

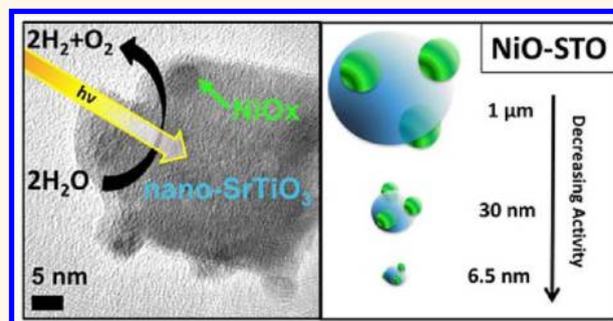
Nanoscale Strontium Titanate Photocatalysts for Overall Water Splitting

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The photocatalytic water splitting reaction is the most promising process to convert photochemical energy from the sun directly into high purity chemical fuel (hydrogen). According to a recent DOE report, suspended heterogeneous photocatalysts operating at 10% solar energy conversion efficiency are theoretically able to produce hydrogen fuel at a cost of \$1.63/kg H₂.¹ This is considerably cheaper than hydrogen produced from photoelectrochemical (PEC) cells or from the combination of photovoltaic (PV) cells with water electrolyzers. Inorganic nanomaterials are particularly interesting as photocatalysts for the water splitting reaction. Because of their small size, full light penetration can be achieved, and even short-lived charge carriers can reach the interface. The positive effects of nanoscaling on photocatalytic activity have been documented for IrO₂,² Fe₂O₃,^{3–7} MnO₂,^{8–10} all of which have been shown to photooxidize water in the presence of an electrical or chemical bias. A disadvantage of nanomaterials is that on this size scale, space charge layers are not effective for separating electron hole pairs, and recombination is enhanced in the absence of a bias. This is one of the reasons that nanoscale catalysts for overall water splitting are very rare. So far only three systems have been reported in the literature, all of which require deep ultraviolet light ($\lambda < 300$ nm) for operation. Kudo's group showed that LiNbO₃ nanowires (70 nm \times 10 μ m) can split water after modification with RuO₂ cocatalyst particles. At 254 nm, the quantum yield was estimated as 0.7%.¹¹ Yan *et al.* reported overall water splitting with RuO₂-modified Zn₂GeO₄ nanorods (100 \times 150 nm) under UV light from a 400 W Hg UV lamp to excite the large bandgap (>4.5 eV) of the material.¹² In 2011, Domen's group reported overall water splitting with aggregates of NiO-loaded NaTaO₃ nanocrystals (20 and 40 nm), but deep

ABSTRACT



SrTiO₃ (STO) is a large band gap (3.2 eV) semiconductor that catalyzes the overall water splitting reaction under UV light irradiation in the presence of a NiO cocatalyst. As we show here, the reactivity persists in nanoscale particles of the material, although the process is less effective at the nanoscale. To reach these conclusions, Bulk STO, 30 \pm 5 nm STO, and 6.5 \pm 1 nm STO were synthesized by three different methods, their crystal structures verified with XRD and their morphology observed with HRTEM before and after NiO deposition. In connection with NiO, all samples split water into stoichiometric mixtures of H₂ and O₂, but the activity is decreasing from 28 μ mol H₂ g⁻¹ h⁻¹ (bulk STO), to 19.4 μ mol H₂ g⁻¹ h⁻¹ (30 nm STO), and 3.0 μ mol H₂ g⁻¹ h⁻¹ (6.5 nm STO). The reasons for this decrease are an increase of the water oxidation overpotential for the smaller particles and reduced light absorption due to a quantum size effect. Overall, these findings establish the first nanoscale titanate photocatalyst for overall water splitting.

KEYWORDS: photocatalyst · photolysis · complete water splitting · solar fuel · nanoscale

ultraviolet light ($\lambda < 300$ nm) was necessary for catalyst operation.¹³ These studies are very encouraging, but significant improvement of the efficiency and an extension of the light absorption range will be required before these systems can ever be useful for true solar energy to fuel conversion. This will require a deeper understanding of the interplay of energetics and the kinetics in such catalysts. Here, we describe such relationships for nano-NiO–SrTiO₃ (NiO–STO)—the first nanoscale titanate photocatalyst for overall water splitting. With its 3.2 eV bandgap, STO allows considerably better

