ag/AgCl) + 0.197 V. Unless otherwise stated, the electrolyte was 0.1 M potassium borate at pH 9.2 (B₃ electrolyte).

**Cyclic Voltammetry.** A 0.07 cm² glassy carbon button electrode was used as the working electrode and Pt wire as the auxiliary electrode. The working electrode was polished with 0.3 and 0.05 μm alumina particles for 30 s each and thoroughly rinsed with reagent-grade water prior to use. Unless otherwise noted, cyclic voltammograms (CVs) were collected with compensated cell resistance (iR) at 50 mV/s and 0.01 or 0.1 mA/V sensitivity in Ni-free B₃ electrolyte, and B₃ electrolytes containing 1 mM NiO₂⁻.

To determine the surface catalyst coverage on a glassy carbon electrode, the integrated charge of the anodic prefeature was taken as a direct estimate of the number of Ni atoms electrodeposited. Monolayer coverage was approximated as a surface density of 5Ni atoms/25 Å² implying that an integrated charge of ~300 μC/cm² corresponds to one monolayer of catalyst.

To verify the necessity of using an electrolyte with good proton-accepting ability, a CV was taken with a glassy carbon electrode of a 0.1 M NaNO₃ solution whose pH was adjusted to 9.2 using concentrated KOH. Upon addition of 1 mM NiO₂⁻ to this solution, the CV trace was identical to that of the background, indicating that the presence of a proton-accepting electrolyte (such as borate) is essential for catalyst formation and activity (Fig. S1).

To determine the dependence of the anodic prefeature with pH, a catalyst film was prepared on a 0.07 cm² glassy carbon electrode by running 13–15 consecutive CV cycles in a 1 mM NiO₂⁻/0.1 M B₃ solution with a switching potential of 1.4 V. The working electrode was then thoroughly rinsed with reagent-grade water and placed in a fresh Ni-free B₃ solution of 0.1 M. This solution had an initial pH of 7.82. The pH was gradually increased by adding small aliquots of KOH solution, and CVs were recorded at each pH point with a scan rate of 50 mV/s. CVs at all pH points were taken with iR compensation using solution resistance values measured prior to each CV.

To verify the stability of a thin catalyst film, a polished electrode was used to record a CV in 1 mM NiO₂⁻ solution with a switching potential of 1.5 V. After the first full scan, the electrode was removed, rinsed with reagent-grade water, and placed into a Ni-free B₃ solution. One hundred consecutive CV scans were recorded with a switching potential of 1.1 V. An insignificant decrease in current density was observed for the anodic prefeature centered at ~1 V (Fig. S6). To verify that a competent catalyst was still present on the electrode after this treatment, the same electrode was cycled past the onset potential (>1.1 V); a large catalytic wave was observed, attesting that the catalyst film is stable to reductive bias on the CV timescale.

**Bulk Electrolysis and In Situ Catalyst Formation.** Bulk electrolyses were performed in a two-compartment electrochemical cell with a glass frit junction of fine porosity. For catalyst electrodeposition, the auxiliary compartment was charged with ~40 mL of 0.1 M B₃ electrolyte and the working compartment was charged with 40 mL of solution obtained by combining 20 mL of 0.2 M B₃ electrolyte and 20 mL of 2 mM NiO₂⁻ solution. The B₃ electrolyte solution became turbid immediately following the addition of NiO₂⁻ solution [likely due to the formation of Ni(OH)₂]. The solution was filtered through a 0.45 μm filter (Pall Acrodisc) prior to electrolysis. The filtered Ni-B₃ solution remained clear for several hours and during the entire duration of electrodeposition experi-
ments. The working electrode was a 1 × 2.5 cm piece of ITO-coated glass cut from a commercially available slide. Typically, a 1 cm² area of the electrode was exposed to the solution. Pt mesh was used as the auxiliary electrode. Electrolysis was carried out at 1.30 V with stirring and without iR compensation and with the reference electrode placed 2–3 mm from the ITO surface.

**Tafel Plot Data Collection.** Current-potential data were obtained by performing controlled potential electrolyses in B₃ electrolytes at a variety of applied potentials in two-compartment cells containing ~40 mL of fresh electrolyte on each side. Prior to data collection, the solution resistance was measured with a clean ITO working electrode using the iR test function. The working ITO electrode and the Ag/AgCl reference electrode were kept in the same configuration while the electrolysis cell was replaced with one containing Ni⁴⁺ in the working compartment, as described above. A 1 cm² catalyst film was prepared via an electrodeposition that passed 300 mC/cm². Working and reference electrodes were subsequently rinsed in reagent-grade water and all electrodes were reimmersed in a two-compartment electrochemical cell containing fresh B₃ electrolyte in both compartments. A current of 950 μA/cm² was applied for ~12 h prior to collecting data for the Tafel plot. Steady-state currents were measured at a variety of applied potentials while the solution was stirred, proceeding in 10–25 mV steps between 1.15 and 0.99 V. Typically, the current reached a steady-state at a particular potential in 2–5 min and current values were read after 7–10 min. Measurements were made twice. The variation in steady-state current between two runs at a particular potential was <5%. The solution resistance measured prior to the data collection (36 Ω) was used to correct the Tafel plot for iR drop (Fig. 4).