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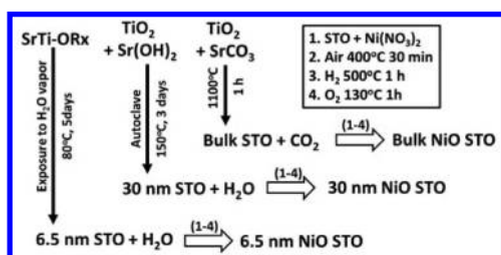
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light absorption than the germanates, tantalates, and niobates mentioned above. Another desirable feature of the NiO–STO system is that no rare elements are required for function. Consequently, there is significant interest in this catalyst type.^{14–28} We find that 30 nm sized NiO–STO composites can split water with an activity only slightly lower ($19.4 \mu\text{mol H}_2 \text{g}^{-1} \text{h}^{-1}$) than that of bulk NiO–SrTiO₃ ($28 \mu\text{mol H}_2 \text{g}^{-1} \text{h}^{-1}$). The estimated quantum efficiency is 0.041% at 315 nm (based on 50 mg of the catalyst). When the size of the titanate particles is reduced further, the activity drops significantly, due to the combination of a quantum size effect leading to lower light absorption, and due to slower water oxidation kinetics. These results establish a lower size limit (30 nm diameter) for photocatalysts based on SrTiO₃, and suggest that improvements of the activity should target water oxidation kinetics and charge separation in this system.

RESULTS AND DISCUSSION

Syntheses of the three types of STO (bulk, 30 nm, and 6.5 nm) are summarized in Scheme 1. Bulk STO was



Scheme 1. Synthesis of Bulk NiO STO, 30 nm NiO STO and 6.5 nm NiO STO with reduction/oxidation pre-treatment to activate the NiO_x ($0 < x < 1$) cocatalyst.

prepared from TiO₂ and SrCO₃ using a high temperature solid-state reaction. As the TEM in Figure 1A shows, the material forms 0.1–1.0 μm particles with ill-defined surface features and of likely polycrystalline structure.²⁹ Nanoscale STO particles were synthesized through a hydrothermal synthesis from TiO₂, Sr(OH)₂, and KOH.³⁰

According to TEM (Figure 1B), particles are mostly of cubic appearance with edge lengths of 25–60 nm (average value of 30 nm). A third variety of STO was obtained by exposing 10 mL of 0.7 M solution of the bimetallic precursor (SrTi-(OCH₂CH(CH₃)OCH₃)₆ in *n*-butanol/3-methoxy-propanol) to 100 mL water vapor under controlled conditions.³¹ This produces agglomerated 6.0 ± 1.0 nm STO nanocrystals (Figure 1C), henceforth referred to as 6.5 nm STO.

The crystal structures of the materials were verified with powder X-ray diffraction (Figure 2). All samples are phase pure, based on the diffraction patterns. For 30 nm STO and 6.5 nm STO, crystal domain sizes were estimated from the diffraction peak widths (at half-maximum) using the Scherrer equation. The calculated values of 30 and 6.5 nm agree well with the TEM images of these compounds. To obtain viable catalysts for overall water splitting, NiO was deposited on each type of STO, according to Scheme 1, by impregnating the solids in aqueous Ni(NO₃)₂, followed by calcination in air (400 °C), hydrogen (500 °C), and then oxygen atmosphere (130 °C) at the indicated temperatures.

TEM images of bulk NiO STO, 30 nm NiO STO, and 6.5 nm NiO STO are shown in Figure 1D–F. Z-contrast imaging for bulk STO verified the presence of NiO_x ($0 \leq x \leq 1$) cocatalyst particles due to the darker contrast. In some cases, Ni(0) particles could also be identified,