**Elemental Analyses.** Microanalyses were performed by Columbia Analytics. The Ni-Bi catalyst was prepared on large surface area (~25 × 25 cm) FTO-coated glass slides using filtered 1 mM Ni²⁺/0.1 M B₃ solutions. Upon termination of the electrolyses, the slides were immediately removed from the solution, rinsed with reagent-grade water, and allowed to dry in air. The electrodeposited material was carefully scraped off using a razor blade and the material was submitted for microanalysis. The elemental composition for a sample prepared as above was as follows: Ni, 43.6 wt.%; H, 2.16 wt.%; B, 2.7 wt.%; and K, 1.1 wt.%. Although an idealized catalyst formula of Ni²⁺O₅(OH)₂/₃(H₄BO)₁/₃ • 1.5H₂O matches these values, it is unlikely that the composition of a dry film corresponds exactly to that of a film under operational conditions.

**Scanning Electron Microscopy.** SEM micrographs were obtained with a JSM-5910 microscope (JEOL). Following electrodeposition, catalyst samples were rinsed with deionized water and allowed to dry in air before loading into the instrument. Images were obtained with an acceleration voltage of 5–10 kV. Fig. S3 displays SEM images of a catalyst prepared by passing 10 C/cm² at 1.3 V.

**Powder X-Ray Diffraction.** A powder X-ray diffraction pattern for a film grown by passing 10 C/cm² was obtained with a Rigaku RU300 rotating anode X-ray diffractometer (185 mm) using Cu Ka radiation (λ = 1.5405 Å). As shown in Fig. S4, the only peaks in the diffraction pattern correspond to those pertaining to the ITO background, indicating that the electrodeposited nickel oxide catalyst is amorphous.

**Spectroelectrochemistry.** Spectra were recorded on a Spectral Instruments 400 series diode array spectrometer. The working electrode consisted of a 2 × 0.8 cm piece of ITO-coated quartz cut from a commercially available slide (Delta Technologies, Inc.). Working, reference, and Pt auxiliary electrodes were fitted into a standard 1-cm path-length UV-Vis cuvette to comprise a one-compartment electrolysis cell. The spectrometer was blanked against a filtered solution of B₃ electrolyte containing Ni²⁺ (1 mM) and spectra were collected periodically while 1.2 V was applied. The spectrum recorded after 9 min of electrolysis is shown in Fig. S7.

**Determination of Faradaic Efficiency.** An Ocean Optics oxygen sensor system was used to detect O₂ quantitatively. The experiment was performed in a custom-built two-compartment gas-tight electrochemical cell with a 14/20 port on each compartment and a Schlenk connection with a Teflon valve on the working compartment. The B₃ electrolyte was degassed by bubbling with high purity N₂ for 12 h with vigorous stirring, and it was transferred to the electrochemical cell under N₂. One compartment contained a Ni foam auxiliary electrode and the other compartment contained the working and Ag/AgCl reference electrodes. The Ni catalyst was prepared from an electrodeposition as described above. The reference electrode was positioned several millimeters from the surface of the catalyst. The 14/20 port of the working compartment was fitted with a FOXY OR125–73 mm O₂ sensing probe connected to a MultiFrequency Phase Fluorometer. The phase shift of the O₂ sensor on the FOXY probe, recorded at 10 s intervals, was converted into the partial pressure of O₂ in the headspace using a two-point calibration curve (air, 20.9% O₂, and high purity N₂, 0% O₂). After recording the partial pressure of O₂ for 1 h in the absence of an applied potential, electrolysis was initiated at 1.3 V without iR compensation.

For determination of Faradic efficiency in the B₃ buffer, electrolysis with O₂ sensing was continued until 53.5 C passed. Upon terminating the electrolysis, the O₂ signal reached a plateau over the course of the next 3 h. During this time, the O₂ level had risen from 0% to 6.98%. At the conclusion of the experiment, the volume of the solution (59.5 mL) and the volume of the headspace (48.0 mL) in the working compartment were measured. The total charge passed in the electrolysis was divided by 4 F to get a theoretical O₂ yield of 138.3 μmol. The measured partial pressure of O₂ was corrected for dissolved O₂ in solution using Henry’s Law and converted, using the ideal gas law, into a measured O₂ yield of 143.4 μmol (103.7% ± 5%) (Fig. S5).

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