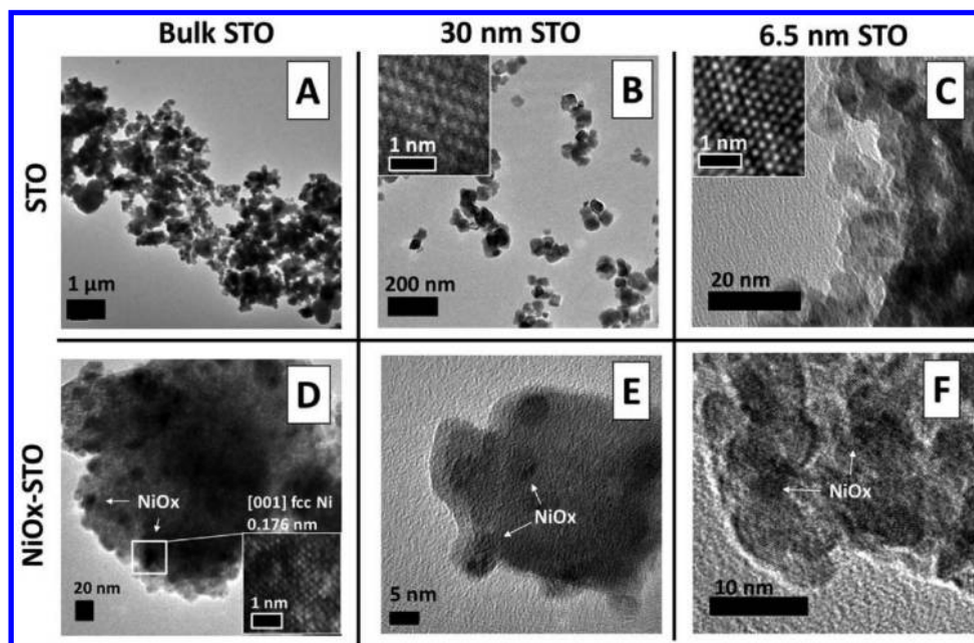


Figure 1. HRTEMs of Bulk STO (A), 30 nm STO (B), 6.5 nm STO (C), bulk NiO STO with inset of Ni particle (D), 30 nm NiO STO (E), and 6.5 nm NiO STO (F).

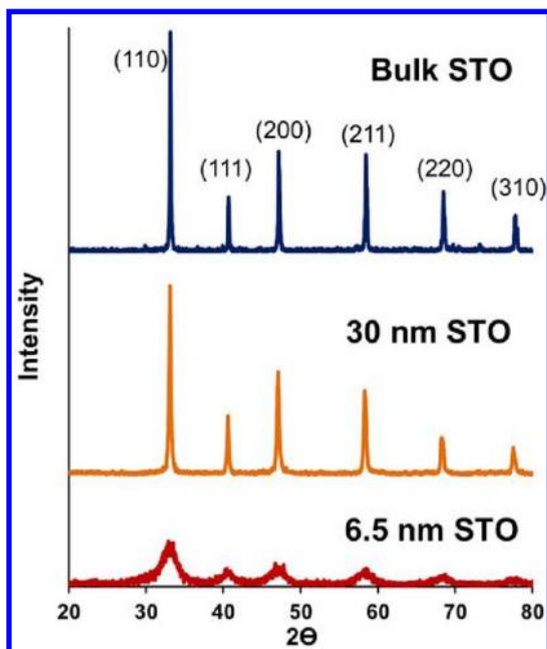


Figure 2. Powder X-ray diffraction patterns of bulk STO, 30 nm STO, and 6.5 nm STO.

based on atomic spacing (Figure 1D). The NiO_x sizes are between 10 and 50 nm (Figure 1D) for bulk STO and 5–10 nm for 30 nm STO (Figure 1E). The decrease in NiO_x particle size is expected from heterogeneous nucleation theory, which predicts an increase in nucleation sites on the larger surface of the smaller STO particles, and consequentially, smaller NiO particles at the end of the growth period. No discrete NiO particles could be observed for the 6.5 nm STO sample (Figure 1F) due to low contrast and agglomeration.

The attachment of NiO_x can also be seen in the optical properties of the materials (Figure 3). NiO_x deposition is accompanied by color change (insert in Figure 3A) due to the broad visible light absorption of nickel and the ultraviolet absorption of NiO (3.2 eV). The smallest 6.5 nm NiO STO appears dark black, 30 nm NiO STO is dark gray, and bulk NiO STO appears light gray (Figure 3 photos). In contrast, the Ni-free materials are white powders with no absorption in the visible. According to the Tauc plots in Figure 3B,C, bulk STO has optical transitions related to a direct bandgap at 3.3 eV and to an indirect bandgap at 3.2 eV. The indirect transition was previously assigned to O 2p and Ti 3d states and the direct transition to O 2p to Sr 4d states.³² 30 nm STO and 6.5 nm STO also show direct and indirect transitions, but all occur at higher energy (at 3.3–3.7 eV) than for the bulk form. This indicates an incipient quantum size effect in the smaller particles. Such an effect would be expected from the value of the Bohr-exciton radius for SrTiO₃ (6.25 nm).³³ In 6.5 nm STO the increase of bandgap is partially obscured by an absorbance tail in the visible region (see Figure 3C). This tail appears to be a result of midgap defect states

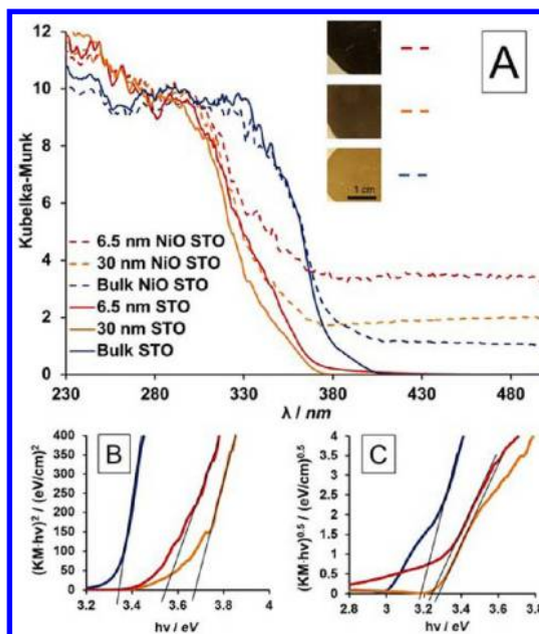


Figure 3. Kubelka–Munk diffuse reflectance of unmodified STO powders and NiO_x attached STO catalysts (A). Tauc plot for allowed direct transitions (B) and allowed indirect transitions (C) of the unmodified STO catalysts.

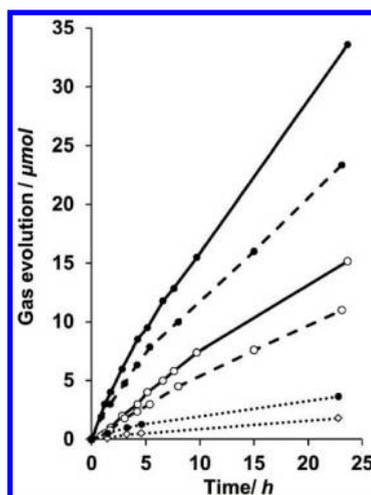


Figure 4. H₂ (●) and O₂ (○) evolution from bulk NiO STO (solid line), 30 nm NiO STO (dashed line), and 6.5 nm NiO STO (dotted line) in 50 mL of water at pH = 7 under full spectrum irradiation ($\sim 26.3 \text{ mW cm}^{-2}$ in the UV from 250 to 380 nm). The catalyst amount was 50 mg (272 μmol based on SrTiO₃). Rates were determined for the 5–24 h time period, when H₂ evolution was linear.

that are due to the greater specific surface area (smaller crystal size) of this material. Crystal defects are also due to the low synthesis temperature (80 °C, compared to 150 and 1100 °C for the other materials).

To assess catalytic activity of the three compounds, equal amounts of these catalysts (50 mg) were suspended in pure water and irradiated with a 300 W Xe arc lamp. Stoichiometric H₂ and O₂ evolution was observed in all cases (Figure 4). Rates slightly decreased after 5 h, but remained stable for the duration

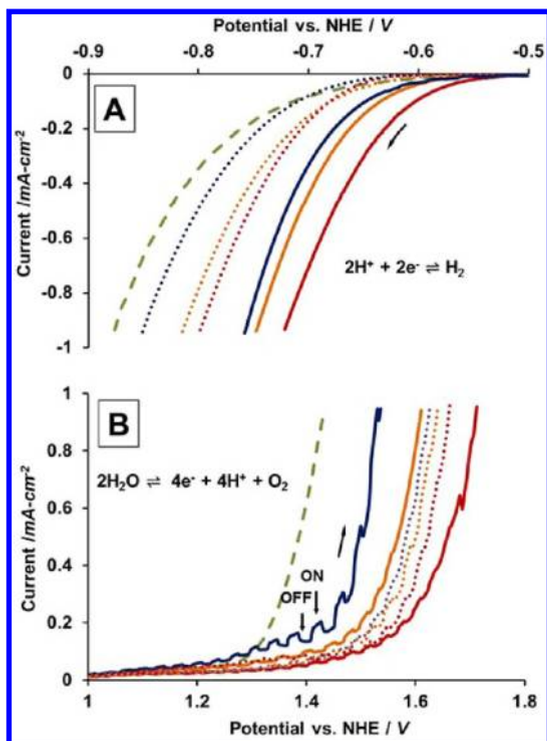


Figure 5. Electrochemical scans (10 mV s^{-1}) on NiO_x -STO films on gold in $0.25 \text{ M Na}_2\text{HPO}_4/\text{NaH}_2\text{PO}_4$ buffer ($\text{pH} = 7$). All potentials are vs NHE. Chopped light during the anodic scans (B) reveals small photocurrents. Dotted blue = bulk STO, solid blue = bulk NiO STO, dotted orange = 30 nm STO, solid orange = 30 nm NiO STO, dotted red = 6.5 nm STO, solid red = 6.5 nm NiO STO, dashed green = plain NiO .

of the experiments (24 h). Bulk NiO STO had the highest photocatalytic activity ($28 \mu\text{mol g}^{-1} \text{ h}^{-1}$), followed by 30 nm NiO STO ($19.4 \mu\text{mol g}^{-1} \text{ h}^{-1}$), and 6.5 nm NiO STO ($3.0 \mu\text{mol g}^{-1} \text{ h}^{-1}$) (Figure 4). Reference experiments using a lower ($1.5 \text{ wt } \%$) and a higher ($20 \text{ wt } \%$) NiO loading on 30 nm NiO -STO resulted in lower water splitting activity (12 and $6.5 \mu\text{mol H}_2 \text{ g}^{-1} \text{ h}^{-1}$, respectively), and no activity for the 6.5 nm NiO STO sample. Therefore, all experiments were conducted with samples containing $3 \text{ wt } \%$ NiO .

To determine the reasons for the decreasing activity with decreasing size, the electrochemical overpotentials of the catalysts for water oxidation and water reduction were measured. Potential–current scans of films of the NiO_x -free and NiO_x -loaded catalysts on gold foil in 0.1 M KCl solution at $\text{pH} = 7$ are shown in Figure 5.

Under these conditions, the proton reduction potentials occur in the following order: Bulk STO (-0.85 V) $>$ 30 nm STO (-0.81 V) $>$ 6.5 nm STO (-0.80 V); that is, the smallest particles reduce protons the easiest. Attachment of NiO_x cocatalysts moves all proton reduction overpotentials to lower values (by 6 – 9 mV), but the overall order remains the same. Interestingly, water oxidation potentials show an inverted trend: 6.5 nm STO ($+1.66 \text{ V}$) $>$ 30 nm STO ($+1.64 \text{ V}$) $>$ bulk STO ($+1.62 \text{ V}$); that is, here the bulk material oxidizes

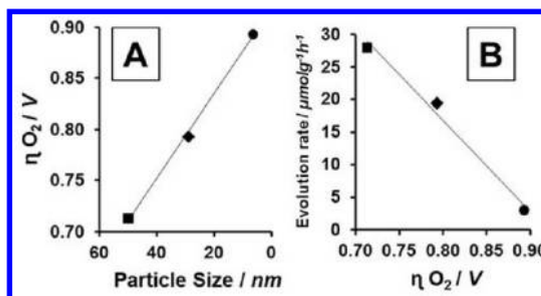


Figure 6. (A) Water oxidation overpotential versus particle size; (B) water oxidation potential versus gas evolution rate. Squares, bulk NiO STO; diamonds, 30 nm NiO STO; circles, 6.5 nm NiO STO. Overpotentials are calculated on the basis of the theoretical water oxidation potential of $+0.817 \text{ V}$ (NHE), at $\text{pH} 7$.

water the easiest. Again, NiO_x attachment does reduce the overpotentials slightly, but does not change the relative reactivity order. This suggests that NiO_x may be directly involved in the water oxidation reaction. This question is currently under investigation in the laboratory.

At present, the reason for the variation of redox potentials with SrTiO_3 particle size is unclear. One possible explanation is a increase of the SrTiO_3 surface acidity with decreasing particle size, as has been recently reported for nanoscale TiO_2 particles.³⁴ Higher surface acidity would translate into higher proton coverage (at fixed solution pH), and lower proton reduction potential (higher water oxidation potential). This would explain the trend observed above.

Figure 6 shows plots of the particle size against water oxidation overpotentials η_{ox} and against photocatalytic H_2/O_2 evolution rates R . The correlation between η_{ox} and R is evident. This proves that water oxidation kinetics are limiting the catalyst performance under illumination. This result is somewhat surprising because photochemical and electrochemical water oxidation proceed by different mechanisms. The former is driven by the photoholes in the valence band, whereas the latter is driven by empty states in the conduction band. The correlation between η_{ox} and R suggests that both routes proceed *via* similar reaction intermediates (*e.g.*, hydroxyl radicals), and that the energetics of those intermediates are limiting the water oxidation rate. Indeed the trend observed here is not unique. We have previously reported that the photocatalytic H_2 and O_2 evolution rates in nanomaterials can be correlated with the respective electrochemical overpotentials.^{35–37}

The photoenergetics of the three SrTiO_3 materials were also investigated, employing photoelectrochemical scans on films in aqueous 0.1 M KCl solution that contained 10% (vol) methanol. Corresponding data is shown in Figure 7 and listed in Table 1. For bulk STO, photocurrents were found to be small ($10 \mu\text{A cm}^{-2}$), but exhibited a clear onset potential E_{ph} at -0.87 V vs NHE. For n-type semiconductors, including SrTiO_3 , the

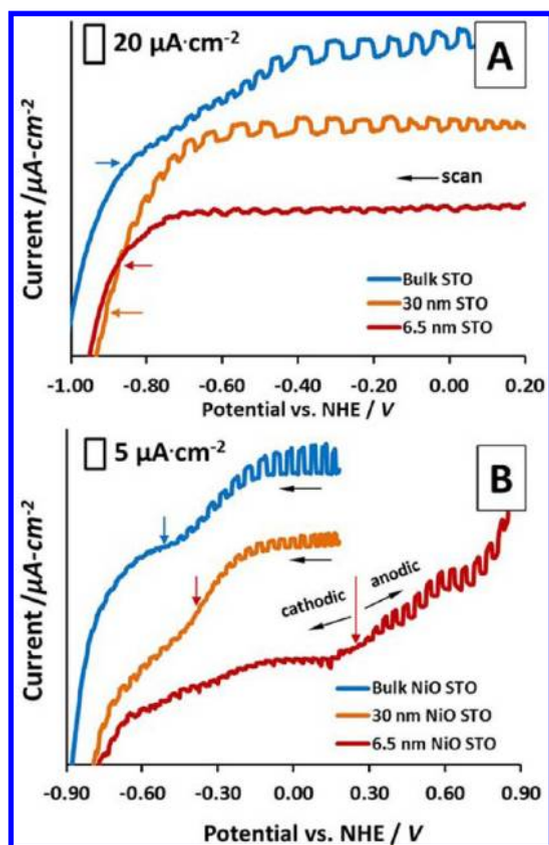


Figure 7. Photocurrent scans of bulk STO, 30 nm STO, 6.5 nm STO (A), and bulk NiO STO, 30 nm NiO STO, 6.5 nm NiO STO (B) films on Au substrate, 10 mV s^{-1} , at $\text{pH} = 7$ in $0.1 \text{ M KCl}/10\% \text{ vol. methanol}$. Arrows indicate photo-onset potentials E_{ph} . In all samples, a reversal of the photocurrent direction occurs at $E_{\text{applied}} < E_{\text{ph}}$.

TABLE 1. Electrochemical Water Reduction/Oxidation Potentials and Photo-onset Potential E_{ph} (V vs. NHE at 0.90 mA cm^{-2}) with Band Gap Energy, E_{G} (eV)

| catalyst | proton reduction | | water oxidation | |
|----------------|------------------|---------------|---------------------|---------------------|
| | potential (V) | potential (V) | E_{ph} (V) | E_{G} (eV) |
| bulk STO | −0.85 | +1.62 | −0.87 | 3.32 |
| 30 nm STO | −0.81 | +1.64 | −0.94 | 3.67 |
| 6.5 nm STO | −0.80 | +1.66 | −0.90 | 3.55 |
| bulk NiO STO | −0.76 | +1.53 | −0.49 | 3.32 |
| 30 nm NiO STO | −0.75 | +1.61 | −0.36 | 3.67 |
| 6.5 nm NiO STO | −0.72 | +1.71 | +0.27 | 3.55 |

photo-onset potential E_{ph} serves as an approximation of the conduction band edge. Indeed the experimental value compares well with the literature for the conduction band edge of SrTiO_3 .³⁸ For 30 nm STO the onset potential value was shifted in the negative direction (−0.94 V). A shift of E_{CB} to more reducing potentials can be taken as the second manifestation of the quantum size effect, in addition to the blue-shifted absorption in the optical spectra. For 6.5 nm STO, a E_{ph} value similar to that of bulk STO is found (−0.90 V). However, the photocurrent for this material is very

weak, possibly obscuring a more negative value, closer to that of 30 nm STO.

When NiO_x cocatalysts are fused onto the STO particles, photocurrents get smaller ($< 5 \mu\text{Acm}^{-2}$) and photo-onsets shift to more positive values for all catalysts (Figure 7). For bulk NiO STO, E_{ph} is at −0.49 V, and for 30 and 6.5 nm, NiO–STO values are found at −0.36 and +0.27 V, respectively. These very positive shifts are due to Fermi level equilibration between STO and NiO. Significant, but smaller (60–240 mV) positive shifts of the photo-onset potentials were previously found in nanoscale niobates and titanates after the binding of Pt and IrO_x nanoparticles.^{35,39} Nickel oxide is a p-type semiconductor with a 3.31 eV band gap and a flatband potential of +0.67 V (NHE).⁴⁰ The experimental E_{ph} values for bulk NiO STO fall between those observed for NiO and STO alone. Due to the p-type character of NiO, the new E_{ph} values can no longer be taken as the conduction band edge position of STO, but must be interpreted as the common Fermi level of the NiO–STO system. Another feature of the electrochemical scans is an inversion from anodic to cathodic photocurrents that occurs negative of the E_{ph} potentials. This inversion is found for all samples, but it is most pronounced for the 6.5 and 30 nm STO particles. In small particles, the space charge layer is not effective for separating charge,^{41–43} and so the sign of the photocurrent is due to other factors. In the present case, we hypothesize that the direction of photocurrent is determined by the polarity of the electrical double layer at the semiconductor electrode. Further studies on the photocurrent inversion effect are underway.

CONCLUSION

We have described overall photocatalytic water splitting with bulk and two types of nanoscale $\text{NiO}_x\text{–SrTiO}_3$ particles. All materials perform under UV light, but the 30 nm particles are about 35% less active than the bulk, and 6.5 nm have only 10% of the activity. The decrease of activity is attributed to a combination of factors; the most important ones are reduced light absorption due to the beginning of a quantum size effect in the 30 and 6.5 nm samples, and an observed increase in water oxidation potentials with the smaller particles. Thus, for full activity under UV light, catalyst particles should be larger than 30 nm. Also, the significant overpotentials for water reduction and oxidation suggest that optimization of activity with SrTiO_3 could be achieved by adding more effective cocatalysts. The positive effect of the NiO_x coating on both water reduction and oxidation potentials implies that NiO_x is directly involved in both reactions. Additional studies on the detailed mechanism of water splitting with NiO–SrTiO_3 are underway in this laboratory. Lastly, we note that 30 nm NiO SrTiO_3 is only the

fourth nanoscale catalyst for overall water splitting, apart from RuO₂–LiNbO₃ nanowires (70 nm × 10 μm),¹¹ RuO₂–Zn₂GeO₄ nanorods (100 × 150 nm),¹² and NiO–NaTaO₃ nanocrystals (20 and 40 nm).¹³ Owing

to its smaller bandgap of 3.2 eV, nano NiO–STO is active under near UV conditions, whereas the other nanoscale catalysts require deep UV light (<300 nm) for operation.

METHODS

Chemicals. KOH 99.9%, Sr(NO₃)₂ 99.9%, Degussa P25 TiO₂, NH₄OH 99.9%, Ni(NO₃)₂·(H₂O)₆ 99.9%, and ammonium hydroxide were purchased from Fisher Scientific, Pittsburgh, PA. Strontium titanium bimetallic alkoxide (SrTi-(OCH₂CH(CH₃)-OCH₃)₆; 0.7 M in *n*-butanol/3-methoxypropanol; Gelest, Inc.) was used as received. Sr(OH)₂ was prepared by precipitation at room temperature from Sr(NO₃)₂ and KOH in water. They were of reagent quality and were used as received. Water was purified to 18 MΩcm resistivity using a *Nanopure* system.

Particulate Synthesis. Bulk-SrTiO₃ particles were synthesized *via* a high temperature solid state process²⁹ where 0.021 mol, 1.74 g of P25 TiO₂ was mixed and sonicated for 10 min with 0.026 mol, 5.53 g Sr(NO₃)₂ in a 1:1.2 molar ratio in 200 mL of water to produce precursors for 4 g of SrTiO₃. 54 mL of aqueous oxalic acid (0.4 M, 1:1 molar ratio with Sr) was added dropwise to the solution under vigorous stirring. Concentrated ammonium hydroxide was added slowly to increase the pH of the solution to 6.5 in order to precipitate strontium oxalate crystals onto the 25 nm TiO₂ particle surfaces *via* heterogeneous nucleation.⁴⁴ When precipitation was complete, the white solid was centrifuged and washed eight times in 50 mL of water, followed by drying in air at 100 °C. This precursor was calcined in a Thermolyne 79300 Tube Furnace to 1100 °C for 1 h with a heating rate of 10 °C min⁻¹. After being cooled to room temperature, the white solid was washed twice in 50 mL of 5 M HNO₃ to remove excess SrCO₃ followed by repeated water washes until the supernatant reached a pH of 7. Phase purity was confirmed by powder X-ray diffraction.

Nano-SrTiO₃ particles (30 nm) were synthesized *via* a hydrothermal autoclave route.³⁰ TiO₂ (0.18 g, 2.3 mmol) was mixed in a 20 mL aqueous solution of 1.26 g, 23 mmol of KOH and 0.508 g, 2.3 mmol of Sr(OH)₂ in a 30 mL Teflon container sealed in a pressure vessel. This mixture was heated to 150 °C for 3 days. The vessel was cooled to room temperature, and the formed white powder was washed with water until the supernatant reached a pH of 7 and then dried at 100 °C in air.

Nano-SrTiO₃ particles (6.5 nm) were synthesized following a vapor diffusion sol–gel technique³¹ where 10 mL of a 0.7 M solution of SrTi-(OCH₂CH(CH₃)OCH₃)₆ in *n*-butanol/3-methoxypropanol (Gelest, Inc.) was heated under N₂ atmosphere at 80 °C, exposed to water vapor from an adjacent flask of water (at 25 °C) for 72 h. The resulting off-white gel was washed in ethanol and dried at 100 °C to yield 6.5 nm STO.

NiO Loading. The SrTiO₃ particles (200 mg, 1.10 mmol) were added to 20 mL of an aqueous Ni(NO₃)₂·6H₂O solution (0.023 g, 3 wt % loading of NiO:STO) and thoroughly mixed in a sonication bath for 10 min. This solution was dried at 100 °C and then calcined for 30 min at 400 °C in air. The air above the solid was flushed with N₂ and then replaced by H₂, and the sample was heated to 500 °C for 1.5 h to reduce nickel. The final product was obtained after another 30 min of heating at 130 °C in an O₂ atmosphere as described in the literature.⁴⁵

High-resolution transmission electron microscopy (HRTEM) images were taken using a JEOL 2500SE 200 kV TEM. To prepare samples, copper grids with a carbon film were dropped into aqueous dispersions of the samples followed by washing with water and air drying. UV–vis diffuse reflectance spectra were recorded on dried powders on white Teflon tape using a Thermo Scientific Evolution 220 spectrometer. The reflectance data were converted to the Kubelka–Munk function as $f(R) = (1 - R)^2(2R)^{-1}$ versus wavelength to correct for scattering, which is plotted in Figure 3. For electrochemical measurements, thin films of the catalysts were prepared on a gold foil electrode (1.0 cm²) by drop coating and drying at 25 °C. A wire was

attached to the bare gold back with conductive carbon tape and sealed with adhesive. The electrode was placed into a N₂-purged 3-electrode cell with a Pt counter electrode and a saturated calomel reference electrode connected to the cell with a KCl salt bridge. The cell was filled to 50 mL with 0.25 M Na₂HPO₄/NaH₂PO₄ buffer solution at pH 7 and purged with N₂ to remove oxygen. Cyclic voltammetry scans were taken at 50 mV/s. The system was calibrated using the redox potential of K₄(Fe(CN)₆) at +0.358 V (NHE). Photoelectrochemical scans were conducted in 0.1 M KCl with 10% (vol) added methanol. The rate of photochemical hydrogen and oxygen evolution from each catalyst was determined by irradiating 50 mg of NiO–STO dispersed in 50 mL of water. Irradiations were performed in a quartz round-bottom flask with a 300 W Xe arc lamp (26.3 mW cm⁻² at the flask λ = 250–380 nm), measured with a GaN photodetector. The airtight irradiation system connects a vacuum pump and a gas chromatograph (Varian 3800) with the sample flask to quantify the amount of gas evolved, using area counts of the peaks and the identity of the gas from the calibrated carrier times. Prior to irradiation, the flask was evacuated down to 5 Torr and purged with argon gas. This cycle was repeated until the chromatogram of the atmosphere above the solution indicated that the sample did not contain hydrogen, oxygen, or nitrogen.

Powder XRD scans were conducted with a Scintag XRD, at a wavelength of λ = 0.154 nm with 2 mm tube slit divergence, 4 mm scatter, 0.5 mm column scatter, and 0.2 mm receiving widths. The full width at half-maximum (fwhm) values of the peaks were calculated with Thermo ARLDMSNT software (version 1.39-1).

Conflict of Interest: The authors declare no competing financial interest.

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